Comments About This draft

This is a set of course notes hoping to someday be a book.

Unfortunately, there is a huge difference between course notes and a book. This is why I need everyone’s help. If there are parts that you think are unclear — please let me know. If there are errors — please let me know (even if they are small and subtle). If the figures are unclear — please let me know. If there are mistakes in grammar — please let me know.

If you don’t get the jokes... well, that is your problem. Seriously though, I need help if this is eventually going to arrive at the Nirvana that is bookdom. Give me feedback please. It will be good for your Karma. 😊
Some thoughts about this book

This book originated as part of a lecture course given at Oxford in the fall of 2016 and then again in 2017, 2018, 2019, 2020, ... and this kept going until I finished the book, which seemed like forever.

The idea of this book is to give a general introduction to topological quantum ideas. This includes topological quantum field theories, topological quantum memories, topological quantum computing, topological matter and topological order — with emphasis given to the examples of toric code, loop gases, string nets, and particularly quantum Hall effects. The book is aimed at a physics audience (i.e., we avoid the language of category theory like the plague!), although some mathematicians may also find the perspectives presented here to be useful.

How to read this book

The book was originally written to be read roughly sequentially. However, you may be able to jump around quite a bit depending on your interests. When the toric code is introduced, it is quite independent of the prior chapters on the general structure of TQFTs. In the course I teach, I am certainly not assigning all of the chapters — I’m not a sadist!

I should also mention that chapter 41 introduces some basic mathematics that many people may know but I thought should be included for completeness.

There are often small hitches and caveats that are swept under the rug in the name of simplifying the discussion. I try to footnote these caveats when they occur. Many technical details are pushed to chapter appendices — often these can be skipped on a first reading.

In a margin note of my previous book (Simon 2013), I said that my next book (i.e., this one) would be about two dimensional electron systems. This topic is covered in the section on fractional quantum Hall effect1.

A list of useful references is given etc.

---

1I also suggested that I might write a thriller about physicists defeating drug smugglers. For those who are interested, I’m still working on it, but I discovered that writing a novel is pretty hard.
Contents

1 Introduction and Ancient History 1

2 Kauffman Bracket Invariant and Relation to Physics 5
  2.1 The Idea of a Knot Invariant 5
  2.2 Relation to Physics 8
    2.2.1 Twist and Spin-Statistics 11
    2.2.2 Blackboard Framing 12
  2.3 Bras and Kets 13
  2.4 Quantum Computation with Knots 15
  2.5 Some Quick Comments about Fractional Quantum Hall Effect 16
  2.6 Appendix: More Knot Theory Basics 18
    2.6.1 Isotopy and Reidemeister Moves 18
    2.6.2 Writhe and Linking 18
Exercises 19

I Anyons and Topological Quantum Field Theories 21

3 Particle Quantum Statistics 23
  3.1 Single Particle Path Integral 23
  3.2 Two Identical Particles 25
  3.3 Many Identical Particles 27
    3.3.1 Paths in 2+1 D, the Braid Group 28
    3.3.2 Paths in 3+1 D, the Permutation Group 29
    3.3.3 Building a Path Integral 30
  3.4 Abelian Examples 31
    3.4.1 3+1 Dimensions 31
    3.4.2 2+1 Dimensions 32
  3.5 Nonabelian Case 32
    3.5.1 Parastatistics in 3+1 Dimensions 34
Exercises 35

4 Aharanov-Bohm Effect and Charge-Flux Composites 39
  4.1 Review of Aharanov-Bohm Effect 39
  4.2 Anyons as Charge-Flux Composites 41
    4.2.1 Fusion of Anyons 42
    4.2.2 Anti-Anyons and the Vacuum Particle 42
  4.3 Anyon Vacuum on a Torus and Quantum Memory 43
8.4 Application of Fusion: Dimension of Hilbert Space on 2-Manifolds 94
8.5 Product Theories 97
8.6 Appendix: Tensor Description of Fusion and Splitting Spaces 98
Exercises 100

9 Change of Basis and $F$-Matrices 103
9.1 Example: Fibonacci Anyons 104
9.2 Example: Ising Anyons 106
9.3 Pentagons 106
9.4 Gauge Transforms 107
9.5 Appendix: $F$-matrix Odds and Ends 108
9.5.1 Product Theories 108
9.5.2 Unitarity of $F$ 109
9.5.3 $F$-matrix with higher fusion multiplicities 109
Exercises 110

10 Exchanging Identical Particles 113
10.1 Introducing the $R$-matrix 113
10.1.1 Locality 115
10.2 Some Examples 116
10.2.1 Fibonacci Anyons 116
10.2.2 Ising Anyons 118
Exercises 119

11 Computing with Anyons 123
11.1 Quantum Computing 123
11.1.1 Universal Quantum Computing in the Quantum Circuit Model 124
11.2 Topological Quantum Computing 125
11.2.1 Hilbert space 125
11.2.2 Measurement (in brief) and initialization 126
11.2.3 Universal Braiding 126
11.3 Fibonacci Example 127
11.3.1 A Single Fibonacci Qubit 127
11.3.2 Topological Quantum Compiling: Single Qubit 129
11.4 Two-Qubit Gates 132
11.4.1 Controlled Gates 133
Exercises 137

III Anyon Diagrammatics (in detail) 139

12 Planar Diagrams 141
12.1 Diagrams as Operators 142
12.1.1 Stacking operators 144
12.2 Basis of States 145
12.2.1 One Particle 146
<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>12.2.2 Two Particles</td>
<td>147</td>
</tr>
<tr>
<td>12.2.3 Three Particles</td>
<td>149</td>
</tr>
<tr>
<td>12.2.4 F-Matrices Again</td>
<td>151</td>
</tr>
<tr>
<td>12.2.5 More Particles</td>
<td>152</td>
</tr>
<tr>
<td>12.3 Causal Isotopy</td>
<td>153</td>
</tr>
<tr>
<td>12.4 Summary of Planar Diagram Rules in Physics Normalization</td>
<td>155</td>
</tr>
<tr>
<td>12.4.1 A Simple Example</td>
<td>156</td>
</tr>
<tr>
<td>12.5 Appendix: Higher Fusion Multiplicities</td>
<td>157</td>
</tr>
<tr>
<td>Exercises</td>
<td>158</td>
</tr>
<tr>
<td>13 Braiding Diagrams</td>
<td>159</td>
</tr>
<tr>
<td>13.1 Three Dimensional Diagrams</td>
<td>159</td>
</tr>
<tr>
<td>13.2 Braiding Non-Identical Particles</td>
<td>160</td>
</tr>
<tr>
<td>13.2.1 Summary of Rules for Evaluating any 2+1 D Diagram with Physics Normalization</td>
<td>161</td>
</tr>
<tr>
<td>13.3 The Hexagon</td>
<td>162</td>
</tr>
<tr>
<td>13.4 R-matrix Odds and Ends</td>
<td>164</td>
</tr>
<tr>
<td>13.4.1 Appendix: Gauge Transforms and R</td>
<td>164</td>
</tr>
<tr>
<td>13.4.2 Product Theories</td>
<td>164</td>
</tr>
<tr>
<td>13.4.3 Appendix: Higher fusion multiplicities</td>
<td>165</td>
</tr>
<tr>
<td>Exercises</td>
<td>165</td>
</tr>
<tr>
<td>14 Seeking Isotopy</td>
<td>167</td>
</tr>
<tr>
<td>14.1 Isotopy Normalization of Diagrams</td>
<td>168</td>
</tr>
<tr>
<td>14.2 Gauge Choice and Frobenius-Schur Indicator</td>
<td>171</td>
</tr>
<tr>
<td>14.3 Isotopy Invariant Unitary Rules</td>
<td>173</td>
</tr>
<tr>
<td>14.3.1 Isn’t Chern-Simons Theory Isotopy Invariant and Unitary?</td>
<td>173</td>
</tr>
<tr>
<td>14.3.2 What Have We Achieved?</td>
<td>176</td>
</tr>
<tr>
<td>14.4 Impediments to Isotopy Invariance in Fusion Diagrams</td>
<td>177</td>
</tr>
<tr>
<td>14.5 Appendix: Bookkeeping Scheme with Negative d</td>
<td>178</td>
</tr>
<tr>
<td>14.6 Appendix: $\left[F^{aa}_a F\right]$ is real</td>
<td>180</td>
</tr>
<tr>
<td>14.7 Appendix: Spin 1/2 Analogy and Why We Have a Frobenius-Schur Sign</td>
<td>180</td>
</tr>
<tr>
<td>14.8 Appendix: Some Additional Properties of Unitary Fusion Categories</td>
<td>181</td>
</tr>
<tr>
<td>14.8.1 Pivotal Property</td>
<td>182</td>
</tr>
<tr>
<td>14.8.2 Spherical Property</td>
<td>183</td>
</tr>
<tr>
<td>14.9 Appendix: Higher Fusion Multiplicities</td>
<td>184</td>
</tr>
<tr>
<td>Exercises</td>
<td>185</td>
</tr>
<tr>
<td>15 Twists</td>
<td>187</td>
</tr>
<tr>
<td>15.1 Relations between $\theta$ and $R$</td>
<td>188</td>
</tr>
<tr>
<td>15.2 Appendix: Higher Fusion Multiplicities</td>
<td>190</td>
</tr>
<tr>
<td>Exercises</td>
<td>190</td>
</tr>
<tr>
<td>16 Theories with Tetrahedral Symmetry (or Full Isotopy)</td>
<td>193</td>
</tr>
<tr>
<td>16.1 Planar Diagrams</td>
<td>194</td>
</tr>
</tbody>
</table>
20 Anyons from Groups

20.1 Fusion as Group Multiplication

20.1.1 Group Cohomology

20.1.2 Simple Examples with $G = \mathbb{Z}_N$

20.1.3 Using Nonabelian Groups?

20.2 Fusion of Group Representations: $\text{Rep}(G)$

20.2.1 Example: Representations of $S_3$

20.2.2 Example: Quaternion Group $Q_8$

20.2.3 $F$-Matrices

20.2.4 Some Simple Braiding for $\text{Rep}(G)$

20.2.5 Continuous (Lie) Group Representations?

20.3 Parastatistics Revisited

20.4 Appendix: Isotopy Invariant Planar Algebras and Anyon Theories from $G = \mathbb{Z}_N$ Cohomology

20.4.1 Trivial Cocycle: $\mathbb{Z}^{(n)}_N$ Anyons

20.4.2 Nontrivial Cocycle: $\mathbb{Z}^{(n)}_{N=2}$

20.5 Appendix: Cocycles for $S_3$

20.6 Details of Working out an $F$-matrix

Exercises

21 State Sum TQFTs

21.1 Simplicial Decomposition and Pachner Moves

21.1.1 Two Dimensions

21.1.2 Three Dimensions

21.2 The Turaev-Viro State Sum

21.2.1 Proof Turaev-Viro is a Manifold Invariant

21.2.2 Some TQFT Properties

21.2.3 Connection to Chern-Simons Theory

21.3 Connections to Quantum Gravity Revisited

21.4 Dijkgraaf-Witten Model

21.4.1 Other Dimensions

21.4.2 Further Comments

Exercises

22 Formal Construction of “Chern-Simons” TQFT: Surgery and More Complicated Manifolds

22.1 Surgery

22.1.1 Simple Example of Surgery on a 2-manifold

22.1.2 Surgery on 3-manifolds

22.2 Representing Manifolds with Knots

22.2.1 Lickorish-Wallace Theorem

22.2.2 Kirby Calculus

22.3 Witten-Reshitikhin-Turaev Invariant

22.3.1 Some examples

22.3.2 Turaev-Viro Revisited: Chain-Mail and the Turaev-Walker-Roberts Theorem

Exercises
23 Anyon Condensation 301
  23.1 Condensing Simple Current Bosons 302
  23.2 Identification Step 303
    23.2.1 Orbits of maximum size 304
  23.3 Confinement Step 305
  23.4 Splitting: Orbits not of maximum size 306
  23.5 Other Features of Condensation 310
  23.6 Cosets 311
  23.7 More General Condensations 312
  23.8 Condensation and Boundary Modes 313
Exercises 314

V Toric Code Basics 317

24 Introducing Quantum Error Correction 319
  24.1 Classical Versus Quantum Information 319
    24.1.1 Memories 319
  24.2 Errors 320
    24.2.1 Classical Error Correction 320
  24.3 Quantum No Cloning Theorem 321
  24.4 Quantum Error Correction 322
    24.4.1 Bit Flip Correcting Code 322
    24.4.2 Nine Qubit Shor Code 323

25 Introducing the Toric Code 327
  25.1 Toric Code Hilbert Space 327
  25.2 Vertex and Plaquette Operators 328
    25.2.1 Operators Commute 330
    25.2.2 Is This a Complete Set of Operators? (Not quite!) 331
  25.3 Building the Code Space 332
  25.4 Errors and Error Correction 335
    25.4.1 $\sigma_x$ Errors 335
    25.4.2 $\sigma_z$ Errors 338
    25.4.3 Combinations of $\sigma_x$ and $\sigma_z$ 340
    25.4.4 Other Errors 341
  25.5 The Toric Code on Different Lattices, and Different Topologies 341
  25.6 $\mathbb{Z}_N$ Toric Code (Briefly) 343
    25.6.1 Code Space 344
    25.6.2 Errors 346
Exercises 348

26 The Toric Code as a Phase of Matter and a TQFT 349
  26.1 Excitations 349
  26.2 Statistical Properties of Vertex and Plaquette Defects 351
    26.2.1 Braiding Vertex Defect with Plaquette Defect 352
    26.2.2 Properties of $f$, the fermion 354
  26.3 $S$ and $T$ matrices 354
29.8 Appendix: Two Expressions for Ground State Degeneracy on a Torus 411
Exercises 413

30 Doubled Semion Model 415
30.1 A microscopic model 416
  30.1.1 Magnetic String Operator 417
30.2 Graphical Analysis 418
30.3 Gauge Choice and Unitary Diagram Algebra 421
30.4 Comments 423
Exercises 423

31 Levin-Wen String Net 425
31.1 Microscopic Hamiltonian 426
  31.1.1 Fat Lattice Construction 426
  31.1.2 Vertex and Plaquette Operators 427
  31.1.3 Levin-Wen Hamiltonian 428
  31.1.4 Ground State is Topological 429
31.2 Braiding of the Input Category and String Operators of the Double 430
  31.2.1 Toric Code 431
  31.2.2 Doubled Semion 432
  31.2.3 More Generally 432
31.3 Detailed Example: Doubled Fibonacci 432
  31.3.1 Excitations 434
  31.3.2 Ground State Degeneracy 435
31.4 Appendix: Explicit Form of Operators 436
  31.4.1 Plaquette Operator 436
  31.4.2 String Operator Example 437
31.5 Appendix: S-matrix for Fibonacci Anyons 438
Exercises 439

VII Symmetries and Entanglement 441

32 Anyon Permuting Symmetry 443
  32.0.1 The Idea of a Symmetry Defect 443
32.1 Symmetric Form of the Toric Code 444
32.2 Symmetry Defects in the Toric Code 446
Exercises 448

33 Topological Entanglement 449
  33.1 Entanglement in the Toric Code 450
    33.1.1 Generalizing To Arbitrary TOFTs 453
  33.2 Topological Entanglement Entropy is Robust 453
Exercises 455

34 Phases of Matter from/with Symmetry 457
  34.1 Symmetry Protection 457
## Contents

34.1.1 On-Site Symmetries 458
34.2 Example: Ising Symmetry 459

**VIII Introduction to Quantum Hall Effects** 461

### 35 Introduction to Quantum Hall — The Integer Effect 463
35.1 Classical Hall Effect 463
35.2 Two-Dimensional Electrons 463
35.3 Phenomenology of Integer Quantum Hall Effect 465
35.4 Transport in Zero Disorder 466
35.5 The Landau Problem 467
35.6 Laughlin’s Quantization Argument 471
35.6.1 Byers and Yang Theorem 471
35.6.2 Quantization of Hall Conductance 472
35.7 Edge States 473
35.7.1 Landau Gauge Edge Picture for Integer Quantum Hall 473
35.8 The Halperin Refinement of Laughlin’s Argument 474
Exercises 476

### 36 Aside: A Rapid Introduction to Topological Insulators 479
36.1 Topological Phases of Matter 479
36.1.1 Gapless Edges 481
36.2 Curvature and Chern Number 481
36.3 Symmetry Protection 482
36.4 Appendix: Chern Number is Hall Conductivity 482

### 37 Introduction to Fractional Quantum Hall Effect 485
37.0.1 Our Model Hamiltonian 487
37.1 Landau Level Wavefunctions in Symmetric Gauge 487
37.1.1 What We Want in a Trial Wavefunction 488
37.2 Laughlin’s Ansatz 489
37.2.1 Exact statements about Laughlin Wavefunction 490
37.2.2 Real Interactions 492
37.3 Quasiparticles 492
37.3.1 Quasiholes 493
37.3.2 Quasielectrons 494
37.3.3 Fractional Charge and Statistics 495
37.4 Digression on Berry’s Phase 495
37.5 Arovas-Schrieffer-Wilczek Calculation of Fractional Statistics 496
37.6 Gauge Choice and Monodromy 498
37.6.1 Fractional Statistics Calculation: Redux 500
37.7 Appendix: Building an Effective (Chern-Simons) Field Theory 501
37.8 Appendix: Quantum Hall Hierarchy 502
Exercises 504
## Contents

38 Fractional Quantum Hall Edges 507
- 38.1 Parabolic Confinement 507
- 38.2 Edges of The Laughlin State 507
  - 38.2.1 Edge Mode Field Theory: Chiral Boson 509
- 38.3 Appendix: Edges and Chern-Simons theory 510

39 Conformal Field Theory Approach to Fractional Quantum Hall Effect 511
- 39.1 The Chiral Boson and The Laughlin State 511
  - 39.1.1 Writing the Laughlin Wavefunction 513
  - 39.1.2 Quasiholes 513
- 39.2 What We Need to Know About Conformal Field Theory 515
  - 39.2.1 Example: Chiral Boson 516
  - 39.2.2 Example: Ising CFT 517
- 39.3 Quantum Hall Wavefunction Based on Ising CFT: The Moore-Read State 518
  - 39.3.1 Some Exact Statements About the Moore-Read Wavefunction 520
- 39.4 Quasiholes of the Moore-Read state 521
- 39.5 Multiple Fusion Channels and Conformal Blocks 524
- 39.6 More Comments on Moore-Read State with Many Quasiholes 527
- 39.7 Generalizing to Other CFTs 527
  - 39.7.1 $\mathbb{Z}_3$ Parafermions (briefly) 528
Exercises 530

40 Appendix: Resources for TQFTs 535

41 Some Mathematical Basics 541
- 41.1 Manifolds 541
  - 41.1.1 Some Simple Examples: Euclidean Spaces and Spheres 541
  - 41.1.2 Unions of Manifolds $M_1 \cup M_2$ 542
  - 41.1.3 Products of Manifolds $M_1 = M_1 \times M_2$ 542
  - 41.1.4 Manifolds with Boundary $M_1 = \partial M_2$ 543
  - 41.1.5 Boundaries of Manifolds $M_1 = \partial M_2$ 544
- 41.2 Groups 545
  - 41.2.1 Some Examples of Groups 545
  - 41.2.2 More Features of Groups 547
  - 41.2.3 Lie Groups and Lie Algebras 548
  - 41.2.4 Representations of Groups 549
- 41.3 Fundamental Group $\pi_1(M)$ 550
  - 41.3.1 Examples of Fundamental Groups 551

42 Commentary on References 553
The field of quantum topology inhabits a beautiful nexus between mathematics, computer science, and physics. Within the field of physics, it has been fundamental to a number of subfields. On the one hand, topology and topological matter are key concepts of modern condensed matter physics. Similarly, in the field of quantum information and quantum computation, topological ideas are extremely prominent. At the same time much of our modern study of topological matter is rooted in ideas of topological quantum field theories that developed from the high energy physics, quantum gravity, and string theory community starting in the 1980s. These earlier works have even earlier precedents in physics and mathematics. Indeed, the historical roots of topology in physics date all the way back to the 1800s which is where we will begin our story.

In 1867 Lord Kelvin and his close friend Peter Tait were interested in a phenomenon of fluid flow known as a smoke ring, configurations of fluid flow where lines of vorticity form closed loops as shown in Fig. 1.1. Peter Tait built a machine that could produce smoke rings, and showed it to Kelvin who had several simultaneous epiphanies. First, he realized that there should be a theorem (now known as Kelvin’s circulation theorem) stating that in a perfectly dissipationless fluid, lines of vorticity are conserved quantities, and the vortex loop configurations should persist for all time. Unfortunately, few dissipationless fluids exist — and the ones we know of now, such as superfluid helium at very low temperatures, were not discovered until the next century. However, at the time, scientists incorrectly believed that the entire universe was filled with a perfect dissipationless fluid, known as Luminiferous Aether, and Kelvin wondered whether one could have vortex loops in the Aether.

At the same time, one of the biggest mysteries in all of science was the discreteness and immutability of the chemical elements. Inspired by Tait’s smoke ring demonstration, Kelvin proposed that different atoms corresponded to different knotting configurations of vortex lines in the Aether. This theory of “vortex atoms” was appealing in that it gave a

---

The 2016 Nobel Prize was awarded to Kosterlitz, Thouless, and Haldane for the introduction of topological ideas into condensed matter physics. The topic of this book is a great-granddaughter of some of those ideas. In chapters 36 and 35 we will discuss some of the key works that this Nobel Prize honored.

1 See chapter 6.

---

Fig. 1.1: A smoke ring or vortex loop is an invisible ring in space where the fluid flows around the invisible ring as shown by the arrows. The whole thing moves out of the plane of the page at you as the fluid circulates.

---

4 Actually, in 1867 he was just William Thomson, but he would later be elevated to the peerage and take the name Lord Kelvin after the River Kelvin that flowed by his laboratory.
reason why atoms are discrete and immutable — on the one hand there
are only so many different knots that one can make. (See for example,
the list of the simplest few knots you can form from one piece of string
shown in Fig. 1.2.) On the other hand, by Kelvin’s circulation theorem,
the knotting of the vortices in a dissipationless fluid (the Aether) should
be conserved for all time. Thus, the particular knot could correspond
to a particular chemical element, and this element should never change
to another one. Hence the atoms should be discrete and immutable!

For several years the vortex theory of the atom was quite popular,
attracting the interest of other great scientists such as Maxwell, Kirch-
hoff, and J. J. Thomson (no relation). However after further research
and failed attempts to extract predictions from this theory, the idea of
the vortex atom lost popularity.

Although initially quite skeptical of the idea, Tait eventually came
to believe that by building a table of all possible knots (knotted config-
uration of strands such that there are no loose ends) he would gain
some insight into the periodic table of the elements, and in a remarkable
series of papers he built a catalogue of all knots with up to 7 crossings
(the first few entries of the table being shown in Fig. 1.2). From his
studies of knots, Tait is viewed as the father of the mathematical theory
of knots, which has been quite a rich field of study since that time (and
particularly during the last fifty years).

During his attempt to build his “periodic table of knots”, Tait posed
what has become perhaps the fundamental question in mathematical
knot theory: how do you know if two pictures of knots are topologically
identical or topologically different. In other words, can two knots be
smoothly deformed into each other without cutting any of the strands.
Although this is still considered to be a difficult mathematical problem,
a powerful tool that helps answer this question is the idea of a “knot
invariant” which we will study in the next chapter. Shortly, it will
become clear how this idea is related to physics.

Although Tait invented a huge amount of mathematics of the theory
of knots and developed a very extensive table of knots, he got no closer
to understanding anything about the periodic table of the atoms. In
his later life he became quite frustrated with his lack of progress in this
direction and he began to realize that understanding atoms was probably
unrelated to understanding knots. Tait died in 1901 not realizing that
his work on the theory of knots would be important in physics, albeit
for entirely different reasons.

7 Some of his conjectures were way ahead of their time — some being proven only in
the 1980s or later! See Stoimenow [2008] for a review of the Tait conjectures proven
after 1985.
8 Peter Tait was also a huge fan of golf and wrote some beautiful papers on the
trajectory of golf balls. His son, Freddie Tait, was a champion amateur golfer, being
the top amateur finisher in the British Open six times and placing as high as third
overall twice. Freddie died very young, at age 30, in the Boer wars in 1900. This
tragedy sent Peter into a deep depression from which he never recovered.
Further Reading

The purpose of this chapter is to introduce you to a few of the key ideas and get you interested in the subject!

2.1 The Idea of a Knot Invariant

**Topological equivalence.** We say two knots are topologically equivalent if they can be deformed smoothly into each other without cutting. For example, the picture of a knot (or more properly, the picture of the link of two strings) on the left of Fig. 2.1 is topologically equivalent to the picture on the right of Fig. 2.1.

It may appear easy to determine whether two simple knots are topologically equivalent and when they are not. However, for complicated knots, it becomes extremely difficult to determine whether two knots are equivalent or inequivalent. It is thus useful to introduce a mathematical tool known as a knot invariant that can help us establish when two knots are topologically inequivalent.

A **Knot Invariant** is a mapping from a knot (or a picture of a knot) to an output via a set of rules which are cooked up in such a way that two topologically equivalent knots must give the same output. (See Fig. 2.2.) So if we put two knots into the set of rules and we get two different outputs, we know immediately that the two knots cannot be continuously deformed into each other without cutting.

To demonstrate how knot invariants work, we will use the example of the Kauffman bracket invariant (See Kauffman [1987]). The Kauffman bracket is due to a common notation where one draws a picture of a knot inside brackets to indicate that one is supposed to evaluate this invariant. We will not draw these brackets.

---

1A few pieces of fine print here. (1) I am not precise about knot versus link. Strictly speaking a knot is a single strand, and a link is more generally made of multiple strands. Physicists call them all knots. In either case no dangling ends are allowed. A knot can be defined as a particular embedding of a circle ($S^1$) into a three dimensional reference manifold such as $\mathbb{R}^3$ (regular 3-dimensional space) with no self-intersections. A link is an embedding of several circles into the three dimensional manifold with no intersections. (2) When I say “topologically equivalent” here, I mean the concept of regular isotopy (See section 2.2.1 and 2.6.1). Two knots are isotopic if there is a continuous smooth family of knots between the initial knot and the final knot — however to be more precise, as we will see below in section 2.4.3, we should think of the knots as being thickened to ribbons and we want a smooth family of ribbons.

2Be warned: there are multiple things named after Kauffman. The particular normalization of the bracket invariant that we use has been named the topological bracket by Kauffman. The more common definition of the bracket is our definition divided by $d$. 
Kauffman Bracket Invariant and Relation to Physics

The Kauffman bracket invariant was essentially invented by Vaughan Jones who won the Fields medal for his work on knot theory [Jones, 1985]. Kauffman’s important contribution to this story (among his many other contributions in the field of knot theory) was to explain Jones’ work in very simple terms.

To define the Kauffman Bracket Invariant, we start with a scalar variable $A$. For now, leave it just a variable, although later we may give it a value. There are then just two rules to the Kauffman bracket invariant. First, a simple loop of string (with nothing going through it) can be removed from the diagram and replaced with the number

$$d = -A^2 - A^{-2} .$$

(2.1)

The second rule replaces a diagram that has a crossing of strings by a sum of two diagrams where these strings don’t cross — where the two possible uncrossings are weighted by $A$ and $A^{-1}$ respectively as shown in Fig. 2.3. This type of replacement rule is known as a skein rule.\(^6\)

\[^4\text{We will eventually see that } d \text{ stands for “dimension”.}\]

\[^5\text{There is a hidden assumption that an empty diagram has value 1. This means that the overall value of a diagram with a single loop is } d, \text{ the overall value of a diagram of two unlinked loops is } d^2, \text{ and so forth.}\]

\[^6\text{The word “skein” is an infrequently used English word meaning loosely coiled yarn, or sometimes meaning an element that forms part of a complicated whole (probably both of these are implied for our mathematical usage). “Skein” also means geese in flight, but I suspect this is unrelated.}\]

The general scheme is to use the second (and third) rule of Fig. 2.3 to remove all crossings of a diagram. In so doing, one generates a sum of many diagrams with various coefficients. Then once all crossings are removed, one is just left with simple loops, and each loop can just be replaced by a factor of $d$.\(^4\)

\[^4\text{We will eventually see that } d \text{ stands for “dimension”.}\]
The Idea of a Knot Invariant

Fig. 2.4 Example of evaluation of the Kauffman bracket invariant for the simple twisted loop in the upper left. The light dotted red circle is meant to draw attention to where we apply the Kauffman crossing rule (the middle line in Fig. 2.3) to get the two diagrams on the right hand side. After applying the Kauffman rules again (the final line in Fig. 2.3), we have removed all crossings and we are left only with simple loops, which each get the value $d$. In the penultimate line we have used the definition of $d$ to replace $A^2 + A^{-2} = -d$. The fact that we get $d$ in the end of the calculation is expected since we know that the original knot is just a simple loop (the so-called “unknot”) and the Kauffman rules tell us that a loop gets a value $d$. 

$$= A_{2}d^2 + d + d^3 + A^{-2}d^2$$

$$= d$$
To give an example of how these rules work we show evaluation of the Kauffman bracket invariant for the simple knot in the upper left of Fig. 2.4. The output of the calculation is that the Kauffman invariant of this knot comes out to be \( d \). This result is expected since we know that the original knot (in the upper left of the figure) is just a simple loop (the so-called “unknot”) and the Kauffman rules tell us that a loop gets a value \( d \). We could have folded over this knot many many times and still the outcome of the Kauffman evaluation would be \( d \).

The idea of a knot invariant seems like a great tool for distinguishing knots from each other. If you have two complicated knots and you do not know if they are topologically equivalent, you just plug them into the Kauffman machinery and if they don’t give the same output then you know immediately that they cannot be deformed into each other without cutting. However, a bit of thought indicates that things still get rapidly difficult for complicated knots. In the example of Fig. 2.4 we have two crossings, and we ended up with 4 diagrams. If we had a knot with \( N \) crossings we would have gotten \( 2^N \) diagrams, which can be huge! While it is very easy to draw a knot with 100 crossings, even the world’s largest computer would not be able to evaluate the Kauffman bracket invariant of this knot! So one might then think that this Kauffman bracket invariant is actually not so useful for complicated knots. We will return to this issue later in Section 2.4.

### 2.2 Relation to Physics

There is a fascinating relationship between knot invariants and quantum physics. For certain types of so-called “topological quantum systems” the amplitudes of space-time processes can be directly calculated via knot invariants such as the Kauffman bracket invariant.

We should first comment that most of what we will discuss in this book corresponds to 2 dimensional systems plus 1 dimension of time. There are topological systems in 3+1 dimension (and higher dimensions as well) but more is known about 2+1 D and we will focus on that at least for now.

Figure 2.5 shows a particular space-time process of particle world lines. At the bottom of the figure is shown the shaded 2 dimensional system (a disk). At some early time there is a pair creation event — a particle-antiparticle appear from the vacuum, then another pair creation event; then one particle walks around another, and the pairs come back together to try to reannihilate. At the end of the process, it is possible that the particles do reannihilate to the vacuum (as shown in the diagram), but it is also possible that (with some probability amplitude) the particle-antiparticle pairs form bound states that do not annihilate back to the vacuum.

In a topological theory, the quantum amplitude for these processes depends on the topology of the world lines, and not on the detailed geometry (i.e., the probability that the particles reannihilate versus form...
bound states). In other words, as long as the topology of the world lines looks like two linked rings, it will have the same quantum amplitude as that shown in Fig. 2.5. It should surprise us that systems exist where amplitudes depend only on topology, as we are used to the idea that amplitudes depend on details of things, like details of the Hamiltonian, how fast the particles move, and how close they come together. But in a topological theory, none of these things matter. What matters is the topology of the space-time paths.

What should be obvious here is that the quantum amplitude of a process is a knot invariant. It is a mapping from a knot (made by the world lines) to an output (the amplitude) which depends only on the topology of the knot. This connection between quantum systems and knot invariants was made famously by Ed Witten, one of the world’s leading string theorists [Witten, 1989]. He won the Fields medal along with Vaughan Jones for this work.

Such topological theories were first considered as an abstract possibility, mainly coming from researchers in quantum gravity (see chapter 6). However, now several systems are known in condensed matter which actually behave like this. While not all topological theories are related to the Kauffman bracket invariant, many of them are. (There are other knot invariants that occur in physical systems as well — including the so-called HOMFLY invariant [Freyd et al., 1985]. See exercise 31) A brief table of some of the physical systems that are believed to be related to nontrivial knot invariants is given in Table 2.1.

In addition there are a host of complicated systems that could in principle be engineered but are much too hard for current technology to contemplate. There are many other quantum hall states that are also topological, but have corresponding knot invariants are fairly trivial, as we will later see in chapter ***.
Kauffman Bracket Invariant and Relation to Physics

(1) $SU(2)_2$ class. For these, the Kauffman bracket invariant gives the quantum amplitude of a process by using the value $A = ie^{-i\pi/(2(2+2))} = i^{3/4}$. This is also known as “Ising” anyons. Possibly physical realizations include

- $\nu = 5/2$ Fractional Quantum Hall Effect (2D electrons at low temperature in high magnetic field). See chapters ***.
- 2D p-wave superconductors.
- 2D Films of $^3$HeA superfluid.
- A host of “engineered” structures that are designed to have these interesting topological properties. Typically these have a combination of spin-orbit coupling, superconductivity, and magnetism of some sort. Recent experiments have been quite promising. See chapter ***?

(2) $SU(2)_3$ class. For this, the Kauffman bracket invariant gives the quantum amplitude of a process by using the value $A = ie^{-i\pi/(2(2+3))} = i^{4/5}$. The only physical system known in this class is the $\nu = 12/5$ fractional quantum hall effect.

(3) $SU(2)_4$ class. For this, the Kauffman bracket invariant gives the quantum amplitude of a process by using the value $A = ie^{-i\pi/(2(2+4))} = i^{5/6}$. It is possible that $\nu = 2 + 2/3$ Fractional quantum hall effect is in this class.

(4) $SU(2)_1$ class Also known as semions. These are proposed to be realized in rotating boson fractional quantum Hall effect (See comments in chapter [50]). This corresponds to a fairly trivial knot invariant as we will see later in section ***.

(5) $SU(3)_2$ class. This corresponds to a case of the HOMFLY knot invariant rather than the Kauffman bracket invariant. It is possible that the unpolarized $\nu = 4/7$ fractional quantum hall effect is in this class.

---

Table 2.1 Table of some interesting topological systems related to knot invariants. Note that these are closely related to, but not precisely the same as $SU(2)_k$ Chern-Simons theory (which we discuss in chapter [50]). The slight differences are related to extra phases that appear in braiding. See also chapter ****. See end of chapter for references ****
2.2.1 Twist and Spin-Statistics

Before moving on, let us do some more careful examination of the Kauffman bracket invariant. To this end, let us examine a small curl in a piece of string (as shown in Fig. 2.6) and try to evaluate its Kauffman bracket invariant.

\[
\begin{align*}
A - A^{-1} = & \left( A \left[ -A^2 - A^{-2} \right] + A^{-1} \right) \\
= & -A^3
\end{align*}
\]

Fig. 2.6 Evaluation of a curl in a string. The dotted lines going off the top and bottom of the diagrams mean that the string will be connected up with itself, but we are not concerned with any part of the knot except for piece shown. The result of this calculation is that removal of the curl incurs a factor of \(-A^3\).

We see from the calculation, that the curl in the string has value of \(-A^3\) compared to a straight string. But this seems to contradict what we said earlier! We claimed earlier that any two knots that can be deformed into each other without cutting should have the same Kauffman bracket invariant, but they don’t!

The issue here is that the curled string on the right and the curled string on the left are, in fact, not topologically equivalent\(^\text{12}\). To see this we should think of the string as not being infinitely thin, but instead having some width, like a garden hose, or a “ribbon”\(^\text{13}\). If we imagine straightening a thick string (not an infinitely thin string) we realize that pulling it straight gives a twisted string (see fig. 2.7) — anyone who has tried to straighten a garden hose will realize this!\(^\text{14}\)

So the curled string is equivalent to a string with a self-twist, and this is then related to a straight string by the factor of \(-A^3\). In fact, this is a result we should expect in quantum theory. The string with a self-twist represents a particle that stays in place but rotates around an axis. In quantum theory, if a particle has a spin, it should accumulate a phase when it does a \(2\pi\) rotation, and indeed this factor of \(-A^3\) is precisely such a phase in any well defined quantum theory.

\(^{12}\)In mathematics we say they are ambient isotopic but not regular isotopic! (See section 2.6.1)

\(^{13}\)We should thus think of our knots as not just being a simple embedding of a circle \(S^1\) into a three manifold \(\mathbb{R}^3\), but rather an embedding of a ribbon. This is equivalent to specifying an orthogonal vector at each point along knot which gives the orientation of the ribbon cross section at each point. When one draws a knot as a line, one must have a convention as to what this means for the orientation of the ribbon. See comment on blackboard framing at the end of this section.

\(^{14}\)If you have not had this experience with a garden hose, you are not paying enough attention to your garden!
In the most interesting case of non-abelian statistics, there may be multiple possible exchange phases for two particles, although this does not effect the equivalence of diagrams stated here. We will discuss this more in chapter 3.

In fact, Fig. 2.7 is a very slick proof of the famous spin statistics theorem. In the left picture with the curl, we have two identical particles that change places. When we pull this straight, we have a single particle that rotates around its own axis. In quantum theory, the phases accumulated by these two processes must be identical. As we will see below in chapter 3 in 2+1 D this phase can be arbitrary (not just +1, or -1), but the exchange phase (statistical phase) and the twist phase (the spin phase) must be the same.

In the most interesting case of non-abelian statistics, there may be multiple possible exchange phases for two particles, although this does not effect the equivalence of diagrams stated here. We will discuss this more in chapter 3.

As a side comment, one can easily construct a knot invariant that treats the curled string on the left of Fig. 2.6 as being the same as the straight piece of string. One just calculates the Kauffman bracket invariant and removes a factor of $-A^3$ for each self twist that occurs.

To properly count the self twists, one calculates the so-called "writhe" of the knot (See section 2.6.2). Give the string an orientation (a direction to walk along the string) and count +1 for each positive crossing and -1 for each negative crossing where a positive crossing is when, traveling in the direction of the string that crosses over, one would have to turn left to switch to the string that crosses under. If we orient the twisted string on the left of Fig. 2.6 as up-going it then has a negative crossing by this definition.

2.2.2 Blackboard Framing

Since it is important to specify when a strand of string has a self-twist (as in the middle of Fig. 2.7) it is a useful convention to use so-called blackboard framing. With this convention we always imagine that the string really represents a ribbon and the ribbon always lies in the plane of the blackboard. An example of this is shown in Fig. 2.8. If we intend a strand to have a self twist, we draw it as a curl as in the left of Fig. 2.7 or the left of Fig. 2.6.

Fig. 2.7 Pulling straight a curl introduces a twist in the string. This twist can be replaced with a factor of $-A^3$.

Fig. 2.8 Blackboard framing. The knot drawn on the left represents the ribbon on the right, where the ribbon always lies flat in the plane of the page (i.e., the plane of the blackboard).
2.3 Bras and Kets

For many topological theories (the so-called nonabelian theories) the physical systems have an interesting, and very unusual property. Imagine we start in a ground state (or vacuum) of some systems and create two particle-hole pairs, and imagine we tell you everything that you can locally measure about these particles (their positions, if their spin, etc etc). For most gapped systems (insulators, superconductors, charge density waves) once you know all of the locally measurable quantities, you know the full wavefunction of the system. But this is not true for topological systems. As an example, see Fig. 2.9.

Particles in topological systems seem to “remember” their space-time history. The reason for this, as we will see in chapter 4 and thereafter, is that this historical information becomes encoded in the properties of the vacuum; i.e., the regions away from the particles.

\[ |0\rangle = \begin{array}{c}
\text{\begin{figure}[h]
\centering
\begin{tikzpicture}
\draw[thick] (0,0) -- (2,0);
\draw[thick] (0,1) -- (2,1);
\end{tikzpicture}
\end{figure}}
\end{array}\]
\[ |1\rangle = \begin{array}{c}
\text{\begin{figure}[h]
\centering
\begin{tikzpicture}
\draw[thick] (0,0) -- (2,0);
\draw[thick] (0,1) -- (2,1);
\end{tikzpicture}
\end{figure}}
\end{array}\]

Fig. 2.9 Two linearly independent quantum states that look identical locally but have different space-time history. The horizontal plane is a space-time slice at fixed time, and the diagrams are all oriented so time runs vertically.

\[ \langle 0 | = \begin{array}{c}
\text{\begin{figure}[h]
\centering
\begin{tikzpicture}
\draw[thick] (0,0) -- (2,0);
\draw[thick] (0,1) -- (2,1);
\end{tikzpicture}
\end{figure}}
\end{array}\]
\[ \langle 1 | = \begin{array}{c}
\text{\begin{figure}[h]
\centering
\begin{tikzpicture}
\draw[thick] (0,0) -- (2,0);
\draw[thick] (0,1) -- (2,1);
\end{tikzpicture}
\end{figure}}
\end{array}\]

Fig. 2.10 Kets are turned into bras by reversing time.

To demonstrate that these two different space-time histories are linearly independent quantum states, we simply take inner products as shown in Fig. 2.11 by gluing together a ket with a bra. Since \( \langle 0 | 0 \rangle = \langle 1 | 1 \rangle = d^2 \) but \( \langle 0 | 1 \rangle = d \), we see that \( |0\rangle \) and \( |1\rangle \) must be linearly independent, at least for \( |d| \neq 1 \). (We also see that the kets here are not properly normalized, we should multiply each bra and ket by \( 1/d \) in order that we have normalized states.)

We can think of the \( |0\rangle \) and \( |1\rangle \) states as being particular operators that produce particle-hole pairs from the vacuum, and (up to the issue of having properly normalized states) the inner product produced by graphical gluing a bra to a ket is precisely the inner product of these two resulting states. So for example, the inner product \( \langle 0 | 1 \rangle \) as shown in the bottom of Fig. 2.11 can be reinterpreted as starting from the vacuum, time evolving with the operator that gives \( |0\rangle \) then time evolving with the inverse of the operator that produces \( |1\rangle \) to return us to the vacuum.
Fig. 2.11 Showing that the kets $|0\rangle$ and $|1\rangle$ are linearly independent. For $|d| \neq 1$ the inner products show they must be linearly independent quantities.

Suppose now we insert a braid between the bra and the ket as shown in Fig. 2.12. The braid makes a unitary operation on the two dimensional vector space spanned by $|0\rangle$ and $|1\rangle$. We can once again evaluate this matrix element by calculating the Kauffman bracket invariant of the resulting knot.

Fig. 2.12 Inserting a braid between the bra and the ket. The braid performs a unitary operation on the two dimensional vector space spanned by $|0\rangle$ and $|1\rangle$. 

\[
\langle 0|0 \rangle = \begin{array}{c}
\begin{tikzpicture}
\draw[fill=red!20] (0,0) circle (1);
\draw[fill=red!20] (1,0) circle (1);
\draw[blue,thick] (0,0) to[out=90,in=90] (1,1);
\draw[blue,thick] (0,1) to[out=-90,in=-90] (1,0);
\end{tikzpicture}
\end{array}
= \begin{array}{c}
\begin{tikzpicture}
\draw[fill=red!20] (0,0) circle (0.5);
\end{tikzpicture}
\end{array}
= d^2
\]

\[
\langle 1|1 \rangle = \begin{array}{c}
\begin{tikzpicture}
\draw[fill=red!20] (0,0) circle (1);
\draw[fill=red!20] (1,0) circle (1);
\draw[blue,thick] (0,0) to[out=90,in=90] (1,1);
\draw[blue,thick] (0,1) to[out=-90,in=-90] (1,0);
\end{tikzpicture}
\end{array}
= \begin{array}{c}
\begin{tikzpicture}
\draw[fill=red!20] (0,0) circle (0.5);
\end{tikzpicture}
\end{array}
= d^2
\]

\[
\langle 0|1 \rangle = \begin{array}{c}
\begin{tikzpicture}
\draw[fill=red!20] (0,0) circle (1);
\draw[fill=red!20] (1,0) circle (1);
\draw[blue,thick] (0,0) to[out=90,in=90] (1,1);
\draw[blue,thick] (0,1) to[out=-90,in=-90] (1,0);
\end{tikzpicture}
\end{array}
= \begin{array}{c}
\begin{tikzpicture}
\draw[fill=red!20] (0,0) circle (0.5);
\end{tikzpicture}
\end{array}
= d
\]
2.4 Quantum Computation with Knots

Why do we care so much about topological systems and knot invariants? A hint is from the fact that we wrote states above as $|0\rangle$ and $|1\rangle$. This notation suggests the idea of qubits, and indeed this is one very good reason to be interested.

It turns out that many topological quantum systems can compute quantities efficiently that classical computers cannot. To prove this, suppose you wanted to calculate the Kauffman invariant of a very complicated knot, say with 100 crossings. As mentioned above, a classical computer would have to evaluate $2^{100}$ diagrams, which is so enormous, that it could never be done. However, suppose you have a topological system of Kauffman type in your laboratory. You could actually arrange to physically measure the Kauffman bracket invariant. The way we do this is to start with a system in the vacuum state, arrange to “pull” particle-hole (particle-antiparticle) pairs out of the vacuum, then drag the particles around in order to form the desired knot, and bring them back together to reannihilate. Some of the particles will reannihilate, and others will refuse to go back to the vacuum (forming bound states instead). The probability that they all reannihilate is (up to a normalization) given by the absolute square of the Kauffman bracket invariant of the knot (since amplitudes are the Kauffman bracket invariant, the square of the Kauffman bracket invariant is the probability). Even estimation of the Kauffman bracket invariant of a large knot is essentially impossible for a classical computer, for almost all values of $A$. However, this is an easy task if you happen to have a topological quantum system in your lab! Thus the topological quantum system has computational ability beyond that of a classical computer.

It turns out that the ability to calculate Kauffman bracket invariant is sufficient to be able to do any quantum computation. One can use this so-called topological quantum computer to run algorithms such as Shor’s famous factoring (i.e., code breaking) algorithm. The idea of using topological systems for quantum computation is due to Michael Freedman and Alexei Kitaev.

So it turns out that these topological systems can do quantum computation. Why is this a good way to do quantum computation? First we must ask about why quantum computing is hard in the first place. In the conventional picture of a quantum computer, we imagine a bunch of two state systems, say spins, which act as our qubits. Now during our computation, if some noise, say a photon, or a phonon, enters the system, if some noise, say a photon, or a phonon, enters the system, it is virtually impossible for a classical computer. Say for 100 spins you would have to find the eigenvalues and eigenvectors of a $2^{100}$ dimensional matrix. But if you had the physical system in your lab, you could just measure its dynamics and answer certain questions. So in that sense the physical quantum system is able to compute certain quantities, i.e., its own equations of motion, that a classical computer cannot. In the following year Feynman started thinking along the same lines and asked the question of whether one quantum system can compute the dynamics of another quantum system — which starts getting close to the ideas of modern quantum computation.
system and interacts with a qubit, it can cause an error or decoherence, which can then ruin your computation. And while it is possible to protect quantum systems from errors (we will see in section *** below how you do this) it is very hard.

Now consider what happens when noise hits a topological quantum computer. In this case, the noise may shake around a particle, as shown in Fig. 2.13. However, as long as the noise does not change the topology of the knot, then no error is introduced. Thus the topological quantum computer is inherently protected from errors. (of course sufficiently strong noise can change the topology of the knot and still cause errors.)

\[ \text{noise} \Rightarrow \]

**Fig. 2.13** The effect of noise on a topological quantum computation. As long as the noise does not change the topology of the knot, then no error is introduced.

### 2.5 Some Quick Comments about Fractional Quantum Hall Effect

There will be chapters later about Fractional Quantum Hall Effect (FQHE). But it is worth saying a few words about FQHE as a topological system now.

FQHE occurs in two dimensional electronic systems\(^{25}\) in high magnetic field at low temperature (typically below 1K). There are many FQHE states which are labeled by their so called filling fraction \( \nu = p/q \) with \( p \) and \( q \) small integers. The filling fraction can be changed in experiment by, for example, varying the applied magnetic field (we will discuss this later in chapter ??). The FQHE state emerges at low temperature and is topological\(^{26}\).

How do we know that the system is topological? There are not a whole lot of experiments that are easy to do on quantum Hall systems, since they are very low temperature and complicated experiments to do. However, one type of experiment is fairly straightforward — a simple electrical resistance measurement, as shown in Figs. 2.14 and 2.15. In Fig. 2.14 the so-called longitudinal resistance is measured — where the current runs roughly parallel to the voltage. In this case the measured voltage is zero — like a superconductor. This shows that this state of matter has no dissipation, no friction.

The measurement in the Fig. 2.15 is more interesting. In this case, the Hall voltage is precisely quantized as \( V = (\hbar/e^2)(1/\nu)I \) where \( I \) is

---

25 Electronic systems can be made two dimensional in several ways. See comments in chapter ??.

26 A comment in comparing this paradigm to the common paradigm of high energy physics: In high energy there is generally the idea that there is some grand unified theory (GUT) at very high energy scale and it is extremely symmetric, but then when the universe cools to low temperature, symmetry breaks (such as electro-weak symmetry) and we obtain the physics of the world around us. The paradigm is opposite here. The electrons in magnetic field at high temperature have no special symmetry. However, as we cool down to lower temperature, a huge symmetry emerges. The topological theory is symmetric under all diffeomorphisms (smooth distortions) of space and time.
2.5 Some Quick Comments about Fractional Quantum Hall Effect

2D electron sample in $B$-field

$V = 0$ Dissipationless Flow

Fig. 2.14 Measurement of longitudinal resistance in FQHE experiment.

$V = (h/e^2)(1/\nu)I$ Quantized Resistance

Fig. 2.15 Measurement of Hall resistance in FQHE experiment.

the current, $h$ is Plank’s constant, $e$ the electron charge and $\nu = p/q$ is a ratio of small integers. This quantization of $V/I$ is extremely precise — to within about a part in $10^{10}$. This is like measuring the distance from London to Los Angeles to within a millimeter. What is most surprising is that the measured voltage does not depend on details, such as the shape of the sample, whether there is disorder in the sample, or where you put the voltage leads or how you attach them as long as the current and voltage leads are topologically crossed, as they are in the Fig. 2.15 but not in Fig. 2.14. We should emphasize that this is extremely unusual. If you were to measure the resistance of a bar of copper, the voltage would depend entirely on how far apart you put the leads and the shape of the sample. This extremely unusual independence of all details is a strong hint that we have something robust and topological happening here.

Finally we can ask about what the particles are that we want to braid around each other in the FQHE case. These so-called quasiparticles are like the point-vortices of the FQHE superfluid. As we might expect for a dissipationless fluid, the vortices are persistent — they will last forever unless annihilated by antivortices.

So in fact, Kelvin was almost right (See chapter 1). He was thinking about vortices knotting in the dissipationless aether. Here we are thinking about point vortices in the dissipationless FQHE fluid, but we move the vortices around in time to form space-time knots!
2.6 Appendix: More Knot Theory Basics

2.6.1 Isotopy and Reidermeister Moves

Two knots (or two pictures of knots) are ambient isotopic if one can be deformed into each other without cutting any of the strands. In order for two pictures of knots to be ambient isotopic they must be related to each other by a series of moves, known as Reidermeister moves\(^ {27}\), as shown in Fig. 2.16.

\(^{27}\)This is a very old result, by Kurt Reidemeister from 1927. Note that it may take many many moves in order to bring a knot into some particular desired form. For example, if there are \(c\) crossings in a diagram which is equivalent to the simple unknot (an unknotted loop), the strongest theorem yet proven is that it can be reduced to the simple unknot with \((236c)^{11}\) moves\(^ {27}\), as shown in Fig. 2.16.

In the context of quantum physics, and as elaborated in section 2.2.1, we are usually concerned with regular isotopy which treats the strands as ribbons. Two knots are regular isotopic if they can related to each other using only type-II and type-III moves. A type-I move inserts a twist in the ribbon (See Fig. 2.7) and results in a different ribbon diagram, whereas type-II and III moves do not twist the ribbon\(^ {28}\).

2.6.2 Writhe and Linking

Let us put arrows on all strands of our knots and links (so now we have directed lines). For each crossing we define a sign \(\epsilon\) as shown in Fig. 2.17.

\[\epsilon = \begin{cases} +1 & \text{if over-under}\ \text{crossing} \\ -1 & \text{if under-over}\ \text{crossing} \end{cases}\]

\[\epsilon = \begin{cases} +1 & \text{if over-under}\ \text{crossing} \\ -1 & \text{if under-over}\ \text{crossing} \end{cases}\]

The writhe \(w\) of an oriented knot (here “knot” means made of a single strand) is the sum of all of the \(\epsilon\) values of the crossings

\[w(knot) = \sum_{\text{crossings}} \epsilon(\text{crossing}) \quad (2.2)\]

Note that type II and III Reidermeister moves preserve the writhe of a knot, whereas type I moves do not. Thus, the writhe is an invariant of regular isotopy but not of ambient isotopy.

For a link made of two strands, the linking number \(lk\) between the two strands is given by

\[lk(link) = \sum_{\text{crossings between two different strands}} \epsilon(\text{crossing}) \quad (2.3)\]
Chapter Summary

- Knot invariants, such as the Kauffman bracket invariant, help distinguish knots from each other.
- The quantum dynamics of certain particles are determined by certain knot invariants.
- Computation of certain knot invariants is computationally “hard” on a classical computer, but not hard using particles whose dynamics is given by knot invariants.
- Computation by braiding these particles is equivalent to any other quantum computer.
- Physical systems which have these particles include fractional quantum Hall effect.

Further Reading

- The book by Kauffman [2001] is a delightful introduction to knot theory and connections to physics. This was the book that got me interested in the subject back when I was in grad school and changed the course of my life.
- I wrote another easy reading introduction, Simon [2010], connecting knots to anyons.
- Some nice introductory books on knots include Adams [1994], and Sossinsky [2002]. A beautiful set of course notes on knot theory is given by Roberts [2015].

Exercises

Exercise 2.1 Trefoil Knot and the Kauffman Bracket
Using the Kauffman rules, calculate the Kauffman bracket invariant of the right and left handed trefoil knots shown in Fig. 2.18. Conclude these two knots are topologically inequivalent. While this statement appears obvious on sight, it was not proved mathematically until 1914 (by Max Dehn). It is trivial using this technique!

Exercise 2.2 Abelian Kauffman Anyons
Anyons described by the Kauffman bracket invariant with certain special values of the constant $A$ are abelian anyons – meaning that an exchange introduces only a simple phase as shown in Fig. 2.19.

29 The word “trefoil” is from the plant trifolium, or clover, which has compound trifoliate leaves.
Exercises

(a) For \( A = \pm e^{i\pi/3} \) (and the complex conjugates of these values), show that the anyons are bosons or fermions respectively (i.e., \( e^{i\theta} = \pm 1 \)).

(b) For \( A = \pm e^{i\pi/6} \) (and the complex conjugates of these values) show the anyons are semions (i.e., \( e^{i\theta} = \pm i \)). In fact these are precisely the anyons that arise for the \( \nu = 1/2 \) fractional quantum Hall effect of bosons (We will discuss this later in this book (See chapter 57). This particular phase of quantum Hall matter has been produced experimentally (Clark et al. 2020), but only in very small puddles so far and it has not been possible to measure braiding statistics as of yet.

HINT: For (a) and (b) show first the identity shown in Fig. 2.20.

If you can’t figure it out, try evaluating the Kauffman bracket invariant for a few knots with these values of \( A \) and see how the result arises.

Exercise 2.3 Reidermeister moves and the Kauffman Bracket

Show that the Kauffman bracket invariant is unchanged under application of Reidermeister move of type II and type III. Thus conclude that the Kauffman invariant is an invariant of regular isotopy.

Exercise 2.4 Jones polynomial

Let us define the Jones polynomial of an oriented knot as

\[
\text{Jones}(knot) = (-A^{-3w(knot)}) \text{ Kauffman}(knot)
\]

where \( w \) is the writhe (We must first orient the knot, meaning we arrows on the strands, in order to define a writhe). Show that this quantity is an invariant of ambient isotopy – that is, it is invariant under all three Reidermeister moves.

Exercise 2.5 HOMFLY Polynomial

The HOMFLY polynomial is a generalization of the Jones polynomial which has two variables \( A \) and \( z \) rather than just one variable. To define the HOMFLY polynomial we must first orient the strings in our knot or link (meaning we put arrows on the lines). The HOMFLY polynomial (Freyd et al. 1985; Przytycki and Traczyk 1987) of an oriented link is then defined in terms of two variables \( A \) and \( z \) by the two rules

\[
\text{Fig. 2.19} \quad \begin{array}{c}
\text{For abelian anyons, exchange gives a phase } e^{i\theta}.
\end{array}
\]

\[
\begin{array}{c}
\text{Fig. 2.20} \quad \begin{array}{c}
\text{For bosons or fermions the sign in this figure is +, for semions the sign is −.}
\end{array}
\end{array}
\]

(a) Show that the HOMFLY polynomial is invariant under type I Reidermeister moves\(^{31}\)

(b) Calculate the HOMFLY polynomial of the right and left handed trefoil knots (shown in Fig. 2.18).

\(^{30}\)HOMFLY is an acronym of the names of the inventors of this polynomial. Sometimes credit is even more distributed and it is called HOMFLYPT.

\(^{31}\)In order for the HOMFLY invariant to represent quantum particles, a phase for this type I move must be inserted by hand.
Part I

Anyons and Topological Quantum Field Theories
In chapter 2 we discussed braiding particles around each other, or exchanging their positions. This is what we call particle statistics (or quantum statistics, or exchange statistics). What we mean by this is “what happens to the many particle wavefunction when particles are exchanged in a certain way.”

We are familiar with bosons and fermions. If we exchange two bosons the wavefunction is unchanged, if we exchange two fermions the wavefunction accumulates a minus sign. Various arguments have been given as to why these are the only possibilities. The argument usually given in introductory books is as follows:

If you exchange a pair of particles then exchange them again, you get back where you started. So the square of the exchange operator should be the identity, or one. There are two square roots of one: +1 and −1, so these are the only two possibilities for the exchange operator.

In the modern era this argument is considered to be incorrect (or at least not really sufficient). To really understand the possibilities in exchange statistics, it is very useful to think about quantum physics from the Feynman path integral point of view.

### 3.1 Single Particle Path Integral

Consider a space-time trajectory of a single non-relativistic particle. We say that we have $x$ moving in $\mathbb{R}^D$ where $D$ is the dimension of space, so we can write $x(t)$ where $t$ is time.

Given that we start at position $x_i$ at the initial time $t_i$ we can define a so-called propagator which gives the amplitude of ending up at position $x_f$ at the final time $t_f$. This can be written as

$$\langle x_f | \hat{U}(t_f, t_i) | x_i \rangle$$

where $\hat{U}$ is the (unitary) time evolution operator.

The propagator can be used to propagate forward in time some arbitrary wavefunction $\psi(x) = \langle x | \psi \rangle$ from $t_i$ to $t_f$ as follows

$$\langle x_f | \psi(t_f) \rangle = \int dx_i \langle x_f | \hat{U}(t_f, t_i) | x_i \rangle \langle x_i | \psi(t_i) \rangle$$

If we are trying to figure out the propagator from some microscopic

---

1 Bose cooked up the current picture of Bose statistics in 1924 in the context of photons and communicated it to Einstein who helped him get it published. Einstein realized the same ideas could be applied to non-photon particles as well.

2 Based on ideas by Pauli, Fermi-Dirac statistics were actually invented by Jordan in 1925. Jordan submitted a paper to a journal, where Max Born was the referee. Born stuck the manuscript in his suitcase and forgot about it for over a year. During that time both Fermi and Dirac published their results. Jordan could have won a Nobel Prize (potentially with Born) for his contributions to quantum physics, but he became a serious Nazi and no one really liked him much after that. Born felt terribly guilty about his mistake later in life, stating “I hate Jordan’s politics, but I can never undo what I did to him.”

3 The error in this argument is that one has to be much more careful about defining what one means about an “exchange”.

4 If you are familiar with path integrals you can certainly skip down to section 3.2. If you are not familiar with path integrals, please do not expect this to be a thorough introduction! What is given here is a minimal introduction to give us what we need to know for our purposes and nothing more! See the Further Reading for this chapter for a better introduction.
calculation, there are two very fundamental properties it must obey. First, it must be unitary — meaning no amplitude is lost along the way (normalized wavefunctions stay normalized). Secondly it must obey composition: propagating from \( t_i \) to \( t_m \) and then from \( t_m \) to \( t_f \) must be the same as propagating from \( t_i \) to \( t_f \). We can express the composition law as

\[
\langle x_f | \hat{U}(t_f, t_i) | x_i \rangle = \int dx_m \langle x_f | \hat{U}(t_f, t_m) | x_m \rangle \langle x_m | \hat{U}(t_m, t_i) | x_i \rangle
\]

The integration over \( x_m \) allows the particle to be at any position at the intermediate time (and it must be at some position). Another way of seeing this statement is to realize that the integral over \( x_m \) is just insertion of a complete set of states at some intermediate time

\[
1 = \int dx_m | x_m \rangle \langle x_m |.
\]

Feynman’s genius was to realize that you can subdivide time into infinitesimally small pieces, and you end up doing lots of integrals over all possible intermediate positions. In order to get the final result, you must sum over all values of all possible intermediate positions, or all possible functions \( x(t) \). Feynman’s final result is that the propagator can be written as

\[
\langle x_f | \hat{U}(t_f, t_i) | x_i \rangle = \mathcal{N} \sum_{\text{paths } x(t) \text{ from } (x_i, t_i) \text{ to } (x_f, t_f)} e^{iS[x(t)]/\hbar} \tag{3.1}
\]

where \( \mathcal{N} \) is some normalization constant. Here \( S[x(t)] \) is the (classical!) action of the path

\[
S = \int_{t_i}^{t_f} dt L[x(t), \dot{x}(t), t]
\]

with \( L \) the Lagrangian.

The sum over paths in Eq. (3.1) is often well defined as a limit of dividing the path into discrete time steps and integrating over \( x \) at each time. We often rewrite this sum over paths figuratively as a so-called path integral

\[
\langle x_f | \hat{U}(t_f, t_i) | x_i \rangle = \mathcal{N} \int_{(x_i, t_i)}^{(x_f, t_f)} \mathcal{D}x(t) \ e^{iS[x(t)]/\hbar} \tag{3.2}
\]

Analogous to when we evaluate regular integrals of things that look like \( \int dx \ e^{iS[x]/\hbar} \), we can approximate the value of this integral in the small \( \hbar \), or classical, limit by saddle point approximation. We do this by looking for a minimum of \( S \) with respect to its argument — this is where the exponent oscillates least, and it becomes the term which dominates the result of the integral. Similarly, with the path integral, the piece that dominates in the small \( \hbar \) limit is the piece where \( S[x(t)] \) is extremized — the function \( x(t) \) which extremizes the action. This is just the classical principle of least action!
3.2 Two Identical Particles

We would now like to generalize the idea of a path integral to systems with multiple identical particles, starting with the case of two particles. If the particles are identical there is no meaning to saying that particle one is at position $x_1$ and particle two is at position $x_2$. This would be the same as saying that they are the other way around. Instead, we can only say that there are particles at both positions $x_1$ and $x_2$. To avoid the appearance of two different states expressed as $|x_1, x_2\rangle$ versus $|x_2, x_1\rangle$ (which are actually the same physical state\(^5\)), it is then useful to simply agree on some convention for which coordinate we will always write first — for example, maybe we always write the leftmost particle first\(^6\). For simplicity, we can assume that $x_1 \neq x_2$, i.e., the particles have hard cores and cannot overlap\(^7\). For these indistinguishable particles, the Hilbert space is then cut in half compared to the case of two distinguishable particles where $|x_1, x_2\rangle$ and $|x_2, x_1\rangle$ mean physically different things.

We call the space of all states the configuration space $\mathcal{C}$. To construct a path integral, we want to think about all possible paths through this configuration space. The key realization is that the space of all paths through the configuration space $\mathcal{C}$ divides up into topologically inequivalent pieces. I.e., certain paths cannot be deformed into other paths by a series of small deformations.

What do these topologically disconnected pieces of our space of paths look like? For example, we might consider the two paths as shown in Fig. 3.1. Here we mean that time runs vertically. It is not possible to continuously deform the path on the left into the path on the right assuming the end points are fixed.

We will call the non-exchange path TYPE +1 (left in Fig. 3.1), and the exchange path TYPE −1 (right in Fig. 3.1). The two sets of paths cannot be continuously deformed into each other assuming the end points are fixed. Note that we may be able to further refine our classification of paths — for example, we may distinguish over- and under-crossings, but for now we will only be concerned with exchanges (TYPE −1) and non-exchanges (TYPE +1).

Paths can be composed with each other. In other words, we can follow one path first, then follow the second. We can write a multiplication table for such composition of paths (the path types form a group, see Section 3.1.2).

\[
\begin{array}{ccc}
\text{TYPE +1} & \text{Followed by} & \text{TYPE +1} & = & \text{TYPE +1} \\
\text{TYPE +1} & \text{Followed by} & \text{TYPE −1} & = & \text{TYPE −1} \\
\text{TYPE −1} & \text{Followed by} & \text{TYPE +1} & = & \text{TYPE −1} \\
\text{TYPE −1} & \text{Followed by} & \text{TYPE −1} & = & \text{TYPE +1} \\
\end{array}
\] (3.3)

So for example, an exchange path (which switches the two particles) followed by another exchange path (which switches again) results in a

\(^5\) Often books define $|x_1, x_2\rangle = -|x_2, x_1\rangle$ for fermions. The two sets describe the same state in the Hilbert space only with a different phase prefactor. We should contrast this to the case of distinguishable particles where $|x_1, x_2\rangle$ and $|x_2, x_1\rangle$ have no overlap for $x_1 \neq x_2$.

\(^6\) This ordering scheme works in one dimension. In two dimensions we would perhaps say, the particle with the smaller $x$ coordinate is written first, but in case of two particles with the same value of $x$, the particle with smaller $y$ coordinate is written first.

\(^7\) It is sometimes even more convenient to declare $|x_1 − x_2| > \epsilon$. 

Fig. 3.1 Two possible sets of paths (paths in configuration space) from the same two starting positions to the same two ending positions (we are implying that time runs vertically). We call the non-exchange path TYPE +1, and the exchange path TYPE −1. Here we mean that time runs vertically. The two sets of paths cannot be continuously deformed into each other assuming the end points are fixed. Note that we may be able to further refine our classification of paths — for example, we may distinguish over and under-crossings, but for now we will only be concerned with exchanges (TYPE −1) and non-exchanges (TYPE +1).
If \( |x_1, x_2\rangle \neq |x_1, x_2\rangle \), i.e., if the initial and final endpoints of the path are not the same, then we need a more general definition of what we call TYPE \( +1 \) versus TYPE \( -1 \). One simple possible definition is to count the number of times the space-time paths cross given a particular fixed viewing angle. For example, we can use the ordering rule of note 6 above and as time evolves, we can count the number of times the ordering changes (which corresponds to a crossing of the world lines as in Fig. 3.1). An even number of such crossings would correspond to a TYPE \( +1 \) path, and an odd number of crossings would correspond to a TYPE \( -1 \) path. Other consistent definitions are also possible as long as the multiplication rule of Eq. 3.3 is maintained.

\[ N \left( \sum_{\text{TYPE } +1 \text{ paths}} e^{iS[\text{path}]/\hbar} + \sum_{\text{TYPE } -1 \text{ paths}} e^{iS[\text{path}]/\hbar} \right) \]

This second line is simply a rewriting of the first having broken the sum into the two different classes of paths.

It turns out however, that it is completely consistent to try something different. Let us instead write

\[ \langle x_1 x_2 | \hat{U}(t_f, t_i) | x_{1f} x_{2f} \rangle = \]

\[ N \left( \sum_{\text{TYPE } +1 \text{ paths}} e^{iS[\text{path}]/\hbar} - \sum_{\text{TYPE } -1 \text{ paths}} e^{iS[\text{path}]/\hbar} \right) \]

Notice the change of sign for the TYPE \( -1 \) paths.

The reason this change is allowed is because it obeys the composition law. To see this, let us check to see if the composition law is still obeyed. Again, we break the time propagation at some intermediate time

\[ \langle x_1 x_2 | \hat{U}(t_f, t_i) | x_{1f} x_{2f} \rangle = \]

\[ \int dx_{1m} dx_{2m} \langle x_{1f} x_{2f} | \hat{U}(t_f, t_m) | x_{1m} x_{2m} \rangle \langle x_{1m} x_{2m} | \hat{U}(t_m, t_i) | x_{1f} x_{2f} \rangle e^{iS[\text{path}]/\hbar} \]

where in the last line we have substituted in Eq. 3.4 for each of the two propagators on the right, and we have used a bit of shorthand in writing the result.

Now, when we compose together subpaths from \( i \to m \) with those from \( m \to f \) to get the overall path, the sub-path types multiply according to our above multiplication table Eq. 3.3. For the full path, there are two ways to obtain a TYPE \( +1 \) path: (1) both sub-paths are TYPE \( +1 \) or (2) both sub-paths are TYPE \( -1 \). In either case, note that the net prefactor of the overall TYPE \( +1 \) path is \( +1 \). (In the case where both
3.3 Many Identical Particles

Generalizing this idea, to figure out what is consistent in quantum mechanics, we must do two things:

(a) Characterize the space of paths through configuration space

(b) Insist on consistency under composition.

Let us first discuss our configuration space. If we had \( N \) distinguishable particles in \( D \) dimensions we would have a configuration space \((\mathbb{R}^D)^N\) representing the coordinates \(\{x_1, x_2, x_3, \ldots, x_N\}\). For simplicity we usually assume all of these coordinates are different (We might imagine that the particles are hard spheres of some very small diameter \(\epsilon\)). Thus we write the configuration space as \([(\mathbb{R}^D)^N - \Delta]\) where \(\Delta\) represents the so-called coincidences where two particles are at the same position\(^{10}\).

In the case of identical particles we want to disregard the order in which we write the coordinates. In other words, we have an equivalence relationship \(\sim\) between the \(N!\) possible orderings of the coordinates

\[
\{x_1, x_2, x_3, \ldots, x_N\} \sim \{x_2, x_3, x_7, \ldots, x_9\} \sim \{x_3, x_N, x_2, \ldots, x_1\} \sim \ldots
\]

Thus for indistinguishable particles the configuration space is

\[
\mathcal{C} = [(\mathbb{R}^D)^N - \Delta] / \sim
\]

where “/ \(\sim\)” means that we are “modding out” by the equivalence relationship \(\sim\). This is just a fancy way to say that the order in which we list the coordinates \(\{x_1, x_2, x_3, \ldots, x_N\}\) does not matter (or as described in section 3.2 we choose some convention for the order, like always writing the left-most first). In the case of 2 identical particles above, this reduced the Hilbert space by a factor of 2. With \(N\) identical particles this will reduce the Hilbert space by a factor of \(N!\). This is the same indistinguishability factor which is familiar from the Gibbs paradox of statistical mechanics.

We would now like to consider all possible paths through this configuration space \(\mathcal{C}\). In other words we want to consider how these \(N\) different points move in time. We can think of this as a set of coordinates moving through time \(\{x_1(t), \ldots, x_N(t)\}\) but we must be careful

---

\(^{10}\)Mathematicians often write \((\mathbb{R}^D)^N \setminus \Delta\) to represent removing \(\Delta\) from the set \((\mathbb{R}^D)^N\).
that the particles are indistinguishable, so the order in which we write the coordinates doesn’t matter. We can think of this as \( N \) directed curves moving in \( ND + 1 \) dimensional space\(^{11}\). Since we want to add up all of these possible paths in a path integral it is useful to try to understand the structure of this space of paths better.

Again, the key realization is that the space of all paths through the configuration space \( \mathcal{C} \) divides up into topologically inequivalent pieces. I.e., certain paths cannot be deformed into other paths by a series of small deformations assuming the endpoints are fixed. The group of paths through \( \mathcal{C} \) is familiar to mathematicians and is known as the first homotopy group \( \Pi_1(\mathcal{C}) \) or fundamental group\(^{12}\). The reason this is a group is that it comes with a natural operation, or multiplication of elements — which is the composition of paths: follow one path, then follow another path.

### 3.3.1 Paths in 2+1 D, the Braid Group

A path through the configuration space of particles in 2 dimensions is known as a braid. An example of a braid is shown in Fig. [3.3].

A few notes about braids:

1. Fixing the endpoints, the braids can be deformed continuously, and so long as we do not cut one string through another, it still represents the same topological class, or the same element of the braid group.
2. We cannot allow the strings to double-back in time as in Fig. [3.2]. This would be pair creation or annihilation, which we will consider later, but not now.

The set of braids have mathematical group structure (See section \[11.2\]: multiplication of two braids is defined by stacking the two braids on top of each other – first do one then do another. It is easy to see that braids can be decomposed into elementary pieces which involve either clockwise or counterclockwise exchange of one strand with its neighbor. These elementary pieces involving single exchanges are known as generators.

The braid group on \( N \) strands is typically notated as \( B_N \). The generators of the braid group on 4 strands are shown in Fig. [3.4]. Any braid can be written as a product of the braid generators and their inverses\(^{13}\).

The “multiplication” of the generators is achieved simply by stacking the generators on top of each other. An expression representing a braid, such as \( \sigma_1 \sigma_2 \sigma_3^{-1} \sigma_1 \) is known as a “braid word.” Typically we read the braid word from right to left (do the operation listed right-most first), although sometimes people use the opposite convention! The important thing is to fix a convention and stick with it!

Note that many different braid words can represent the same braid. An example of this is shown for \( B_4 \) in Fig. [3.5]. Although a braid can be written in many different ways\(^{14}\), it is possible to define invariants of the braid which do not change under deformation of the braid — so long
3.3 Many Identical Particles

\[ \sigma_1 = \quad \sigma_2 = \quad \sigma_3 = \]

\[ \sigma_1^{-1} = \quad \sigma_2^{-1} = \quad \sigma_3^{-1} = \]

**Fig. 3.4** The three generating elements \( \sigma_1, \sigma_2, \sigma_3 \) of the braid group on 4 strands, \( B_4 \), and their inverses \( \sigma_1^{-1}, \sigma_2^{-1}, \sigma_3^{-1} \). Any braid on four strands (any element of \( B_4 \)) can be written as a product of the braid generators and their inverses by simply stacking these generators together (See Fig. 3.5 for examples).

\[
\sigma_1^{-1} \sigma_2^{-1} \sigma_1 = \begin{array}{c}
\text{third} \\
\text{second} \\
\text{first}
\end{array} = \begin{array}{c}
\text{third} \\
\text{second} \\
\text{first}
\end{array} = \sigma_2 \sigma_1^{-1} \sigma_2^{-1}
\]

**Fig. 3.5** Two braid words in \( B_4 \) that represent the same braid. The figure on the left can be continuously deformed to the one on the right, keeping endpoints fixed. The braid words are read from right to left indicating stacking the generators from bottom to top. \(^{14}\)

as the braid is topologically unchanged. One very useful braid invariant is given by the so-called winding number

\[
W = \text{Winding Number} = (\text{# of overcrossings}) - (\text{# of undercrossings})
\]

where an overcrossing is a \( \sigma \) and an undercrossing is a \( \sigma^{-1} \). As can be checked in Fig. 3.5, the winding number is independent of the particular way we represent the braid. As long as we do not cut one strand through another or move the endpoints (or double-back strands) the winding number, a braid invariant, remains the same.

14All braid word equivalences can be derived from the identity

\[ \sigma_n \sigma_{n+1} \sigma_n = \sigma_{n+1} \sigma_n \sigma_{n+1} \]

For example, try deriving Fig. 3.5 from this. See also exercise 3.1.

The observant reader will see the similarity here to Reidermeister moves of type-III discussed in section 2.6.1. Similarly \( \sigma_i \sigma_i^{-1} = 1 \) is a type-II move.

3.3.2 Paths in 3+1 D, the Permutation Group

We now turn to consider physics in 3+1 dimensions. A key fact is that it is not possible to knot a one-dimensional world-line that lives in a four-dimensional space. If this is not obvious consider the following lower dimensional analogue shown in Fig. 3.6. In one dimension, two points cannot cross through each other without hitting each other. But if we allow the points to move in 2D they can move around each other without touching each other. Analogously we can consider strings forming knots or braids in 3D space. When we try to push these strings through each other it would be very convenient to be able to draw a diagram in four dimensions! \(^{16}\)

\[ \text{In one dimension:} \]

Two objects cannot cross

\[ \text{In two dimensions:} \]

Two objects can go around each other

**Fig. 3.6** Top: In one dimension, two points cannot cross through each other without hitting each other. Bottom: However, if we allow the points to move in two dimensions they can get around each other without touching. This is supposed to show you that one-dimensional world-lines cannot form knots in four-dimensional space.
other, they bump into each other and get entangled. However, if we allow the strings to move into the fourth dimension, we can move one string a bit off into the fourth dimension so that it can move past the other string, and we discover that the strings can get by each other without ever touching each other! Hence there are no knots of one dimensional objects embedded in four dimensions.

Given that in 3+1 D world-lines cannot form knots, the only thing that is important in determining the topological classes of paths is where the strings start and where they end. In other words, we can draw things that look a bit like braid-diagrams but now there is no meaning to an over or under-crossing. If the world line lives in 3+1 dimensions, everything can be unentangled without cutting any of the world lines until the diagram looks like Fig. 3.7 indicating only where lines start and end. This is precisely describing the permutation group, or symmetric group \( S_N \) (see section 31.2.1).

### 3.3.3 Building a Path Integral

We now return to the issue of building a path integral. We will follow the intuition we gained in the two particle case, but now we will include the information we have discovered about the group of paths through configuration space.

Using the notation \( \{x\} \) to denote all of the \( N \) particle coordinates, we construct the path integral as

\[
\langle \{x\}_f | \hat{U}(f,t_i) | \{x\}_i \rangle = \mathcal{N} \sum_{g \in G} \rho(g) \sum_{\text{paths } i \rightarrow f \in g} e^{iS[\text{path}]/\hbar} \tag{3.5}
\]

Here \( G \) is the group of paths (the fundamental group — or the set of classes of topologically different paths). This is the symmetric group \( S_N \) for 3+1 dimensions and is the braid group \( B_N \) for 2+1 dimensions. Here we have split the sum over paths into the different classes — the outer sum being a sum over the classes \( g \) and the inner sum being the sum over all paths of type \( g \), i.e., a set of paths that can be continuously deformed into each other. We have also introduced a factor of \( \rho(g) \) out front where \( \rho \) is a unitary representation of the group \( G \). (See section 41.2.4 on group theory).

In the case where the initial set of position \( |\{x\}_i\rangle \) and the final set of position \( |\{x\}_f\rangle \) are not the same (similar to the case mentioned in note 8 above), the definition 3.5 can still be used, although strictly speaking these are not precisely what we would call braids or permutations (for which initial and final positions are supposed to match). Nonetheless we can associate an element \( g \) of the braid or permutation group to each space-time path by viewing the motion from some fixed angle and

---

**Fig. 3.7** Paths in 3+1 D are elements of the permutation group (or symmetric group) \( S_N \) (See section 31.2.1). Shown here is an element of \( S_4 \).

---

17 One way to think about the relationship between the symmetric group and the braid group is to say that the symmetric group \( S_N \) is a “truncation” of the braid group \( B_N \), meaning that it obeys the same group properties, except that in \( S_N \), the element \( \sigma_i^2 \) has been identified with the identity.

18 In the nonabelian case discussed in section 3.5 below the ket \( |\{x\}\rangle \) is given an additional index to become \( |n,\{x\}\rangle \) with \( n = 1 \ldots M \). This then implies a basis choice for the \( M \)-dimensional space, and this basis choice for one set of positions \( \{x\} \) can be chosen independently of the basis choice for a different set of positions. When the initial and final positions are not the same we can make two independent basis choices and changing these choices simply pre- or post- multiplies the representation \( \rho \) by the appropriate basis changing unitaries. This caution is related to notes 8 and 12 above.
smoothly deforming the paths such that start and endpoints are at some reference positions without introducing any new crossings to the paths\footnote{A crossing can be defined as a reordering of coordinates as in note \ref{note:crossing} above.}.

To show that Eq. 3.5 is allowed by the laws of quantum mechanics, we need only check that it obeys the composition law – we should be able to construct all paths from $i$ to $f$ in terms of all paths from $i$ to $m$ and all paths from $m$ to $f$.

\[
\langle \{x\}_f | \hat{U}(t_f, t_i) | \{x\}_i \rangle = \int d\{x\}_m \langle \{x\}_f | \hat{U}(t_f, t_m) | \{x\}_m \rangle \langle \{x\}_m | \hat{U}(t_m, t_i) | \{x\}_i \rangle e^{iS[\text{path}]/\hbar}
\]

So we have constructed all possible paths from $i$ to $f$ and split them into class $g_2$ in the region $i$ to $m$ and then class $g_1$ in the region $m$ to $f$. When we compose these paths we will get a path of type $g_1g_2$. The prefactors of the paths $\rho(g_1)$ and $\rho(g_2)$ then multiply and we get $\rho(g_1)\rho(g_2) = \rho(g_1g_2)$ since $\rho$ is a representation (the preservation of multiplication is the definition of being a representation! See section \ref{sec:rep}). So the prefactor of a given path from $i$ to $f$ is correctly given by $\rho(g)$ where $g$ is the topological class of the path. In other words, the form shown in Eq. 3.5 is properly preserved under composition, which is what is required in quantum mechanics!

### 3.4 Abelian Examples

Let us consider the case where the representation $\rho$ of our group $G$ of paths through configuration space is one dimensional — in other words it is a mapping from $g$ to a complex phase.\footnote{We call these cases \textit{abelian} since the group $G$ is commutative.}

This case seems to be most applicable in the quantum mechanics we know, because this representation is acting on the wavefunction of our system — and we are quite familiar with the idea of wavefunctions accumulating a complex phase.

#### 3.4.1 3+1 Dimensions

In 3+1 D, the group $G$ of paths through configuration space is the symmetric group $S_N$. It turns out that there are \textit{only two possible}\footnote{See exercise \ref{ex:rep} This is a fairly short proof!} one-dimensional representations of $S_N$:

- **Trivial rep:** In this case $\rho(g) = 1$ for all $g$. This corresponds to bosons. The path integral is just a simple sum over all possible paths with no factors inserted.
- **Alternating (or sign) rep:** In this case $\rho(g) = +1$ or $-1$ depending on whether $g$ represents an even or odd number of exchanges.
In this case the sum over all paths gets a positive sign for an even number of exchanges and a negative sign for an odd number. This is obviously fermions and is the generalization of the two particle example we considered above in section 3.2 where the exchange was assigned a $-1$.

### 3.4.2 2+1 Dimensions

In 2+1 D, the group $G$ of paths through configuration space is the braid group $B_N$. We can describe the possible one-dimensional representations by a single parameter $\theta$. We write the representation

$$\rho(g) = e^{i\theta W(g)}$$

where $W$ is the winding number of the braid $g$. In other words, a clockwise exchange accumulates a phase of $e^{i\theta}$ whereas a counterclockwise exchange accumulates a phase of $e^{-i\theta}$.

- For $\theta = 0$ there is no phase, and we simply recover bosons.
- For $\theta = \pi$ we accumulate a phase of $-1$ for each exchange no matter the direction of the exchange (since $e^{i\pi} = e^{-i\pi}$). This is fermions.
- Any other value of $\theta$ is also allowed. This is known as anyons, or fractional statistics. They are also known as abelian anyons in contrast with the nonabelian case which we will discuss in a moment.

The fact that this fractional statistics is consistent in quantum mechanics was first point out by Leinaas and Myrheim [1977] and popularized by Wilczek [1982]. Soon thereafter, Halperin [1984] showed theoretically that anyons really occur in fractional quantum Hall systems. We will examine these physical systems in detail starting in chapter ??.

### 3.5 Nonabelian Case

Can we do something more interesting and exotic by using a higher dimensional representation of the group $G = B_N$ of paths in configuration space? Generally in quantum mechanics, higher dimensional representations correspond to degeneracies, and indeed this is what is necessary.

Suppose we have a system with $N$ particles at a set of positions $\{x\}$. Even once we fix the positions (as well as the values of any local quantum numbers, like any “color” or “flavor” or ”spin” degree of freedom associated with the particle), suppose there still remains an $M$-fold degeneracy of the state of the system. We might describe the $M$ states as $|n; \{x\}\rangle$ for $n = 1, \ldots, M$. An arbitrary wavefunction of the system can then be expressed as:

$$|\psi\rangle = \sum_n A_n |n; \{x\}\rangle$$

If we want $|\psi\rangle$ normalized then there is a normalization condition on the $A_n$ coefficients. For example, if the $|n; \{x\}\rangle$’s are orthonormal then we need

$$\sum_n |A_n|^2 = 1$$

in order that $|\psi\rangle$ is normalized.

---

22There is no reason why this should not have been discovered in the 1930s, but no one bothered to think about it. It is a lucky coincidence that an experimental system of anyons was discovered so soon after the theoretical proposal (fractional quantum Hall effect, discovered by Tsui, Stormer, and Gossard [1982, see chapter ??], since the original theoretical work was entirely abstract, and they were not thinking about any particular experiment.

23The use of the braid group for describing statistics in two dimension dates back to Goldin et al. [1983]. The use of the symmetric group for understanding statistics in three dimensions appears to go back further to Laidlaw and DeWitt [1971].

24Among other things, Wilczek coined the term anyon. (He also won a Nobel Prize for asymptotic freedom.)

25If we want $|\psi\rangle$ normalized then there is a normalization condition on the $A_n$ coefficients. For example, if the $|n; \{x\}\rangle$’s are orthonormal then we need $\sum_n |A_n|^2 = 1$ in order that $|\psi\rangle$ is normalized.
with the $A_n$'s being some complex coefficients. Given the $N$ positions $\{x\}$, a general wavefunction should be thought of as a vector in $M$ dimensional complex space. Now that we have a vector, we can use an $M$-dimensional representation of the braid group in our path integral! We thus identify that $\rho(g)$ in Eq. 3.5 is an $M$ by $M$ unitary matrix

$$\rho(g) \rightarrow [U(g)]_{n,n'}$$

which is a representation of $G$ and must also be unitary so as to assure that probability is conserved. The propagator in Eq. 3.5 should now be thought of as a propagator between the initial ket $|n; \{x\}\rangle$ and the final bra $\langle n'; \{x\}|$. The unitary matrix $U(g)$ will act on the coefficients $A_n$ (which is a vector) in Eq. 3.6.

Let us now consider the process shown in Fig. 3.8. Here an initial wavefunction is represented as shown in Eq. 3.6 as a vector $A_n^{(i)}$ multiplying basis states $|n; \{x\}\rangle$ as in Eq. 3.6. We braid the particles around each other in some braid $g$ and bring them back to the same positions. After braiding the wavefunction should still be composed of the same basis states $|n; \{x\}\rangle$ since the particles are at the same positions and thus can be written in the form of Eq. 3.6 with a vector $A_n^{(f)}$. The final vector is obtained from the initial vector simply by multiplying by the unitary operator which is the representation of our braid group element $g$

$$A_n^{(f)} = [U(g)]_{n,n'} A_n^{(i)}$$

A particle that obeys this type of braiding statistics is known as a nonabelian anyon, or nonabelion. The word “nonabelian” means non-commutative, and the term is used since generically matrices (in this case the $U$ matrices) don’t commute.

In general the Hilbert space dimension $M$ will be exponentially large in the number of particles $N$. We define a quantity $d$, known as the quantum dimension such that

$$M \sim d^N$$

where the $\sim$ means that it scales this way in the limit of large $N$. We will see a lot more of this quantity $d$ later. It is not coincidence that we used the symbol $d$ previously in the context of Kauffman anyons! (See Eq. 2.1) We will see in section 17.1 that (up to a possible sign) this quantum dimension $d$ is actually the value $d$ of the unknot.

3.5 Nonabelian Case

Some Quick Comments on Quantum Computing:

Quantum Computing is nothing more than the controlled application of unitary operations to a Hilbert space. Unitary operations is exactly what we can do by braiding nonabelions around each other! I.e., we are

\[
|\psi(x)\rangle = \sum_{n=1}^{M} A_n |n; \{x\}\rangle
\]
multiplying a vector by a unitary matrix. Thus we see how braiding of particles, as discussed in chapter 2 can implement quantum computation. In chapter 11 we will give some more explicit descriptions of how one does quantum computation by braiding anyons.

3.5.1 Parastatistics in 3+1 Dimensions

Is it possible to have exotic nonabelian statistics in 3+1 dimensions? Indeed, there do exist higher dimensional representations of the symmetric group, so one can think about particles that obey more complicated statistics even in 3+1 dimensions — which is often known as parastatistics. However, it turns out that, subject to some “additional constraints”, it is essentially not possible to get anything fundamentally new — all we get is bosons and fermions and possibly some internal additional degrees of freedom. The proof of this statement is due to Doplicher et al. [1971, 1974] and took some 200 pages when it was first proven.

However, we should realize that in making statements like this, the fine print is important. As I mentioned in the previous paragraph we want to add some “additional constraints” and these are what really limit us to just bosons and fermions. What are these additional constraints?

1. We want to be able to pair create and annihilate. This means we are not just considering the braid group, but rather a more complicated structure that allows not just braiding particles around each other, but also creating and annihilating and even merging particles by bringing them together. This structure is given by category theory, some parts of which we will encounter (in simplified language) starting in chapter 8.

2. We also want some degree of locality. If we do an experiment on Earth, while off on Jupiter someone creates a particle-antiparticle pair, we would not want the particles on Jupiter to effect the result of our experiment on earth at all.

These two restrictions are crucial to reducing the 3+1 D case to only bosons and fermions. We will not go through the full details of how this happens. However, once we see the full structure of anyons in 2+1 dimensions, it ends up being fairly clear why 3+1 dimensions will be so restrictive. We return to this issue in section 20.3 where we will give further discussion.

We should note that despite this important result, 3+1 D is certainly not boring — but in order to get “interesting” examples, we have to relax some of our constraints. For example, if we relax the condition that “particles” are pointlike, but consider string-like objects instead, then we can have exotic statistics that describe what happens when one loop of string moves through another (or when a point-like particle moves through a loop of string). We would then need to consider the topology of the world-sheets describing loops moving through time.
Chapter Summary

- The path integral formulation of quantum mechanics requires us to add up all possible paths in space-time.
- We can add all of these paths in any way that preserves the composition law and the different possibilities allow for different types of particle statistics.
- The topologically different paths of \(N\) particles in space-time form a group structure (the fundamental group of the configuration space) which is the permutation group \(S_N\) in 3+1 dimensions, but is the braid group \(B_N\) in 2+1 dimensions.
- Particle braiding statistics must be a representation of this group.
- In 3+1 dimensions we can only have bosons and fermions, but in 2+1 dimensions we can have nontrivial braiding statistics which may be abelian (or “fractional”) or nonabelian.
- Quantum computation can be performed by braiding with certain nonabelian representations.

Further Reading

- For more discussion of particle statistics, a nice albeit somewhat dated book is Wilczek [1990].
- A good review discussing many aspects of exotic statistics is Nayak et al. [2008].

For a basic primer on path integrals see

- The classic reference on the subject is Feynman and Hibbs [1965].

Exercises

Exercise 3.1 About the Braid Group

(a) Convince yourself geometrically that the defining relations of the braid group on \(M\) particles \(B_M\) are:

\[
\sigma_i \sigma_{i+1} \sigma_i = \sigma_{i+1} \sigma_i \sigma_{i+1} \quad 1 \leq i \leq M - 2 \quad (3.9)
\]

\[
\sigma_i \sigma_j = \sigma_j \sigma_i \quad \text{for } |i - j| > 1, \quad 1 \leq i, j \leq M - 1 \quad (3.10)
\]

(b) Instead of thinking about particles on a plane, let us think about particles on the surface of a sphere. In this case, the braid group of \(M\) strands on the sphere is written as \(B_M(S^2)\). To think about braids on a sphere, it
is useful to think of time as being the radial direction of the sphere, so that
braids are drawn as in Fig. 3.9.

The braid generators on the sphere still obey Eqns. \(3.9\) and \(3.10\) but they
also obey one additional identity

\[
\sigma_1 \sigma_2 \ldots \sigma_{M-2} \sigma_{M-1} \sigma_{M-1} \sigma_{M-2} \ldots \sigma_2 \sigma_1 = I
\]

(3.11)

where \(I\) is the identity (or trivial) braid. What does this additional identity
mean geometrically?

[In fact, for understanding the properties of anyons on a sphere, Eq. \(3.11\)
is not quite enough. We will try to figure out below why this is so by using
Ising Anyons as an example.]

**Exercise 3.2 About the Symmetric Group**

Show that Eqs. \(3.9\) and \(3.10\) also hold for the generators of the symmetric

**Exercise 3.3 Ising Anyons and Majorana Fermions**

The most commonly discussed type of nonabelian anyon is the Ising anyon
(we will discuss this in more depth later). Ising anyons occurs in the Moore-
Read quantum Hall state \( (\nu = 5/2)\), as well as in any chiral \(p\)-wave supercon-
derator and in recently experimentally relevant so called “Majorana” systems.

The nonabelian statistics of these anyons may be described in terms of
Majorana fermions by attaching a Majorana operator to each anyon. The
Hamiltonian for these Majoranas is zero – they are completely noninteracting.

In case you haven’t seen them before, Majorana Fermions \(\gamma_j\) satisfy the
anticommutation relation

\[
\{\gamma_i, \gamma_j\} \equiv \gamma_i \gamma_j + \gamma_j \gamma_i = 2 \delta_{ij}
\]

(3.12)
as well as being self conjugate \(\gamma_i^\dagger = \gamma_i\).

(a) Show that the ground state degeneracy of a system with \(2N\) Majoranas
is \(2^N\) if the Hamiltonian is zero. Thus conclude that each pair of Ising anyons
is a two-state system. Hint: Construct a regular (Dirac) fermion operator
from two Majorana fermion operators. For example,

\[
c^\dagger = \frac{1}{2} (\gamma_1 + i \gamma_2)
\]

will then satisfy the usual fermion anti-commutation \(\{c, c^\dagger\} = cc^\dagger + c^\dagger c = 1\).

(If you haven’t run into fermion creation operators yet, you might want to
read up on this first!) There is more discussion of this transformation in later
exercises 9.7 and 10.2.

(b) When anyon \(i\) is exchanged clockwise with anyon \(j\), the unitary trans-
formation that occurs on the ground state is

\[
U_{ij} = \frac{e^{i\alpha}}{\sqrt{2}} [1 + \gamma_i \gamma_j]
\]

\(i < j\).

(3.13)

for some real value of \(\alpha\). Show that these unitary operators form a representation
of the braid group. (Refer back to the previous problem, “About the
Braid Group”). In other words we must show that replacing \(\sigma_i\) with \(U_{i,i+1}\)
Exercises 37

in Eqns. 3.9 and 3.10 yields equalities. This representation is $2^N$ dimensional since the ground state degeneracy is $2^N$.

(c) Consider the operator
\[
\gamma^\text{FIVE} = (i)^N \gamma_1 \gamma_2 \ldots \gamma_{2N}
\]
(3.14)
(the notation FIVE is in analogy with the $\gamma^5$ of the Dirac gamma matrices).

Show that the eigenvalues of $\gamma^\text{FIVE}$ are $\pm 1$. Further show that this eigenvalue remains unchanged under any braid operation. Conclude that we actually have two $2^{N-1}$ dimensional representations of the braid group. We will assume that any particular system of Ising anyons is in one of these two representations.

(d) Thus, 4 Ising anyons on a sphere comprise a single 2-state system, or a qubit. Show that by only braiding these four Ising anyons one cannot obtain all possible unitary operation on this qubit. Indeed, braiding Ising anyons is not sufficient to build a quantum computer. [Part (d) is not required to solve parts (e) and (f)]

(e) [bit harder] Now consider $2^N$ Ising anyons on a sphere (See above problem "About the braid group" for information about the braid group on a sphere). Show that in order for either one of the $2^{N-1}$ dimensional representations of the braid group to satisfy the sphere relation, Eqn. 5.11 one must choose the right abelian phase $\alpha$ in Eq. 5.13 Determine this phase.

(f) [a bit harder] The value you just determined is not quite right. It should look a bit unnatural as the abelian phase associated with a braid depends on the number of anyons in the system. Go back to Eqn. 6.11 and insert an additional abelian phase on the right hand side which will make the final result of part (e) independent of the number of anyons in the system. In fact, there should be such an additional factor — to figure out where it comes from, go back and look again at the geometric “proof” of Eqn. 3.11 Note that the proof involves a self-twist of one of the anyon world lines. The additional phase you added is associated with one particle twisting around itself. The relation between self-rotation of a single particle and exchange of two particles is a generalized spin-statistics theorem.

Exercise 3.4 Small Numbers of Anyons on a Sphere

On the plane, the braid group of two particles is an infinite group (the group of integers describing the number of twists!). However, this is not true on a sphere.

First review the problem “About the Braid Group” about braiding on a sphere.

(a) Now consider the case of two particles on a sphere. Determine the full structure of the braid group. Show it is a well known finite discrete group. What group is it?

(b) [Harder] Now consider three particles on a sphere. Determine the full structure of the braid group. Show that it is a finite discrete group. [Even Harder] What group is it? It is “well known” only to people who know a lot of group theory. But you can google to find information about it on the web with some work. It may be useful to list all the subgroups of the group and the multiplication table of the group elements.

(c) Suppose we have two (or three) anyons on a sphere. Suppose the ground state is two-fold degenerate (or more generally $N$-fold degenerate for some finite $N$). Since the braid group is discrete, conclude that no type of anyon statistics can allow us to do arbitrary $SU(2)$ (or $SU(N)$) rotations on this degenerate ground state by braiding
Aharanov-Bohm Effect and Charge-Flux Composites

This chapter introduces a simple model of how fractional statistics anyons can arise. After reviewing Aharanov-Bohm effect, we describe these exotic particles as charge-flux composites and explore some of their properties. Finally we see how this fits into the framework of abelian Chern-Simons theory and briefly discuss its nonabelian generalization.

4.1 Review of Aharanov-Bohm Effect

Let us consider the two slit interference experiment shown in Fig. 4.1. We all know the result of the two slit experiment but let us rewrite the calculation in the language of a path integral. We can write

\[ \sum_{\text{paths}} e^{iS/h} = \sum_{\text{paths, slit 1}} e^{iS/h} + \sum_{\text{paths, slit 2}} e^{iS/h} \sim e^{ikL_1} + e^{ikL_2} \]

where \( L_1 \) and \( L_2 \) are the path lengths through the two respective slits to whichever point is being measured on the output screen, and \( k \) is the wavevector of the incoming wave. In other words, we get the usual two slit calculation pioneered by Thomas Young in the early 1800s.

Now let us change the experiment to that shown in Fig. 4.2. Here we assume the particle being sent into the interferometer is a charged particle, such as an electron. In this case a magnetic field is added inside the middle box between the two paths. No magnetic field is allowed to leak out of the box, so the particle never experiences the magnetic field. Further the magnetic field is kept constant so the particle does not feel a Faraday effect either. The surprising result is that the presence of the magnetic field nonetheless changes the interference pattern obtained on the observation screen! This effect, named the Aharanov-Bohm effect, was predicted by Ehrenberg and Siday [1949], then re-predicted independently by Aharonov and Bohm [1959].

\[ \text{Possibly the reason it is named after the later authors is that they realized the importance of the effect, whereas the earlier authors pointed it out, but did not emphasize as much how strange it is! The first experimental observation of the effect was by Chambers [1960], although many more careful experiments have been done since.} \]
So why does this strange effect occur? There are several ways to understand it, but for our purpose it will be best to stay with the idea of path integrals and consider the Lagrangian description of particle motion.

We must recall how a charged particle couples to an electromagnetic field in the Lagrangian description of mechanics. We write the magnetic field and electric field in terms of a vector potential
\[
B = \nabla \times A
\]
\[
E = -\nabla A_0 - dA/dt
\]
where \(A_0\) is the electrostatic potential. We can then write the particle Lagrangian as
\[
L = \frac{1}{2m} \dot{x}^2 + q(A(x) \cdot \dot{x} - A_0)
\]
where \(q\) is the particle charge. It is an easy exercise to check that the Euler-Lagrange equations of motion that result from this Lagrangian correctly gives motion under the Lorentz force as we should expect for a charged particle in an electromagnetic field.

Here are the steps: Start with the Euler-Lagrange equations
\[
\frac{d}{dt} \frac{\partial L}{\partial \dot{x}_k} = \frac{\partial L}{\partial x_k}
\]
This gives us
\[
\frac{d}{dt}(m\dot{x}_k + qA_k) = m\ddot{x}_k + q \frac{d}{dt}A_k + q\dot{x}_j \frac{\partial}{\partial x_j} A_k
\]
So that
\[
m\ddot{x}_k = q(E + \dot{x} \times B)_k
\]

We are interested in a situation where we add a static magnetic field to the system. Thus, we need only include \(qA(x) \cdot \dot{x}\) in the Lagrangian. The action then gets changed by
\[
S \rightarrow S_0 + q \int dt \dot{x} \cdot A
\]
where \(S_0\) is the action in the absence of the magnetic field and the integral on the far right is a line integral along the path taken by the particle.

Returning now to the two slit experiment. The amplitude of the process in the presence of the vector potential can be now rewritten as
\[
\sum_{\text{paths, slit 1}} e^{iS_0/h + iq/h \int \text{d}l \cdot A} + \sum_{\text{paths, slit 2}} e^{iS_0/h + iq/h \int \text{d}l \cdot A}
\]
where \(S_0\) is again the action of the path in the absence of the vector potential.

The physically important quantity is the difference in accumulated phases between the two paths. This difference is given by
\[
\exp \left[ i \frac{q}{\hbar} \int_{\text{slit 1}} \text{d}l \cdot A - i \frac{q}{\hbar} \int_{\text{slit 2}} \text{d}l \cdot A \right] = \exp \left[ i \frac{q}{\hbar} \oint \text{d}l \cdot A \right]
\]
where the integral on the right is around a loop that goes forward through slit 1 and then backwards through slit 2.

Using Stokes’ theorem, we have
\[
\frac{iq}{\hbar} \oint \text{d}l \cdot A = \frac{iq}{\hbar} \int \text{d}S \cdot (\nabla \times A) = \frac{iq}{\hbar} \Phi_{\text{enclosed}}
\]
4.2 Anyons as Charge-Flux Composites

Anyons as Charge-Flux Composites

We will now consider a simple model of abelian anyons as charge-flux composites. Imagine we have a two dimensional system with charges \( q \) in them, where each charge is bound to an infinitely thin flux tube through the plane, with each tube having flux \( \Phi \) as shown in Fig. 4.3. We will notate this charge-flux composite object as a \((q, \Phi)\) particle. If we drag one such particle around another, we then accumulate a phase due to the Aharonov-Bohm effect. The phase from the charge of particle 1 going around the flux of particle 2 is \( e^{i q \Phi / \hbar} \), whereas the phase for dragging the flux of 1 around the charge of 2 is also \( e^{i q \Phi / \hbar} \), thus the total phase for dragging 1 around 2 is given by

\[
\text{(Phase of charge-flux composite 1 encircling 2)} = e^{2i q \Phi / \hbar}
\]

Thus we have (as shown in Fig. 4.4)

\[
\text{(Phase for exchange of two charge-flux composites)} = e^{i q \Phi / \hbar}
\]

and we correspondingly call these particles \( \theta \)-anyons, with \( \theta = q \Phi / \hbar \). Obviously \( \theta = 0 \) is bosons, \( \theta = \pi \) is fermions, but other values of \( \theta \) are also allowed, giving us abelian anyons as discussed in chapter 3.

Note that the same type of calculation would show us that taking a composite particle with charge \( q_1 \) and flux \( \Phi_1 \) all the way around a composite particle with charge \( q_2 \) and flux \( \Phi_2 \) would accumulate a phase of \( e^{i \varphi} \) with \( \varphi = (q_1 \Phi_2 + q_2 \Phi_1) / \hbar \).

\[\text{Fig. 4.3 Abelian anyons represented as charges bound to flux tubes through the plane. The charge of each particle is } q, \text{ the flux of each tube is } \Phi. \text{ Dragging one particle around another incurs a phase both because charge is moving around a flux, but also because flux is moving around a charge.}\]

\[\text{Fig. 4.4 An exchange. Two exchanges is the same as dragging one particle all the way around the other as shown in Fig. 4.3.}\]
Aharanov-Bohm Effect and Charge-Flux Composites

Spin of an anyon

Let us see if we can determine the spin of these anyons. Spin refers to properties of the rotation operator, so we need to physically rotate the anyon on its axis. To do this we must think about how the flux is tied to the charge — we must have some microscopic description of exactly where the flux is and where the charge is. It is easiest to put the charge and flux at very slightly different positions as shown in Fig. 4.5.

4 Almost any prescription for attaching flux to charge (for example, break the flux into four pieces and attach one piece on each of four side of the charge) will give the same result. However, if we try to put the flux and charge at exactly the same position, we get infinities that we don’t know how to handle!

This fits very nicely with the spin statistics theorem — the phase obtained by exchanging two identical particles should be the same as the phase obtained by rotating one around its own axis. (See the discussion of Fig. 2.7).

4.2.1 Fusion of Anyons

We can consider pushing two anyons together to try to form a new particle. We expect that the fluxes will add and the charges will add. This makes some sense as the total charge and total flux in a region should be conserved (this is an important principle that we will encounter frequently!). We sometimes will draw a “fusion diagram” as in Fig. 4.6 to show that two anyons have come together to form a composite particle.

A simple example of this is pushing together two particles both having the same charge and flux \((q, \Phi)\). In this case we will obtain a single particle with charge \((2q)\) and flux \((2\Phi)\).

4.2.2 Anti-Anyons and the Vacuum Particle

We now introduce the concept of an anti-anyon. This is a charge-flux composite which instead of having charge and flux \((q, \Phi)\) has charge and flux \((-q, -\Phi)\). Fusing an anyon with its anti-anyon results in pair annihilation — the two particles come together to form the vacuum (which we sometimes refer to as the identity \(I\)) which has zero total charge and zero total flux, as shown in Fig. 4.7. It may seem a bit odd to call the absence of any charge or any flux a “particle”. However, this is often convenient since it allows us to think of pair annihilation (as in the left of Fig. 4.7) in the language of fusion.

In the right of Fig. 4.7 we show that it is sometimes convenient not to indicate the vacuum particle. In this case, we have written the anti-anyon moving forward in time as an anyon moving backwards in time.

5 The vacuum or identity particle can be denoted \(1\), or \(I\) or \(0\) or \(1\) depending on the context. This nomenclatural problem stems from a similar problem in group theory, see section 41.2.

\[
e^{iq\Phi/h} = e^{i\theta}
\]
flux multiplies negative charge!). However, the phase of dragging an anyon clockwise around an anti-anyon is $-2\theta$.

### 4.3 Anyon Vacuum on a Torus and Quantum Memory

A rather remarkable feature of topological models is that the ground state somehow “knows” what kind of anyons exist in the model (i.e., those that could be created), even when they are not actually present. To see this, consider the ground state of an anyon model on torus (the surface of a doughnut).

We can draw the torus as a square with opposite edges identified as shown in Fig. 4.8. The two cycles around the torus are marked as $C_1$ and $C_2$.

Let us now construct operators that do the following complicated operations:

- $T_1$ is the operator that creates a particle-antiparticle pair, moves the two in opposite directions around the $C_1$ cycle of the torus until they meet on the opposite side of the torus and reannihilate.

- $T_2$ is the operator that creates a particle-antiparticle pair, moves the two in opposite directions around the $C_2$ cycle of the torus until they meet on the opposite side of the torus and reannihilate.

Both of these operators are unitary because they can be implemented (in principle) with some time-dependent Hamiltonian. However, the two operators do not commute. To see this let us consider the operator $T_2^{-1}T_1^{-1}T_2T_1$, where we read time from right to left. This can be interpreted as two particles being created, braiding around each other, and then reannihilating. This procedure is shown in Fig. 4.9.

So what we have now is two operators $T_1$ and $T_2$ which do not commute with each other. Indeed, we have:

$$T_2T_1 = e^{-2i\theta}T_1T_2$$

But both $T_1$ and $T_2$ commute with the Hamiltonian (since they start and end with states of exactly the same energy). Whenever you have two operators that don’t commute with each other but do commute with the Hamiltonian, it means you have degenerate eigenstates. Let us see how this happens.

Since $T_1$ is unitary, its eigenvalues must have unit modulus (i.e., they are just a complex phase). Considering the space of possible ground states, let us write a ground state eigenstate of $T_1$ as

$$T_1|\alpha\rangle = e^{i\alpha}|\alpha\rangle.$$

Note that we are labeling the ket $|\alpha\rangle$ by its eigenvalue under the application of $T_1$.
Aharanov-Bohm Effect and Charge-Flux Composites

The torus is drawn as a horizontal rectangle with opposite ends identified. Time runs vertically. First create a particle-antiparticle pair at the center of the rectangle and move them in opposite directions, right and left, until they meet at the edges of the rectangle to reannihilate. Note that a particle moving to the right or an antiparticle moving to the left are both drawn as a rightpointed arrow. Next create a particle-antiparticle pair in the center of the torus and move them to the front and back walls (which are the same point) to reannihilate. Then the two processes are reversed to give $T_2^{-1} T_1^{-1} T_2 T_1$. This procedure can be reduced to one particle wrapping around another which gives a phase of $e^{-2i\theta}$. Note that to make the figure on the left look like the linked rings, we should not quite annihilate the particles at the end of the first and second step (turning the dotted lines into solid lines). This is allowed since bringing a particle-anti-particle pair close together looks like they have fused together to the vacuum if we view it from far away (This is true for abelian anyons as it is a special case of the identity in Fig. 12.19.)

Let us suppose we have a system where the anyons have statistical phase angle $\theta = \pi p/m$ where $p$ and $m$ are relatively prime integers (i.e., $p/m$ is an irreducible fraction). Starting with the ground state $|\alpha\rangle$ we can generate a series of ground states by successive application of $T_2$,

$$|\alpha\rangle, |\alpha + 2\pi p/m\rangle, |\alpha + 4\pi p/m\rangle, \ldots, |\alpha + 2\pi (m-1)/m\rangle$$

When we try to generate yet another state, we get the phase $\alpha + 2\pi$ which is equivalent to $\alpha$ since it is describing a complex phase, so we are back to the original state. So we now have $m$ independent ground states.$^{10}$ Note in particular that the ground state degeneracy of the system with no anyons in it is related to the statistical angle $\theta$ of the anyons if they were to be created.
4.3.1 Quantum Memory and Higher Genus

The degenerate ground state on the torus can be thought of as a quantum memory. If there are \( m \) different ground states, the most general wavefunction we can have is some linear superposition of the multiple ground states

\[
|\Psi\rangle = \sum_{n=0}^{m-1} A_n |\alpha + 2\pi np/m\rangle
\]

where the coefficients \( A_n \) form an arbitrary (but normalized) complex vector. We can initialize the system in some particular superposition (i.e., some vector \( A_n \)) and we can expect that the system remains in this superposition. The only way that this superposition can change is if a \( T_1 \) or \( T_2 \) operation is performed, or some combination thereof — i.e., if a pair of anyons appears from the vacuum moves around a nontrivial cycle of the torus and then reannihilates. Such a process can be extremely unlikely when the energy gap for creating excitations is large\(^{11}\). Hence the quantum superposition is “topologically protected”.

In fact, one does not even need to have a system on a torus in order to have a degenerate ground state. It is often sufficient to have an annulus geometry (a disk with a big hole in the middle as shown in Fig. 4.10). In this case, \( T_1 \) could correspond to moving an anyon around the loop of the annulus and \( T_2 \) could correspond to moving an anyon from the inside to the outside edge.\(^{12}\)

One can consider more complicated geometries, such as a torus with multiple handles, or a disk with multiple holes cut in the middle. For a theory of abelian anyons (fractional statistics) the ground state degeneracy for a surface with genus \( g \) (meaning \( g \) handles, or \( g \) holes) is \( m^g \) (See exercise 4.1). Thus by using high genus one can obtain very very large Hilbert spaces in which to store quantum information.

4.3.2 Number of Species of Anyons

Having established multiple vacuum states on a torus, let us now return to study the anyons that we could create in such a system. Again let us consider anyons of statistical angle \( \theta = \pi p/m \) with \( p \) and \( m \) relatively prime. We can describe such anyons\(^{13}\) with a charge-flux composite \((q, \Phi) = (\pi p/m, 1)\). Fusion of \( n \) of these elementary anyons will have charge and flux given by\(^{14}\)

\[
\text{Fusion of } n \text{ elementary anyons } = |"n"\rangle = (nq, n\Phi) = (n\pi p/m, n)
\]

Something special happens when we have a cluster of \( m \) of these elementary anyons:

\[
|"m"\rangle = (\pi p, m)
\]

If we braid an arbitrary cluster \(|"n"\rangle = (n\pi p/m, n)\) around one of these \(|"m"\rangle = (\pi p, m)\) clusters, we obtain a net phase\(^{15}\) of \(2n\pi p\) which is

\[^{11}\text{Strictly speaking, at any finite temperature for any size system there is a finite time for this process to occur, although it might be very long.}\]

\[^{12}\text{In this case it is often not precisely true that the ground states are entirely degenerate (since there is a non-zero net result of having moved a particle from inside to outside, and therefore one is not necessarily in the precise ground state) but under certain conditions it can be extremely close to degenerate nonetheless. A classic example of this is discussed by Gefen and Thouless [1993].}\]

\[^{13}\text{By this time I’m sick of writing } \hbar \text{ and I’m going to set it equal to 1.}\]

\[^{14}\text{It is only a slight abuse of notation to write the ket } |"n"\rangle \text{ to mean a cluster of } n \text{ elementary anyons.}\]

\[^{15}\text{As mentioned at the beginning of section 4.2 the total phase is given by } q_1\Phi_2 + q_2\Phi_1 = (n\pi p/m)m + (\pi p)n.\]
equivalent to no phase at all! Thus we conclude that the cluster of \( m \) elementary anyons is equivalent to the vacuum in the sense that all particles get trivial phase if they braid all the way around \(|m\rangle\). Sometimes one says that \(|m\rangle\) is a \textit{transparent} particle.

We might be tempted to conclude that there are exactly \( m \) different anyon species in the system. Indeed, this conclusion is often true. However, there is an exception. If both \( p \) and \( m \) are odd, one obtains a nontrivial sign for exchanging (half braiding, as in Fig. 4.4) a \(|m\rangle\) = \((\pi p, m)\) particle with another \(|m\rangle\) = \((\pi p, m)\) particle. To see this note that exchange gives a phase \( \pi pm \) since it is half of the \( 2\pi pm \) phase for wrapping one particle all the way around the other (as in Fig. 4.3). This means the \(|m\rangle\) particle is a fermion. In fact, this case of \( p \) and \( m \) both odd is a bit of an anomalous case and is a bit more difficult to handle.

Neglecting this more complicated case with transparent particles, we are correct to conclude that we have exactly \( m \) different species of anyons — and also \( m \) different ground states on the torus as calculated above. This connection will occur in any well behaved topological theory — the number of ground states on the torus will match the number of different species of particles.

\[\text{Chapter Summary}\]

- The Charge-Flux composite model describes abelian anyons — with the braiding phase coming from Aharonov-Bohm effect.
- We introduced idea of fusion, antiparticles, and spin
- The vacuum for a system of anyons is nontrivial and can be a quantum memory.

\[\text{Further Reading}\]

A good reference for the charge-flux composite model is John Preskill’s lecture notes (Preskill [2004]).

\[\text{Exercises}\]

\text{Exercise 4.1 Abelian Anyon Vacuum on a Two-Handle Torus}

Using similar technique as in section 4.3 show that the ground state vacuum degeneracy on a two handle torus is \( m^2 \) for a system of abelian anyons with statistical angle \( \theta = \pi pm \) for integers \( p \) and \( m \) relatively prime. Hint: Consider what the independent nontrivial cycles are on a two-handled torus and determine the commutation relations for operators \( T_i \) that take anyon-antianyon pairs around these cycles.
Chern-Simons Theory Basics

5.1 Abelian Chern-Simons Theory

It is useful to see how charge-flux binding occurs in a microscopic field theory description of a physical system. The type of field theory we will study, so-called “Chern-Simons” field theory\(^1\), is the main paradigm for topological quantum field theories.

In the current section we will consider the simplest type of Chern-Simons theory which is the abelian type (i.e., it generates abelian anyons, or simple fractional statistics particles). We start by imagining a gauge field \(a_\alpha\), known as the Chern-Simons vector potential, analogous to the vector potential \(A_\alpha\) we know from regular electromagnetism. Here we should realize that \(a_\alpha\) is not the real electromagnetic vector potential because it lives only in our 2-dimensional plane. We should think of it instead as some emergent effective quantity for whatever two dimensional system we are working with.

Let us write the Lagrangian of our system

\[
L = L_0 + \int d^2 x \, \mathcal{L}
\]

Here we have written \(L_0\) to be the Lagrangian of our particles without considering the coupling to the (Chern-Simons) vector potential. This might be nothing more than the Lagrangian for free particles — although we could put other things into this part too, such as inter-particle interaction, if we like.

The second term is the integral of a Lagrangian density — and this will be the term that is relevant for the flux-binding and the exchange statistics of the particles. The form of the Lagrangian density is

\[
\mathcal{L} = \frac{\mu}{2} \epsilon^{\alpha\beta\gamma} a_\alpha \partial_\beta a_\gamma - j^\alpha a_\alpha
\]

where \(j^\alpha\) is the particle current, \(\mu\) is some coupling constant, and \(\epsilon\) is the antisymmetric tensor\(^2\). The indices \(\alpha, \beta, \gamma\) take values 0, 1, 2 where 0 indicates the time direction and 1, 2 are the space directions (and \(j^0\) is the particle density).

The first term in Eq. 5.1 is the Lagrangian density of the Chern-Simons vector potential itself. (It is sometimes known as the “Chern-Simons term”). The second term in Eq. 5.1 couples the Chern-Simons vector potential to the particles in the system. Its form, \(j^\alpha a_\alpha\), may look unfamiliar but it is actually just the expected coupling of the charged particles with the potential.

\(^1\)S. S. Chern was one of the most important mathematicians of the 20th century. Jim Simons was a prominent mathematician who wrote the key first paper on what became known as Chern-Simons theory in 1974. Simons was the head of the math department at Stonybrook university at the time. In 1982, he decided to change careers and start a hedge fund. His fund, Renaissance Technologies, became one of the most successful hedge funds in the world. Simons’ wealth is now estimated at over 20 billion dollars (as of 2018). More recently he has become a prominent philanthropist, and has donated huge amounts of money to physics and mathematics — now being one of the major sources of funds for the best scientists in the world.

\(^2\)The antisymmetric tensor is given by

\[
\epsilon^{012} = \epsilon^{120} = \epsilon^{201} = 1 \quad \text{and} \quad \epsilon^{102} = \epsilon^{210} = -1.
\]
Chern-Simons Theory Basics

particles to a vector potential analogous to what we used when we discussed Aharonov-Bohm effect in section 4.1. To see this, let us carefully define the particle current $j^\alpha$. If we have $N$ particles then the current is

\[ j^0(x) = \sum_{n=1}^{N} q_n \delta(x - x_n) \]

\[ j(x) = \sum_{n=1}^{N} q_n \dot{x}_n \delta(x - x_n) \]

The $j^0$ component, the charge density\(^3\), is just a delta function peak at the position of each particle with value given by the particle charge $q$. The 1 and 2 component, $j$ is a delta function at the position of each particle with prefactor given by the velocity of the particle times its charge. Now when $-j^\alpha a_\alpha$ is integrated over all of space we get

\[ \sum_{n=1}^{N} q_n [a(x_n) \cdot \dot{x}_n - a_0(x_n)] \]

exactly as in Eq. [4.1]. So this is nothing more than the regular coupling of a system of charged particles to a vector potential.

As is usual for a gauge theory, the coupling of the particles to the gauge field is gauge invariant once one integrates the particle motion over some closed path (one measures only the flux enclosed, as with the Aharonov-Bohm effect). The Chern-Simons term (the first term in Eq. 5.1) is also gauge invariant, at least on a closed manifold\(^4\) if we can integrate by parts. To see this, make an arbitrary gauge transformation $a_\mu \rightarrow a_\mu + \partial_\mu \chi$ (5.3)

for any function $\chi$. Then integrating the Chern-Simons term (by parts if necessary) all terms can be brought to the form $\epsilon^{\alpha\beta\gamma} \partial_\alpha \partial_\beta a_\gamma$ which vanishes by antisymmetry. Note that this gauge invariance holds for any closed manifold, although for a manifold with boundaries, we have to be careful when we integrate by parts as we can get a physically important boundary term. (We will discuss these later in section *** but for now, let us just think about closed space-time manifolds).

To determine what the Chern-Simons term does we need to look at the Euler-Lagrange equations of motion. We have

\[ \frac{\partial L}{\partial a_\alpha} = \partial_\beta \left( \frac{\partial L}{\partial (\partial_\beta a_\alpha)} \right) \]

which generates the equations of motion\(^5\)

\[ j^\alpha = \mu \epsilon^{\alpha\beta\gamma} \partial_\beta a_\gamma \]

This equation of motion demonstrates flux binding. To see this, let us

\(^3\)Again not the real electromagnetic charge, but rather the charge that couples to the Chern-Simons vector potential $a_\alpha$. Later in this chapter we will set $q = 1$ along with $\hbar = 1$ for simplicity of notation.

\(^4\)We use the term closed manifold frequently. This means a manifold without boundary containing all its limit point. More detailed discussions and examples are given in section [5.1].

\(^5\)It may look like the right result would have $\mu/2$ on the right hand side, given that it is $\mu/2$ in Eq. [5.1]. However, note that when we differentiate with respect to $a_\alpha$ on the left hand side of Eq. [5.3] we also generate an identical factor of $\mu/2$ and these two add up.
look at the 0th component of this equation. We have

\[ j^0 = \sum_{n=1}^{N} q_n \delta(x - x_n) = \mu(\nabla \times a) = \mu b \]  

(5.6)

where we have defined a “Chern-Simons” magnetic field \( b \) to be the curl of the the Chern-Simons vector potential. In other words this equation attaches a delta function (infinitely thin) flux tube with flux \( q_n/\mu \) at the position of each charge \( q_n \). So we have achieved charge-flux binding!

Formally when we write a functional integral we mean over all possible values of

divide space and time into little boxes and within each box integrate all configurations in space and time so it is effectively a path integral

\[ \int \text{d}[a] \]  

and integrated out a different variable or combination thereof. For this reason we still write the subsequent equation Eq. 5.9 as a strict constraint. Once \( a_0 \) is integrated out, it is no longer a variable of the problem and we can think of this as choosing a \( a_0 = 0 \) gauge thereafter.

\( \delta \) and \( \theta \) as being a Lagrange multiplier which enforces Eq. 5.6. Similarly in the functional integral when we integrate out \( a_0 \) it enforces that equation of motion as a strict constraint. Once \( a_0 \) is integrated out, it is no longer a variable of the problem and we can think of this as choosing a \( a_0 = 0 \) gauge thereafter.

\( \delta \) could have chosen a different gauge and integrated out a different variable or combination thereof. For this reason we still write the subsequent equation Eq. 5.9 in gauge invariant form, although strictly speaking we should realize that once we have enforced the constraint there are fewer degrees of freedom.

\( \theta \) as choosing a \( a_0 = 0 \) gauge thereafter.

\[ \int \text{d}[a] \]

6. One might worry about whether we are actually allowed to plug the equations of motion back into the Lagrangian when we integrate out \( a_0 \). For example, classically we can think of \( a_0 = 0 \) as being a Lagrange multiplier which enforces Eq. 5.6. Similarly in the functional integral when we integrate out \( a_0 \) it enforces that equation of motion as a strict constraint. Once \( a_0 \) is integrated out, it is no longer a variable of the problem and we can think of this as choosing a \( a_0 = 0 \) gauge thereafter.

\[ \delta \] and \( \theta \) as being a Lagrange multiplier which enforces Eq. 5.6. Similarly in the functional integral when we integrate out \( a_0 \) it enforces that equation of motion as a strict constraint. Once \( a_0 \) is integrated out, it is no longer a variable of the problem and we can think of this as choosing a \( a_0 = 0 \) gauge thereafter.

\[ \delta \] could have chosen a different gauge and integrated out a different variable or combination thereof. For this reason we still write the subsequent equation Eq. 5.9 as a strict constraint. Once \( a_0 \) is integrated out, it is no longer a variable of the problem and we can think of this as choosing a \( a_0 = 0 \) gauge thereafter.

\( \delta \) could have chosen a different gauge and integrated out a different variable or combination thereof. For this reason we still write the subsequent equation Eq. 5.9 as a strict constraint. Once \( a_0 \) is integrated out, it is no longer a variable of the problem and we can think of this as choosing a \( a_0 = 0 \) gauge thereafter.

\[ \delta \] could have chosen a different gauge and integrated out a different variable or combination thereof. For this reason we still write the subsequent equation Eq. 5.9 as a strict constraint. Once \( a_0 \) is integrated out, it is no longer a variable of the problem and we can think of this as choosing a \( a_0 = 0 \) gauge thereafter.

\[ \delta \] could have chosen a different gauge and integrated out a different variable or combination thereof. For this reason we still write the subsequent equation Eq. 5.9 as a strict constraint. Once \( a_0 \) is integrated out, it is no longer a variable of the problem and we can think of this as choosing a \( a_0 = 0 \) gauge thereafter.

\[ \delta \] could have chosen a different gauge and integrated out a different variable or combination thereof. For this reason we still write the subsequent equation Eq. 5.9 as a strict constraint. Once \( a_0 \) is integrated out, it is no longer a variable of the problem and we can think of this as choosing a \( a_0 = 0 \) gauge thereafter.
where $S_0$ is the action of the particles following the path but not interacting with the gauge field, $S_{CS}$ is the action of the Chern-Simons gauge field alone (from the first term in Eq. 5.1). The final exponential in Eq. 5.9 represents the coupling (from the second term of Eq. 5.1) of the gauge field to the path of the particles — it is an integral that follows the path of the particles and integrates the vector potential along the path (see also Eq. 5.2). This is precisely the phase accumulated by a particle in the vector potential. It is an example of a Wilson-line operator, which we will see again shortly in section 5.2.

Once the integration over the Chern-Simons field is done, we obtain

$$
\sum_{\text{paths } \{x(t)\}} e^{iS_0/\hbar + i\theta W(\text{path})}
$$

where $W$ is the winding number of the path and $\theta$ is the anyon statistical angle. In other words, integrating out the Chern-Simons gauge field implements fractional statistics for the particles in the system, inserting a phase $e^{\pm i\theta}$ for each exchange!

### Vacuum Abelian Chern-Simons Theory

Something we have pointed out above in section 4.3 is that the vacuum of an anyon theory knows about the statistics of the particles, even when the particles are not present (i.e., the ground state degeneracy on a torus matches the number of particle species). Thus, in the absence of particles, we will be interested in

$$
Z(M) = \int_M \mathcal{D}a_{\mu}(x) e^{iS_{CS}/\hbar}
$$

where $M$ is the space-time manifold we are considering.

If we are consider a three dimensional manifold of the form $M = \Sigma \times S^1$ for a 2D manifold $\Sigma$ and $S^1$ represents time (compactified), this integral gives exactly the ground state degeneracy of the system. As we might expect, this quantity will be a topological invariant of the space-time manifold. That is, smooth deformations of $M$ do not change its value. (See chapter appendix, particularly section 5.3.2). This quantity $Z(M)$, often known as the partition function of the theory for the manifold $M$, will be of crucial importance as we learn more about topological theories in general in Chapter 7 below.

### 5.2 Nonabelian Chern-Simons theory: The paradigm of TQFT

Among 2+1 dimensional topological quantum systems, pretty much everything of interest is somehow related to Chern-Simon theory — however, we don’t generally have the luxury of working with abelian theory as we have been doing so far.
We can generalize abelian Chern-Simons theory by promoting the gauge field $a_\mu$ to be not just a vector of numbers, but rather a vector of matrices. More precisely, to construct a nonabelian Chern-Simons theory, we consider a vector potential that takes values in a Lie algebra. For example, if we choose to work with the Lie algebra of $SU(2)$ in the fundamental representation we can write a general element of this algebra as a sum of the three generators (proportional to $\sigma_x, \sigma_y, \sigma_z$) so that our Lie algebra valued gauge field is then

$$a_\mu(x) = a_\mu^a(x) \left( \frac{\sigma_a}{2i} \right)$$ (5.10)

where $\sigma_a$ are the Pauli matrices. Now that $a_\mu$ is matrix valued it becomes noncommutative and we have to be very careful about the order in which we write factors of $a_\mu$.

The fundamental quantity that we need to think about is the Wilson loop operators:

$$W_L = \text{Tr} \left[ P \exp \left( \int_L d\mu^a a_\mu^a \right) \right]$$ (5.11)

where here the integral follows some closed path $L$. This object, being the exponential of an integral of a vector potential, is essentially the nonabelian analogue of the Aharonov-Bohm phase of Eq. (4.3). In Eq. (5.11) the $P$ symbol indicates path ordering — analogous to the usual time ordering of quantum mechanics. The complication here is that $a_\mu(x)$ is a matrix, so when we try to do the integral and exponentiate, we have a problem that $a_\mu(x)$ and $a_\mu(x')$ do not commute. The proper interpretation of the path ordered integral is then to divide the path into tiny pieces of length $dl$. We then have

$$P \exp(\int_L d\mu^a a_\mu^a) = \ldots [1 + a_\mu(x_3)d\mu^a(x_3)] [1 + a_\mu(x_2)d\mu^a(x_2)] [1 + a_\mu(x_1)d\mu^a(x_1)] \ldots$$ (5.12)

where $x_1, x_2, x_3, \ldots$ are the small steps along the path $L$ (it does not matter if they are equally spaced or not). If it is a closed path as in Eq. (5.11) the trace (which is invariant under cyclic permutation) will give the same answer independent of where on the closed path we start.

The proper gauge transformation in the case of a nonabelian gauge field is given by

$$a_\mu \rightarrow U^{-1} a_\mu U + U^{-1} \partial_\mu U$$ (5.13)

Where $U(x)$ is a matrix (which is a function of position and time) which acts on the matrix part of $a_\mu$. Note that this is just the nonabelian analogue of the gauge transformation in Eq. (5.13). To see that this gauge transformation leaves the Wilson loop operators invariant (and hence is the right way to define a gauge transformation!) see section 5.3.10.

11If you have studied Yang-Mills theory, you already know about non-abelian vector potentials.

12See the introduction to Lie groups and Lie algebras in section 41.2. In brief: A Lie group is a group which is also a continuous manifold. A Lie algebra is the algebra of infinitesimal changes in this group. A prime example is the Lie group $SU(2)$ with algebra generated by $\sigma_j$ with $\sigma_j$’s being the Pauli operators. We write group elements as exponentials of the algebra $g = e^{i a^a \sigma^a}$.

13For general Lie algebras, we want to write $a_\mu = a_\mu^a T_a$ where $T_a$ are the Hermitian generators of the Lie algebra with $T_a = -T_a^\dagger$. This means that $[T_a, T_b] = f^{abc} T_c$, with $f$ the so-called structure constants of the Lie group, and $\text{Tr}[T_a T_b] = \frac{1}{2} \delta_{ab}$. In case of $SU(2)$ in the fundamental representation we have $T_a = -i a^a / 2$ with $f^{abc} = \epsilon^{abc}$. Be warned that other normalization conventions do exist, and changing conventions will insert seemingly random factors of $2$ or $i$ or worse.

14These are named for Ken Wilson, who won a Nobel Prize for his work on the renormalization group and critical phenomena. There is a legend that Wilson had very very few publications when he came up for tenure as a professor at Cornell. Only due to the strong recommendation of his senior colleague Hans Bethe (already a Nobel Laureate at the time) did he manage to keep his job. Bethe knew what Wilson had been working on, and vouched that it would be extremely important. His ground-breaking work on renormalization group was published the next year. Everything worked out for him in the end, but the strategy of not publishing is not recommended for young academics trying to get tenure.

15The factor of $i$ we usually have in the exponential of the Aharonov-Bohm phase (Eq. (4.3)) is missing because it has been absorbed into $a_\mu$ in Eq. (5.10) (See comment in note 13). The factors of $q$ and $h$ are missing because we have set them to one as every theorist should do.

16Here take $U = e^{ix}$ and note that a factor of $i$ is absorbed into the vector potential as mentioned in note 15.
Chern-Simons Theory Basics

The nonabelian Chern-Simons action is given by

\[ S_{CS} = \frac{k}{4\pi} \int_{M} d^3x \; \epsilon^{\alpha\beta\gamma} \; \text{Tr} \left[ a_{\alpha} \partial_{\beta} a_{\gamma} + \frac{2}{3} a_{\alpha} a_{\beta} a_{\gamma} \right] \]  

(5.14)

Note that the second term in the brackets would be zero if the \( a_{\alpha} \) were commutative. (In the abelian case above, we have no such term! See Eq. 5.1). We have not derived Eq. 5.14, but we will explain in a moment why it is the only expression we could have written down for the nonabelian generalization of the Chern-Simons action.

The Chern-Simons action is metric independent, which we show explicitly in the chapter appendix section 5.3.2. This means that space and time can be deformed continuously and the value of the action does not change. While this may not be obvious from looking at the form of the action, a large hint is that the action is written without any reference to the usual space-time metric \( g_{\mu\nu} \).

Since Chern-Simons theory is also a gauge theory, we would like the action to be gauge invariant. It turns out that the action is almost gauge invariant, as we will discuss momentarily. At any rate it is close enough to gauge invariant to be of use for us!

It turns out that the Chern-Simons action is actually unique in being both metric independent and also (at least almost) gauge invariant. In 2+1 dimensions, no other action can be written down which involves only one gauge field and has these two properties: topological invariance and gauge invariance. This is what makes Chern-Simons theory such a crucial paradigm for topological theories in 2+1 dimensions.

Let us now return to this issue of how the Chern-Simons action is only almost gauge invariant. First of all, if the manifold has a boundary, we will run into non-gauge invariant terms as mentioned below Eq. 5.3. For now, let us just assume that our manifold has no boundaries.

More crucially there is another issue with gauge invariance. Under gauge transformation (at least on a closed manifold) as in Eq. 5.13 the Chern-Simons action transforms to (See section 5.3.3 for more detail)

\[ S_{CS} \rightarrow S_{CS} + 2\pi \nu k \]  

(5.15)

where

\[ \nu = \frac{1}{24\pi^2} \int_{M} d^3x \; \epsilon^{\alpha\beta\gamma} \; \text{Tr} \left[ (U^{-1}\partial_{\alpha} U)(U^{-1}\partial_{\beta} U)(U^{-1}\partial_{\gamma} U) \right] \]  

(5.16)

Surprisingly the complicated expression in Eq. 5.16 (sometimes known as the Pontryagin index) is always an integer (See section 5.3.3 for more detail). The integer \( \nu \) gives the winding number of the map \( U(x) \) from the manifold into the gauge group\(^{17}\).

It may now look problematic that our Chern-Simons action is not a true gauge invariant (Eq. 5.15), but we note that the only thing entering our functional integral is \( e^{iS_{CS}} \), not the Chern-Simons action itself. Thus, so long as we choose \( k \), the so-called “level”, as an integer (and since the winding number \( \nu \) is also an integer), then we have a well
defined functional integral of the form

\[ Z(M) = \int_M D\alpha(x) \ e^{iS_{CS}} \]  

(5.17)

where the result \( Z(M) \) turns out to be a manifold invariant (see chapter appendix, section 5.3.2), meaning that smooth deformations of space and time do not change its value.

The insertion of the Wilson loop operator into the path integral gives a knot invariant of the link \( L \) that the Wilson loop follows. The fact that the result should be a topological invariant should not be surprising given the fact that the Chern-Simons action itself is metric independent and therefore independent under deformations of space and time\(^\text{18}\). Often we will think about our link as being embedded in a simple manifold like the three sphere, which we denote as \( S^3 \) (see section 4.1.1 for definition of \( S^3 \)).

So for example, to find the link invariant corresponding to the two linked strings in Fig. 5.1 we have

\[ \text{Knot Invariant} = \frac{Z(S^3, L_1, L_2)}{Z(S^3)} = \frac{\int_{S^3} D\alpha(x) \ W_{L_1} W_{L_2} \ e^{iS_{CS}}} {\int_{S^3} D\alpha(x) \ e^{iS_{CS}}} \]  

(5.18)

with \( W_L \) being the Wilson loop operators as in Eq. 5.11. Indeed, if we choose to work with the gauge group \( SU(2) \) at level \( k \) (working with the spin 1/2 representation of the group, i.e., with Pauli matrices) we obtain the Kauffman invariant of the knot with

\[ A = \frac{-1}{k+2}. \]

If we keep the same gauge group, but work with a different representation (for example, spin 1, rather than spin 1/2 in Eq. 5.10), we will obtain different “particle types” of the theory.

One can also choose to work with different gauge groups. Using \( SU(N) \) and choosing a level \( k \) one obtains the two parameter HOMFLY knot polynomial (the two parameters here being \( N \) and \( k \)). Similarly, using \( SO(N) \) at level \( k \) gives a two parameter Kauffman polynomial (not to be confused with the Kauffman bracket). Typically a Chern-Simons theory with gauge group \( G \) at level \( k \) is notated as \( G_k \) (For example, using \( SU(2) \) at level 2 we write the theory as \( SU(2)_2 \)). Changing the sign of \( k \) corresponds to taking the “mirror image” of the theory (the partition function is complex conjugated).

5.3 Appendix: Odds and Ends about Chern Simons Theory

5.3.1 Gauge Transforms with Nonabelian Gauge Fields

Let us define a Wilson-line operator, similar to the Wilson loop but not forming a closed loop, i.e., going along a curve \( C \) from space-time point
y to space-time point $x$, and we do not take the trace here

$$W_C(x, y) = P \exp \left( \int_C d\mu \, a_\mu \right)$$

Under a gauge transformation function $U(x)$ we intend that the Wilson line operator transform as

$$\tilde{W}_C(x, y) \rightarrow U(x)^{-1} \tilde{W}_C(x, y) \, U(y) \quad (5.19)$$

Clearly this obeys composition of paths, and will correctly give a gauge invariant result for a closed Wilson loop. Now let us see what is required for the gauge field $a_\mu$ such that Eq. (5.19) holds. We consider

$$\tilde{W}_C(x, x + dx) = 1 + a_\mu dx^\mu \quad (5.20)$$

and its transformation should be

$$\tilde{W}_C(x, x + dx) \rightarrow U(x)^{-1} \tilde{W}_C(x, x + dx) U(x + dx)$$
$$= U(x)^{-1} [1 + a_\mu dx^\mu] U(x + dx)$$
$$= U(x)^{-1} [1 + a_\mu dx^\mu] [U(x) + dx^\mu \partial_\mu U(x)]$$
$$= 1 + [U^{-1} a_\mu U + U^{-1} \partial_\mu U] dx^\mu \quad (5.21)$$

By comparing Eq. (5.20) and Eq. (5.21) we see that the gauge transform rule Eq. (5.13) correctly gives a gauge invariant Wilson loop operator.

### 5.3.2 Chern Simons Action is Metric Independent

You will often see books state that Eq. (5.14) must be metric independent because you don’t see the metric $g_{\mu\nu}$ written anywhere. But that kind of misses the point!

A differential geometer would see that one can write the Chern-Simons action in differential form notation

$$S_{CS} = \frac{k}{4\pi} \int (a \wedge da + \frac{2}{3} a \wedge a \wedge a)$$

which then makes it “obvious” that this is metric independent being the integral of a 3-form.

In more detail however, we must first declare how the gauge field transforms under changes of metric. It is a “1-form” meaning it is meant to be integrated along a line to give a reparameterization invariant result, such as in the Wilson loops. In other words, we are allowed to bend and stretch the space-time manifold, but the flux through a loop should stay constant. Under reparametrization of coordinates we have

$$\int da = \int dx^\mu a_\mu(x) = \int dx'^\mu \partial x'^\nu a_\nu(x')$$
This means that under reparameterization \( x'(x) \) we have

\[
a_\mu(x) = \frac{\partial x^\nu}{\partial x'^\mu} a_\nu(x')
\]
such that the line integral remains invariant under a reparameterization of the space.

Now, if we make this change on all of the \( a \)'s in the the Chern-Simons action we obtain

\[
\epsilon^{\alpha\beta\gamma} \text{Tr} \left[ a_\alpha \partial_\beta a_\gamma - \frac{2i}{3} a_\alpha a_\beta a_\gamma \right] \rightarrow \epsilon^{\alpha'\beta'\gamma'} \text{det} \left[ \frac{\partial x}{\partial x'} \right]
\]

But notice that the prefactor, including the \( \epsilon \), is precisely the Jacobian determinant and can be rewritten as

\[
\epsilon^{\alpha'\beta'\gamma'} \text{det} \left[ \frac{\partial x}{\partial x'} \right]
\]

Thus the three-dimensional Chern-Simons action integral can be changed to the \( dx' \) variables and the form of the integral is completely unchanged and thus depends only on the topological properties of the manifold.

In fact, this feature of the Chern-Simons Lagrangian is fairly unique. Given that we have a single gauge field \( a_\mu(x) \) this is the only (3-form) gauge invariant Lagrangian density we can write down which will give a topological invariant!

### 5.3.3 Winding Number: The Pontryagin Index

We would like to show that the integral in Eq. [5.16] is indeed always an integer. While doing this rigorously is difficult, it is not too hard to see roughly how it must be done. First, we note that, like the Chern-Simons action, it is the integral of a three form so it does not care about the metric on the manifold (this is not surprising being that this winding number arose from the Chern-Simons action). One can then reparameterize the manifold in terms of coordinates within the group, and convert the integral over space into an integral over the group. The only thing that is left unclear is then in the mapping \( U(x) : M \rightarrow G \) how many times the group is covered in this mapping. We then have immediately that the given definition of the winding number must be an integer times some constant. By construction of a few examples, one can see that the constant is indeed unity (See exercise 5.4). A more detailed discussion of this issue is given in [Vandoren and van Nieuwenhuizen 2008] and [Rajaraman 1982].
5.3.4 Framing of the Manifold — or Doubling the Theory

There is a bit of a glitch in Chern-Simons theory. We want the Chern-Simons functional $Z(\mathcal{M})$ to be a function of the topology of $\mathcal{M}$ only. This is almost true — it is true up to a phase. In order to get the phase, you need to specify one more piece of information which can be provided in several ways (often called a 2-framing\textsuperscript{19}). This additional piece of information is most easily described by saying that you need to specify a bit of information about the topology of the 4-manifold $\mathcal{N}$ that $\mathcal{M}$ bounds $\mathcal{M} = \partial \mathcal{N}$. It is a fact that all orientable closed 3-manifolds are the boundary of some 4-manifold — in fact, of many possible 4-manifolds. The phase of $Z(\mathcal{M})$ is sensitive only to the so-called “signature” of the 4-manifold $\mathcal{N}$. (Consult a book on 4-manifold topology if you are interested!)

The fact that the Chern-Simons theory should depend on some information about the 4-manifold that $\mathcal{M}$ bounds may sound a bit strange. It is in fact a sign that the Chern-Simons theory is “anomalous”. That is, it is not really well defined in 3-dimensions. If you try to make sense of the functional integral $\int \mathcal{D}a_\mu$, you discover that there is no well-defined limit by which you can break up space-time into little boxes and integrate over $a_\mu$ in each of these boxes. However, if you extend the theory into 4-dimensions, then the theory becomes well behaved. This is not unusual. We are familiar with lots of cases of this sort. Perhaps the most famous example is the fermion doubling problem. You cannot write down a time reversal invariant theory for a single chirality fermion in $D$ dimensions without somehow getting the other chirality. However, you can think of a system extended into $D + 1$ dimensions where one chirality ends up on one of the $D$-dimensional boundaries and the other chirality ends up on the other $D$ dimensional boundary\textsuperscript{20}. So to make Chern-Simons theory well-defined, you must either extend into 4D, or you can “cancel” the anomaly in 3D by, for example, considering two, opposite chirality Chern-Simons theories coupled together (so-called “doubled” Chern-Simons theory). The corresponding manifold invariant of a doubled theory gets $Z(\mathcal{M})$ from the righthanded theory and its complex conjugate from the left handed theory, thus giving an end result of $|Z(\mathcal{M})|^2$ which obviously won’t care about the phase anyway!

5.3.5 Chern Simons Theory as Boundary of a Four Dimensional Topological Theory

With the considerations of the previous section \textsuperscript{[5.3.4]} it is interesting to express Chern-Simons theory as the boundary theory of a 4D topological theory. To do this let us define the field strength tensor

$$F_{\mu \nu} = \partial_\mu A_\nu - \partial_\nu A_\mu + [A_\mu, A_\nu]$$
This definition matches the expression for the electromagnetic field strength in the case where the fields are abelian such that the commutator vanishes. However, more generally for nonabelian gauge theories (including Yang-Mills theory) the additional commutator term must be added.

In 4D we can define the dual field strength

\[ *F_{\mu\nu} = \frac{1}{2} \epsilon^{\mu\nu\lambda\rho} F_{\lambda\rho} \]

where \( \epsilon \) is the antisymmetric tensor. We now consider the following topological action on a 4D manifold \( N \)

\[ S = \frac{\theta}{16\pi^2} \int_N d^4x \ Tr [F_{\mu\nu} * F^{\mu\nu}] \]

This 4D action is well defined and non-anomalous, meaning it can be regularized and/or treated properly on a lattice.

With a bit of algebra the action can be rewritten as

\[ S = \frac{\theta}{8\pi^2} \int_N d^4x \ \partial_\mu G_\mu \]

where

\[ G_\mu = 2\epsilon_{\mu\nu\lambda\rho} Tr \left[ A_\nu \partial_\lambda A_\rho + \frac{2}{3} A_\nu A_\lambda A_\rho \right] \] (5.22)

Since the action can be written as the integral of a total derivative, it should give zero when integrated over a closed manifold \( N \). However when the manifold has a boundary one obtains

\[ S = \frac{\theta}{8\pi^2} \int_{\partial N} d^3x \ G_\mu v^\mu \]

where \( v^\mu \) is the unit vector normal to the boundary. Examining the form of Eq. 5.22 we realize that the action is precisely the Chern-Simons action on the 3D boundary manifold \( \partial N \).

### 5.3.6 Chern Simons Canonical Quantization for the Abelian Case

One can consider the Chern-Simons theory as a quantum mechanical theory with wavefunctions and operators (i.e., not in path integral language). To do this, we need to find the commutation relations. Working in the gauge \( a_0 = 0 \), in the Chern-Simons Lagrangian terms like \( \partial_\mu a_\mu \) multiply \( a_\mu \) and vice versa\(^{21}\). This means that \( a_\mu(x) \) is the momentum conjugate to \( a_x(x) \) and vice versa. We thus have the commutation relations

\[ [a_x(\vec{x}), a_y(\vec{x}'')] = i\hbar \frac{\mu}{\delta(\vec{x} - \vec{x}'')} \]

\(^{21}\)Note that for nonabelian Chern-Simons theories working in the \( a_0 = 0 \) gauge makes the \( a^3 \) term of the action vanish!
The arguments \( \vec{x} \) here live in 2 dimensions. Consider now the Wilson loop operators around two different nontrivial cycles of a torus

\[
W_j = \exp \left( i \frac{q}{\hbar} \oint_{L_j} d\vec{l} \cdot \vec{a} \right)
\]

where here \( j \) indicates we have a loop around either cycle 1 \((L_1)\) or cycle 2 \((L_2)\) of our torus. The two paths must intersect at one point and therefore, due to the above commutations, do not commute with each other. We can use the identity that

\[
e^A e^B = e^{B[A]}
\]

which holds when \([A, B]\) is a number not an operator. This then gives us

\[
W_1 W_2 = e^{iq/\hbar} W_2 W_1 = e^{i\theta} W_2 W_1
\]

where \( \theta \) is the statistical angle of the theory. Thus the Wilson loop operators act just like operators \( T_1 \) and \( T_2 \) in section 4.3 which created particle-hole pairs and moved them around a nontrivial cycle then reannihilated. So even without discussing particles, the ground state wavefunction of the Chern-Simons theory is degenerate!

**Chapter Summary**

- The Charge-Flux model can be realized in an abelian Chern-Simons theory.
- We introduced some ideas of general nonabelian Chern-Simons theory, including manifold invariants and turning Wilson loop operators into knot invariants.

A good reference for abelian Chern-Simons theory is


Some good references on nonabelian Chern-Simons theory are

- E. Witten, *Quantum Field Theory and the Jones Polynomial* Comm. Math. Phys. Volume 121, Number 3 (1989), 351-399; available online here https://projecteuclid.org/euclid.cmp/1104178138. This is the paper that won a Fields’ medal!
Exercises

Exercise 5.1 Polyakov Representation of the Linking Number
Consider a link made of two strands, \( L_1 \) and \( L_2 \). Consider the double line integral

\[
\Phi(L_1, L_2) = \frac{\epsilon_{ijk}}{4\pi} \oint_{L_1} dx^i \oint_{L_2} dx^j \frac{x^k - y^k}{|x - y|^3}
\]

(a) Show that \( \Phi \) is equal to the phase accumulated by letting a unit of flux run along one strand, and moving a unit charged particle along the path of the other strand.

(b) Show that the resulting phase is the topological invariant known as the linking number — the number of times one strand wraps around the other, see section 2.6.2.

This integral representation of linking was known to Gauss.

Exercise 5.2 Gauge Transforming the Chern-Simons Action
Make the gauge transform Eq. 5.13 on the Chern-Simons action 5.9 and show that it results in the change 5.15. Note that there will be an additional term that shows up which is a total derivative and will therefore vanish when integrated over the whole manifold \( \mathcal{M} \).

Exercise 5.3 Winding Numbers of Groups in Manifolds
Consider the mapping of \( U(x) \in SU(2) \to S^3 \). Construct an example of a map with winding number \( n \) for arbitrary \( n \). I.e., find a representative of each group element of \( \Pi_3(SU(2)) \) (See note 17).

Exercise 5.4 Quantization of Winding Number
Let us consider the manifold \( S^3 \) which we consider as \( \mathbb{R}^3 \) plus a point at infinity. Consider the gauge transform function defined

\[
U(x) = \exp \left( \frac{i\pi N x \cdot \sigma}{\sqrt{|x|^2 + R^2}} \right)
\]

where \( x \) is a point in \( \mathbb{R}^3 \), and \( \sigma \) represents the Pauli matrices with \( R \) an arbitrary length scale. Show the winding number Eq. 5.16 gives the integer \( N \). Why does \( N \) need to be an integer here?
6.0.1 Why This Is Hard

Little is known about quantum gravity with any certainty at all. What we do know for sure is the value of some of the fundamental constants that must come into play: the gravitational constant $G$, the speed of light $c$ and of course Planck’s constant $\hbar$. From these we can put together an energy scale, known as the Planck Scale

$$E_{\text{Planck}} = \sqrt{\frac{\hbar c^5}{G}} \approx 10^{28} \text{ eV.}$$

The temperature of the world around us is about 0.03 eV. Chemistry, visible light, and biology occur on the scale of 1 eV. The LHC accelerator probes physics on the scale of roughly $10^{13}$ eV. This means trying to guess anything about the Planck scale is trying to guess physics on an energy scale 15 orders of magnitude beyond what any accelerator experiment has ever probed. We must surely accept the possibility that any physical principle we hold dear from all of our experiments on low energy scales could no longer hold true at the Planck scale! The only thing that is really required is that the effective low energy theory matches that which we can see at the low energies in the world around us.

6.0.2 Which Approach?

There are several approaches to quantum gravity. While I will not make any statement about which approaches are promising, and which approaches are crazy and overpublicized, I am comfortable stating that many of these investigations have led to incredibly interesting and important things being discovered. While in some cases (maybe in most cases) the discoveries may be more about math than about physics, they are nonetheless worthwhile investigations that I am enthusiastic about.

6.1 Some General Principles?

We have to choose general principles that we believe will always hold, despite the fact that we are considering scales of energy and length 15 orders of magnitude away from anything we have ever observed or measured. Much of the community feels that the most fundamental...
thing to hold onto is the Feynman picture of quantum mechanics — that all space-time histories must be allowed. We might write a quantum partition function of the form

\[ Z = \sum_{\text{All universes}} e^{iS/\hbar} \]  

where the sum is now over everything that could happen in all possible histories of the universe — it is the ultimate sum over histories! Obviously such a thing is hard to even contemplate. Several key simplifications will make contemplation easier:

1. Let us ignore matter. Let us (at least to begin with) try to model only universes which are completely devoid of substance and only contain vacuum.

Thus the universe contains only the space-time metric. Doing this, the Einstein-Hilbert action for gravity takes the form

\[ S_{\text{Einstein}} \sim \int_{\mathcal{M}} dx \sqrt{-\det(g)} \]

where the integration is over the entire space-time manifold \(\mathcal{M}\), where \(g\) is the space-time metric tensor and \(R\) is the Ricci scalar. One might imagine that we could construct a theory of quantum gravity by plugging the Einstein-Hilbert action into the path integral form of Eq. 6.1. We obtain

\[ Z = \int Dg(x) e^{iS_{\text{Einstein}}[g(x)]/\hbar}, \]  

thus summing (or integrating) over all possible space-time metrics. Even without matter in the universe, the model is very nontrivial because the space-time metric can fluctuate — these fluctuations are just gravity waves. Even in this limit no one has fully made sense of this type of path integral without many additional assumptions.

2. Let us simplify even more by considering a 2+1 dimensional universe.

We are used to the idea that many things simplify when we go to lower dimension. Indeed, that is what happens here. In 2+1 dimension, there is an enormous simplification that there are no gravity waves! Why not? In short, there are just not enough degrees of freedom in a 2+1 dimensional metric to allow for gravity waves. (For more information about this fact see the appendix to this chapter, section 6.2.) As a result, the only classical solution of the Einstein equations in the vacuum is that \(R = 0\) and that is all! I.e., the universe is flat and there are no fluctuations. (One can also have a cosmological constant \(\Lambda\) in which case \(R = 2\Lambda g\) is the solution).

One might think that this means that gravity in 2+1D is completely trivial. However, it is not. The space-time manifold, although every-

---

4Written down first by Hilbert in 1915.

5If you are rusty with your general relativity, recall that the metric tensor \(g\) defines the relativistically invariant line element via \(ds^2 = g_{\mu\nu}dx^\mu dx^\nu\), and the Ricci scalar \(R\), which is a complicated function of \(g\), is a measure of the curvature of a manifold which compares the volume a small ball to the volume it would have in flat Euclidean space. In particular for a \(D\) dimensional manifold \(\mathcal{M}\) we would consider a \(D\)-dimensional ball \(B^D\) of radius \(\epsilon\) and we have

\[ \frac{V(B^D) \subset \mathcal{M}}{V(B^D) \subset \mathbb{R}^D} = 1 - \frac{\epsilon^2 R}{6(D + 2)} + \ldots \]

6Observation of gravity waves by the LIGO experiment won the 2017 Nobel Prize. Long before this we had very strong indirect observation of gravity waves from observation of the Hulse-Taylor binary pulsar which earned a Nobel Prize in 1993.
where curvature free, still has the possibility of having a nontrivial topology. Thus what we are interested in is actually the different topologies that our space-time manifold might have!

We thus rewrite Eq. 6.1 as

$$Z = \sum_{\text{manifolds } M} \int_M Dg(x) \ e^{iS[g(x)]/\hbar} = \sum_{\text{manifolds } M} Z(M)$$

where $S[g(x)]$ is the Einstein-Hilbert action for a flat universe with metric $g$, the sum is over all different topologies of manifolds the universe might have, and the integration $Dg$ is an integration over all metrics subject to the condition that the manifold’s topology is fixed to be $M$.

Why would we be interested in such a quantity? In short, suppose we know what the topology is of our ($d$-dimensional universe) at a fixed time $t$. We want to know the amplitudes that the topology changes as $t$ develops. I.e., is the space-time manifold of our universe of the form $M = \Sigma \times \text{time}$ or does the space-time manifold split analogous to that shown in Fig. 6.1.

Here is the surprise: the function $Z(M)$ is precisely the Chern-Simons partition function discussed above in section 5.2 for an appropriately chosen gauge group! This connection is very roughly sketched in the chapter appendix section 6.3.

6.1.1 Further Comments on Connections to Quantum Gravity

In the “this is not string-theory” school of thought for quantum gravity, evaluation of Eq. 6.2 is the main goal. Crucially one needs some variables to describe the metric of the universe. Several different approaches to this seem to converge on some similar structures. One interesting approach, known as loop quantum gravity, uses Wilson loop operators as the elementary variables of the theory (once one has reformulated gravity to look like a gauge theory). Another approach discretizes space-time and sums over the different possible discretizations. With certain assumptions these approaches appear to be very closely related! In section 21.3 we will return to the issue of discretizing space-time and how this can result in topological gravity.

---

7 This was first noted by Achúcarro and Townsend [1986] and then was developed further by Witten [1988] and many others.

8 Indeed at length scales as small as the Planck length $l_{\text{Planck}} = \sqrt{\hbar G/c^3} = h\epsilon/E_{\text{Planck}} \approx 1.6 \times 10^{-35}$ m, there is no reason to believe space-time resembles our macroscopic idea of a smooth manifold. The ratio of the radius of the sun to the radius of an atom is roughly the same as the ratio of the radius of an atom to the Planck length!
6.2 Appendix: No Gravity Waves in 2+1 D

Why are there no gravity waves in 2+1 dimension? The short argument for this is as follows (taken from Carlip [2005]):

In $n$ dimensions, the phase space of general relativity is parametrized by a spatial metric at constant time, which has $n(n-1)/2$ components, and its conjugate momentum, which adds another $n(n-1)/2$ components. But $n$ of the Einstein field equations are constraints rather than dynamical equations, and $n$ more degrees of freedom can be eliminated by coordinate choices. We are thus left with $n(n-1)-2n=n(n-3)$ physical degrees of freedom per spacetime point. In four dimensions, this gives the usual four phase space degrees of freedom, two gravitational wave polarizations and their conjugate momenta. If $n=3$, there are no local degrees of freedom.

Let us put a bit more detail on this argument. If we write the flat metric as $\eta_{\mu,\nu} = \text{diag}[-1,1,1,...]$ in any dimension, and we consider small deviations from a flat universe $g = \eta + h$, we can construct the trace-reversed

$$\bar{h}_{\mu,\nu} = h_{\mu,\nu} - \frac{1}{2} \eta_{\mu,\nu} \eta^{\rho,\sigma} h_{\rho,\sigma}.$$ 

In any dimension, gravitational waves in vacuum take the form

$$\bar{h}_{\mu,\nu,\nu} = 0$$

and

$$\Box \bar{h}_{\mu,\nu} = 0$$

where the comma notation indicates derivatives, and indices are raised and lowered with $\eta$.

In any dimension we will have the gravitational wave of the form

$$\bar{h}_{\mu,\nu} = \epsilon_{\mu,\nu} e^{ik^\rho x_{\rho}}$$

where the polarization $\epsilon_{\mu,\nu}$ is orthogonal to the lightlike propagation wavevector, $k^\mu k_{\mu} = 0$, meaning

$$\epsilon_{\mu,\nu} k^\nu = 0. \quad (6.3)$$

However, one must also worry about gauge freedoms. We can redefine our coordinates and change the form of the metric without changing any of the spatial curvatures. In particular, making a coordinate transform $x \rightarrow x - \xi$, we have

$$\bar{h}_{\mu,\nu} \rightarrow \bar{h}_{\mu,\nu} - \xi_{\rho,\mu} - \xi_{\mu,\nu} + \eta_{\mu,\nu} \xi_{\alpha,\alpha}$$

Now here is the key: In 2+1 D for any matrix $\epsilon$ you choose, you can
always find a

$$\xi_\mu = A_\mu e^{ik^\rho x_\rho}$$

such that

$$\bar{h}_{\mu\nu} = \epsilon_{\mu\nu} e^{ik^\rho x_\rho} = \xi_{\nu,\mu} + \xi_{\mu,\nu} - \eta_{\mu,\nu} \xi_{\alpha,\alpha}$$

This means that the wave is pure gauge, and the system remains perfectly flat! I.e., if you calculate the curvature with this form of $\bar{h}$, you will find zero curvature.

To be more precise, we find

$$\epsilon_{\mu,\nu} = A_\mu k_\nu - A_\nu k_\mu + \eta_{\mu\nu} A^\sigma k_\sigma$$

and any $\epsilon$ that satisfies Eq. 6.3 can be represented with some vector $A$.

It is easy to check this by counting degrees of freedom. $\epsilon$ has 6 degrees of freedom in 2+1 D, but Eq. 6.3 is 3 constraints, and $A$ has three parameters, so we should always be able to solve the equation for $A$ given $\epsilon$.

6.3 Appendix: Relation of 2+1D GR to Chern-Simons Theory (In Brief)

Let us start with a Chern-Simons Lagrangian for $SU(2)_k \otimes SU(2)_{-k}$.

Here we will use a very shorthand notation

$$\mathcal{L} = \frac{k}{4\pi} \int_M (A_+ dA_+ + \frac{2}{3} A_+^3) - \frac{k}{4\pi} \int_M (A_- dA_- + \frac{2}{3} A_-^3)$$

Making the transformation

$$\omega = \frac{1}{2} (A_+ + A_-) \quad e = \frac{k}{8\pi} (A_+ - A_-)$$

one obtains the Lagrangian (using differential form notation)

$$\mathcal{L} = \int (e \wedge R + \frac{\lambda}{3} e \wedge e \wedge e)$$

(6.4)

Here $e$ is interpreted as the dreibein of general relativity which is related to the metric by (returning appropriate indices to vectors)

$$g_{\mu\nu} = e^a_\mu e^a_\nu \eta_{ab}$$

with $\eta_{ab}$ the flat metric in 2+1 D, and $\omega$ is a spin connection which has an equation of motion that dictates it is torsion free, and the remaining Lagrangian Eq. 6.3 is precisely the 2+1D Einstein-Hilbert Lagrangian in the so-called Palatini form. In that equation

$$\lambda = (4\pi/k)^2$$
is the cosmological constant. The calculation here has been given for a Euclidean form of gravity. For Lorenzian gravity one needs to work with $SO(2,1)$ Chern-Simons theory which is a bit more complicated.

More details of the relationship between 2+1D general relativity and Chern-Simons theory are provided in the further reading, listed below.

**Further Reading**

- For a huge amount of information on 2+1 dimensional quantum gravity, see [Carlip 2005](#).
- The relationship of 2+1D gravity to Chern-Simons theory was first developed by Ana Achúcarro and Paul Townsend ([Achúcarro and Townsend, 1986](#))
- The relationship was further developed by Edward Witten ([Witten 1988](#))
- Years later, the question was revisited by Witten [2007](#), where doubt is raised as to whether Chern-Simons theory is sufficient to fully describe gravity in 2+1 dimensions.
- A (potentially biased) history of various approaches to quantum gravity is given by Rovelli [2000](#).
- Reviews of loop quantum gravity are given by Rovelli [2008](#) and Nicolai et al. [2005](#).
- Discussions of discretization approaches to quantum gravity are given by Regge and Williams [2000](#) and Lorente [2006](#).
- The article by Nicolai and Peeters [2007](#) covers the connections between the loop and discretization approach fairly clearly.

Note that none of these references are particularly easy to digest!
Defining Topological Quantum Field Theory

We already have a rough picture of a Topological Quantum Field Theory (TQFT) as a quantum theory that depends on topological properties as opposed to depending on geometric properties. For example, it matters that particle 1 traveled around particle 2, but it doesn’t matter how far apart they are.

We can formalize these ideas by saying that the theory should be independent of small deformations of the space-time metric. We might say that

$$\frac{\delta}{\delta g_{\mu\nu}} \langle \text{any correlator} \rangle = 0.$$  

This is a completely valid way to define a TQFT, but is often not very useful.

Another way to define a (2+1 dimensional) TQFT is that it is a set of rules that takes an input of a labeled link embedded in a three-manifold and gives an output of a complex number in a way that is invariant under smooth deformations. This definition is quite analogous to our definition of a knot invariant, with two key differences. First, we allow for the lines to be labeled with a “particle type” (and our rules for evaluating the end result will depend on the particular particle type labels). Secondly, the link can be embedded in some arbitrarily complicated three-manifold.

This type of mapping (see Fig. 7.1) is precisely the sort of thing that one gets as an output of Chern-Simons theory which we called $Z(\mathcal{M},\text{links})$ as we discussed in section 5.2. The advantage of thinking in this language is that strictly speaking, the functional integrals of Chern-Simons theory

$$\mathcal{M} \rightarrow Z(\mathcal{M},a,b)$$

Fig. 7.1 A (2+1) dimensional TQFT takes an input of a labeled link in a manifold and produces an output of a complex number in a manner which is topologically invariant.

1 Many students find this chapter frighteningly abstract. While this chapter sets the stage for a number of ideas that come later, it can also be skipped to a large extent if it seems too difficult. While it may seem a bit cruel to include such a chapter early in the book, I’ve included it here because it gives the best definition of what a TQFT actually is — which, in one form or another, is what we are studying for the remainder of the book.

2 Particularly condensed matter physicists might start to wonder why we need to start talking about arbitrary, and potentially bizarre sounding, three dimensional manifolds — what could they possibly have to do with real physical systems? However (besides just being a beautiful digression) pursuing this direction allows us to understand some of the strong constraints on topological models and their mathematical structure, and this turns out to be important for the analysis of even fairly simple physical systems.

3 We may also allow world lines of anyons to fuse into other species as discussed in section 4.2.
are often not well defined mathematically. Instead, here we bypass the Chern-Simons field theory altogether and define a TQFT simply as a mapping from a manifold with a link to an output.

A closely related but more formal definition of TQFTs is given by a set of Axioms by Sir Michael Atiyah [Atiyah 1988] which are in some sense much more informative.

7.1 Paraphrasing of Atiyah’s Axioms

Here I’m going to give a rough interpretation of Aityah’s axioms of TQFT, suitable for physicists. To begin with, we will consider space-time manifolds with no particles in them. As we have found above, TQFTs are nontrivial even in the absence of any particles. Later on in section 7.2 we will discuss adding particles and moving them around in space-time too.

We will consider a \( D + 1 \) dimensional space-time manifold\(^5\) which we call \( \mathcal{M} \), and \( D \) dimensional oriented slice \( \Sigma \) — we can often think of this slice as being the \( D \)-dimensional space at a fixed time. Almost always we will be thinking of \( D = 2 \), although the axioms are quite general and can be applied to any \( D \).

**AXIOM 1:** A \( D \)-dimensional space \( \Sigma \) is associated with a Hilbert space \( V(\Sigma) \) which depends only on the topology\(^6\) of \( \Sigma \).

We call the space \( V \), which stands for vector space, although sometimes people call it \( H \) for Hilbert space.

As an example of what we mean, we have seen that if \( \Sigma \) is a torus, there is a nontrivial Hilbert space coming from the ground state degeneracy. This degenerate space is the space \( V(\Sigma) \). The space \( V(\Sigma) \) will depend on the particular anyon theory we are considering. For example in the case of abelian anyons in section 4.3 we found a degeneracy of \( m \) for a system on a torus with statistical angle \( \theta = \pi p/m \).

Note that when we add particles to the system (we will do this in section 7.2), if the particles are nonabelian, then there will also be a Hilbert space associated with the additional degeneracy that comes with such nonabelian particles.

**AXIOM 2:** the disjoint union of two \( D \)-dimensional spaces \( \Sigma_1 \) and \( \Sigma_2 \) will be associated with a Hilbert space which is the tensor product of the Hilbert spaces associated with each space\(^7\). I.e.,

\[
V(\Sigma_1 \cup \Sigma_2) = V(\Sigma_1) \otimes V(\Sigma_2)
\]

In particular this means that the vector space associated with the null or empty space \( \emptyset \) must be just the complex numbers. Let us state this mathematically.

**Axiom 2 Implies:**

\[
V(\emptyset) = C
\]
The reason this must be true is because $\emptyset \cup \Sigma = \Sigma$ and $C \otimes V(\Sigma) = V(\Sigma)$ so the result follows\(^8\).

**AXIOM 3:** If $\mathcal{M}$ is a $(D+1)$-dimensional manifold with $D$-dimensional boundary\(^9\) $\Sigma = \partial \mathcal{M}$, then we associate a particular element of the vector space $V(\Sigma)$ with this manifold. We write
\[
Z(\mathcal{M}) \in V(\partial \mathcal{M})
\]
where the association (i.e., which particular state in the vector space is chosen) again depends only on the topology of $\mathcal{M}$.

Here we might think of $\partial \mathcal{M}$ as being the space-like slice of the system at a fixed time, and $V(\partial \mathcal{M})$ as being the possible Hilbert space of ground states. The rest of $\mathcal{M}$ (the interior, not the boundary) is the space-time history of the system, and $Z(\mathcal{M})$ is the particular wavefunction that is picked out by this given space-time history (See Fig. 7.2).

\[\text{Fig. 7.2 Two depictions of a space-time manifold } \mathcal{M} \text{ with boundary } \partial \mathcal{M}. \text{ The left depiction is problematic because the only boundary of the manifold is supposed to be the red top surface } \partial \mathcal{M} \text{ (the black outline of } \mathcal{M} \text{ really should not be there, but we can't draw a closed three manifold!). The right depiction is more accurate in this sense, although it depicts a 2D } \mathcal{M} \text{ and 1D } \partial \mathcal{M}.\]

The point of this axiom is to state that the particular wavefunction of a system $Z(\mathcal{M})$ which is chosen from the available vector space depends on the space-time history of the system. We have seen this principle before several times. For example, we know that if a particle-antiparticle pair is taken around a nontrivial cycle, this changes which wavefunction we are looking at — this process would be part of the space-time history.

**Axiom 3 Implies:** For $\mathcal{M}$ closed, we have $\partial \mathcal{M} = \emptyset$, the empty space, so
\[
Z(\mathcal{M}) \in C
\]
i.e., the TQFT must assign a manifold a topological invariant which is a complex number. This is exactly what we found from Chern-Simons theory.

\(^8\) If this sounds confusing, remember the space $C$ is just the space of length 1 complex vectors, and tensoring a length $n$ vector with a length $m$ vector gives a size $n \times m$ matrix, so tensoring a vector of length $n$ with a length 1 vector gives back a vector of length $n$.

\(^9\) We use the $\partial$ to denote boundary. See section 6.1.1. Note that if $\Sigma = \partial \mathcal{M}$ then $\partial \Sigma = \emptyset$. 

Fig. 7.3 In this picture $\mathcal{M}$ and $\mathcal{M}'$ are meant to fit together since they have a common boundary but with opposite orientation $\Sigma = \partial \mathcal{M} = \partial \mathcal{M}'$. Here $\langle \psi' | \in V(\Sigma^*)$ lives in the dual space of $| \psi \rangle = Z(\mathcal{M}) \in V(\Sigma)$. Note that the normals are oppositely directed.

AXIOM 4: Reversing Orientation

$$V(\Sigma^*) = V^*(\Sigma)$$

where by $\Sigma^*$ we mean the same surface with reversed orientation, whereas by $V^*$ we mean the dual space — i.e., we turn kets into bras. It is a useful convention to keep in mind that the orientation of the normal of $\partial \mathcal{M}$ should be pointing out of $\mathcal{M}$. See Fig. 7.3.

GLUING: If we have two manifolds $\mathcal{M}$ and $\mathcal{M}'$ which have a common boundary $\partial \mathcal{M} = (\partial \mathcal{M}')^*$ we can glue these two manifolds together by taking inner products of the corresponding states as shown in Fig. 7.4.

Here we have $\Sigma = \partial \mathcal{M} = (\partial \mathcal{M}')^*$ so we can glue together the two manifolds along their common boundary to give \(^{10,11}\)

$$Z(\mathcal{M} \cup_\Sigma \mathcal{M}') = \langle Z(\mathcal{M}') | Z(\mathcal{M}) \rangle$$  \hspace{1cm} (7.1)

COBORDISM: Two manifolds $\Sigma_1$ and $\Sigma_2$ are called “cobordant” if their disjoint union is the boundary of a manifold $\mathcal{M}$.

$$\partial \mathcal{M} = \Sigma_1 \cup \Sigma_2$$

We say that $\mathcal{M}$ is a cobordism between $\Sigma_1$ and $\Sigma_2$. See Fig. 7.5 for
an example. The mathematical structure of cobordisms provides a very powerful abstract method of defining topological quantum field theories, which we describe very briefly in appendix ??.

We thus have $Z(\mathcal{M}) \in V(\Sigma_1^*) \otimes V(\Sigma_2)$, so that we can write

$$Z(\mathcal{M}) = \sum_{\alpha\beta} U^{\alpha\beta} |\psi_{\Sigma_2,\alpha}\rangle \otimes \langle \psi_{\Sigma_1,\beta}|$$

where $|\psi_{\Sigma_2,\alpha}\rangle$ is the basis of states for $V(\Sigma_2)$ and $\langle \psi_{\Sigma_1,\beta}|$ is the basis of states for $V(\Sigma_1^*)$. We can thus think of the cobordism $\mathcal{M}$ as being an evolution\footnote{This evolution may or may not be unitary — indeed, the dimensions of $V(\Sigma_1)$ and $V(\Sigma_2)$ may not even match if $\Sigma_1 \neq \Sigma_2$.} similar to that shown in Fig. 7.5.

\[\Sigma_2\]
\[\mathcal{M}\]
\[\Sigma_1\]
\[\partial \mathcal{M} = \Sigma_1^* \cup \Sigma_2\]

\textbf{Fig. 7.5} $\mathcal{M}$ is the cobordism between $\Sigma_1^*$ and $\Sigma_2$. I.e., $\partial \mathcal{M} = \Sigma_1^* \cup \Sigma_2$. Note that we have reversed orientation of $\Sigma_1$ here.
IDENTITY COBORDISM: If we have $M = \Sigma \times I$ where $I$ is the one dimensional interval (We could call it the 1-disk, $D^1$ also) then the boundaries are $\Sigma$ and $\Sigma^*$ (See Fig. 7.6), and the cobordism implements a map between $V(\Sigma)$ and $V(\Sigma)$. Since the interval can be topologically contracted to nothing (or infinitesimal thickness), we can take this map to be the identity:

$$Z(\Sigma \times I) = \sum_{\alpha} |\psi_{\Sigma,\alpha}\rangle \otimes \langle \psi_{\Sigma,\alpha}| = \text{identity}.$$  

where the sum is over the entire basis of states of $V(\Sigma)$.

Fig. 7.6 A cobordism that can be topologically contracted to nothing acts as the identity on the Hilbert space $V(\Sigma)$.

As we have discussed above in section 4.3, for the torus $T^2$ we have

$$\text{Dim} V(T^2) = \text{number of particle species}$$  

which we argued (at least for modular abelian anyon models) based on non-commutativity of taking anyons around the nontrivial cycles of the torus, and we will justify for nonabelian anyons as well in section 7.2.1.

Similarly, for a 2-sphere $S^2$, we have

$$\text{Dim} V(S^2) = 1$$  

since there are no noncontractible loops, and this will also hold for both abelian and nonabelian theories. See section 4.3.1 for discussion of the ground state degeneracy of abelian theories on higher genus surfaces.

### 7.2 Adding Particles

We now consider extending the ideas of TQFT to space-time manifolds with particle world-lines in them.\textsuperscript{13}

Let us imagine that there are different anyon types which we can label as $a$, $b$, $c$, and so forth. The corresponding antianyons are labeled with overbar $\bar{a}$, $\bar{b}$ and so forth as in section 4.2.2. We now imagine a 2-manifold with some marked and labeled points as shown in Fig. 7.8.

We call the combination of the 2-manifold with the marked points $\Sigma$ for brevity. As with the case without particles (AXIOM 1, in section 7.1), $\Sigma$ is associated with a Hilbert space $V(\Sigma)$. The dimension of this Hilbert space depends on the number and type of particles in the manifold (We expect for nonabelian particles, the dimension will grow exponentially with the number of particles). We can span the space $V(\Sigma)$ with some

Fig. 7.8 A 2-manifold with particles in it, which are marked and labeled points. We now call the combination (the manifold and the marked points) $\Sigma$ for brevity.
basis states \( |\psi_\alpha\rangle \) which will get rotated into each other if we move the marked points around within the manifold (i.e., if we braid the particles around each other).

Similarly a 3-manifold \( \mathcal{M} \) is now supplemented with labeled links indicating the world lines of the particles. The world-lines should be directed unless the particles are their own antiparticles. The world lines are allowed to end on the boundary of the manifold \( \partial \mathcal{M} \). See left of Fig. 7.9. Analogously we may sometimes call the combination of the manifold with its world lines \( \mathcal{M} \), although sometimes we will write this as \( \mathcal{M}; L \) where \( L \) indicates the “link” (or knot) of the world lines.

As in the above discussion of axiom 3, the spacetime history specifies exactly which wavefunction

\[
|\psi\rangle = Z(\mathcal{M}) \in V(\partial \mathcal{M})
\]

is realized on the boundary \( \Sigma = \partial \mathcal{M} \). If a basis of \( V(\partial M) \) is given by wavefunctions \( |\psi_\alpha\rangle \) then we can generally write the particular wavefunction \( |\psi\rangle \) in this basis

\[
|\psi\rangle = \sum_\alpha c_\alpha |\psi_\alpha\rangle.
\]
We can now think about how we would braid particles around each other. To do this we glue another manifold $M'$ to $\partial M$ to continue the time evolution, as shown in the right of Fig. 7.9. The final wavefunction is written as 

$$|\psi'\rangle = Z(M \cup M') \in V(\Sigma')$$

If we put the positions of the particles in $\Sigma'$ at the same positions as the particles in $\Sigma$, then the Hilbert spaces, $V(\Sigma')$ is the same as $V(\Sigma)$, and we can write $|\psi'\rangle$ in the same basis as $|\psi\rangle$

$$|\psi'\rangle = \sum_\alpha c'_\alpha |\psi_\alpha\rangle.$$

We can then think of $Z(M')$ as giving us a unitary transformation on this Hilbert space — which is exactly what we think of as nonabelian statistics. We can write explicitly the unitary transformation

$$Z(M') = \sum_{\alpha\beta} U^{\alpha\beta} |\psi_{\Sigma',\alpha}\rangle \otimes \langle \psi_{\Sigma,\beta}|$$

or equivalently

$$c'_\alpha = \sum_\beta U^{\alpha\beta} c_\beta.$$

Note that if the particles stay fixed in their positions (or move in topologically trivial ways) then $M'$ can be contracted to infinitesimal thickness and we can think of the unitary transformation as being the identity. As with the identity cobordism discussed in section 7.1, we can take such an identity transformation, glue the top to the bottom and obtain

$$Z(\Sigma \times S^1) = \dim [V(\Sigma)] \quad (7.5)$$

I.e., the partition function $Z$ is just the dimension of the Hilbert space of the wavefunction. This holds true even when $\Sigma$ has marked points, or particles, in it.

### 7.2.1 Particles or No-Particles

In the same way that the ground state of a topological system “knows” about the types of anyons that can exist in the system, it is also the case that the TQFT in the absence of particles actually carries the same information as in the presence of particles\textsuperscript{14}. To see this consider a manifold $M$ with labeled and directed world-lines $L_i$ in them, as shown in Fig. 7.10. Now consider removing the world lines along with a hollow tubular neighborhood surrounding the paths that the world-lines follow as shown in the figure. We now have a manifold with a solid torus removed for each world-line loop. (Think of a worm having eaten a path out of the manifold.) In this configuration, the boundary $\partial M$ of the manifold $M$ now contains the surface of these empty tubes — i.e, the surface of a torus $T^2$ for each world-line loop. Note that the empty tube is topologically a solid torus $D^2 \times S^1$ even if the world-line forms some

\textsuperscript{14}Up to here our discussion has been applicable to TQFTs in any dimension. From here on we specialize to the most interesting case of $D = 2$, that is 2+1 dimensions.
7.2 Adding Particles

Fig. 7.10 Removing the world-lines on the left along with a thickened tube. Imagine a worm burrowing along the path of the world lines and leaving a hollow hole (colored red).

Fig. 7.11 The possible wavefunctions $|\psi_a\rangle$ that we can have on the surface of the torus can be realized by having a world-line of a particle of type $a$ going around the handle of the torus. We can call these $Z(\text{solid torus with } a \text{ running around handle}) = |\psi_a\rangle$

The partition function of the manifold with the tori excised from it (the right of Fig. 7.10) contains all of the information necessary to determine the partition function for the left of Fig. 7.10 for any particle types that we choose to follow the given world lines. For the manifold on the right there are two surfaces (the two surfaces on the inside of the solid torus) and $S^1$ again is the circle.
holes left where we excised the two tori), so we have
\[ Z(\mathcal{M}) = \sum_{i,j} Z(\mathcal{M}; i, j) \langle \psi_{L1,i} | \otimes \langle \psi_{L2,j} | \]
where \( Z(\mathcal{M}; i, j) \) is the partition function for the torus with two particle types \( i, j \) following the two world line loops \( L_1 \) and \( L_2 \), and the two wavefunctions are the corresponding boundary condition. Thus, if we want to extract \( Z(\mathcal{M}; a, b) \), where the particle lines are labeled with \( a, b \) we simply glue in the wavefunction \( |\psi_{L1,a}\rangle \otimes |\psi_{L2,b}\rangle \) representing the boundary condition on the two surfaces.

7.3 Building Simple 3-Manifolds

7.3.1 \( S^3 \) and the Modular \( S \)-matrix

We will now consider building up 3-manifolds from pieces by gluing objects together using the gluing axiom from section 7.1. The simplest 3-manifold to assemble is the three sphere \( S^3 \). Remember that \( S^3 \) can be thought of as \( \mathbb{R}^3 \) compactified with a single point at infinity (the same way that \( S^2 \) is a plane, closed up at infinity — think of stereographic projection. See the discussion in section 41.1). Recall also that a solid torus should be thought of as a disk crossed with a circle \( D^2 \times S^1 \). I claim that we can assemble \( S^3 \) from two solid tori\(^\text{16}\)

\[ S^3 = (S^1 \times D^2) \cup_{T^2} (D^2 \times S^1) \]

The notation here is that the two pieces \( S^1 \times D^2 \) and \( D^2 \times S^1 \) are joined together on their common boundary which is \( T^2 \) (the torus surface).

There is a very elegant proof of this decomposition. Consider the 4-ball \( B^4 \). Topologically we have\(^\text{17}\)

\[ B^4 = D^2 \times D^2 \]

Now applying the boundary operator \( \partial \) and using the fact that the boundary operator obeys the Leibniz rule (i.e., it distributes like a derivative), we have
\[ S^3 = \partial B^4 = \partial(D^2 \times D^2) = (\partial D^2 \times D^2) \cup (D^2 \times \partial D^2) = (S^1 \times D^2) \cup_{T^2} (D^2 \times S^1) \]

where we have used the fact that the boundary of a disk is a circle, \( \partial D^2 = S^1 \). Note that the two solid tori differ in that they have the opposite \( D^2 \) filled in. Note that the two solid tori here are glued together along a common \( T^2 = S^1 \times S^1 \) boundary. To see this note that
\[ \partial(S^1 \times D^2) = S^1 \times S^1 = \partial(D^2 \times S^1). \]

The two tori are glued together meridian-to-longitude and longitude-to-
This quantity \( S \) below is an important quantity in topological theories as we shall see in chapter 17 of the torus surface where the two solid tori are glued together.

Let us think about the partition function of these two solid tori which are glued together on their boundaries to make up \( S^3 \). We write the partition function as the overlap between wavefunctions on the outside and inside tori:

\[
Z(S^3) = \langle Z(S^1 \times D^2) | Z(D^2 \times S^1) \rangle = \langle \psi_{\text{outside}} | \psi_{\text{inside}} \rangle
\]

where the \( \psi \)'s are the wavefunctions on the surface of the torus.

We can further consider including world lines around the non-contractible loops of the solid torus, as in Fig. 7.11. There is a different state on the surface of the torus for each particle type we have running around the handle. We then assemble \( S^3 \) with these new solid tori and get an \( S^3 \) with two particle world lines linked together as shown in Fig. 7.13. Gluing the two tori together we get:

\[
Z(S^3; a \text{ loop linking } b \text{ loop}) = \langle Z(S^1 \times D^2; b) | Z(D^2 \times S^1; a) \rangle \equiv S_{ab}
\]

This quantity \( S_{ab} \) is known as the modular \( S \)-matrix, and it is a very important quantity in topological theories as we shall see in chapter 17 below.

\[
\langle Z(S^1 \times D^2; b) | Z(D^2 \times S^1; a) \rangle = Z(S^3, a \text{ link } b) = S_{ab}
\]

Fig. 7.13 Here we assemble a partition function for \( S^3 \) with world lines of a linking \( b \) embedded in the \( S^3 \). To do this we glue together two solid tori each with a world line running around the handle. The green line marked \( b \) runs around the handle of the “outside” torus. The end result is known as the modular \( S \)-matrix, and it gives a basis transform converting between the two bases which both span the Hilbert space of the torus surface where the two solid tori are glued together.

Note that the \( S \)-matrix is unitary, since it is simply a basis transformation between the two sets of wavefunction which both span the vector

\[
\langle \psi_{\text{outside}} | \psi_{\text{inside}} \rangle
\]

meridian. (I.e., the contractible direction of one torus is glued to the non-contractible direction of the other, and vice versa.) A sketch of how the two solid tori are assembled together to make \( S^3 \) is given in Fig. 7.12

Let us think about the partition function of these two solid tori which are glued together on their boundaries to make up \( S^3 \). We write the partition function as the overlap between wavefunctions on the outside and inside tori:

\[
Z(S^3) = \langle Z(S^1 \times D^2) | Z(D^2 \times S^1) \rangle = \langle \psi_{\text{outside}} | \psi_{\text{inside}} \rangle
\]

where the \( \psi \)'s are the wavefunctions on the surface of the torus.

We can further consider including world lines around the noncontractible loops of the solid torus, as in Fig. 7.11. There is a different state on the surface of the torus for each particle type we have running around the handle. We then assemble \( S^3 \) with these new solid tori and get an \( S^3 \) with two particle world lines linked together as shown in Fig. 7.13. Gluing the two tori together we get:

\[
Z(S^3; a \text{ loop linking } b \text{ loop}) = \langle Z(S^1 \times D^2; b) | Z(D^2 \times S^1; a) \rangle \equiv S_{ab}
\]

This quantity \( S_{ab} \) is known as the modular \( S \)-matrix, and it is a very important quantity in topological theories as we shall see in chapter 17 below.

\[
\langle Z(S^1 \times D^2; b) | Z(D^2 \times S^1; a) \rangle = Z(S^3, a \text{ link } b) = S_{ab}
\]

Fig. 7.13 Here we assemble a partition function for \( S^3 \) with world lines of a linking \( b \) embedded in the \( S^3 \). To do this we glue together two solid tori each with a world line running around the handle. The green line marked \( b \) runs around the handle of the “outside” torus. The end result is known as the modular \( S \)-matrix, and it gives a basis transform converting between the two bases which both span the Hilbert space of the torus surface where the two solid tori are glued together.

Note that the \( S \)-matrix is unitary, since it is simply a basis transformation between the two sets of wavefunction which both span the vector

\[
\langle \psi_{\text{outside}} | \psi_{\text{inside}} \rangle
\]

We can further consider including world lines around the noncontractible loops of the solid torus, as in Fig. 7.11. There is a different state on the surface of the torus for each particle type we have running around the handle. We then assemble \( S^3 \) with these new solid tori and get an \( S^3 \) with two particle world lines linked together as shown in Fig. 7.13. Gluing the two tori together we get:

\[
Z(S^3; a \text{ loop linking } b \text{ loop}) = \langle Z(S^1 \times D^2; b) | Z(D^2 \times S^1; a) \rangle \equiv S_{ab}
\]

This quantity \( S_{ab} \) is known as the modular \( S \)-matrix, and it is a very important quantity in topological theories as we shall see in chapter 17 below.

\[
\langle Z(S^1 \times D^2; b) | Z(D^2 \times S^1; a) \rangle = Z(S^3, a \text{ link } b) = S_{ab}
\]

Fig. 7.13 Here we assemble a partition function for \( S^3 \) with world lines of a linking \( b \) embedded in the \( S^3 \). To do this we glue together two solid tori each with a world line running around the handle. The green line marked \( b \) runs around the handle of the “outside” torus. The end result is known as the modular \( S \)-matrix, and it gives a basis transform converting between the two bases which both span the Hilbert space of the torus surface where the two solid tori are glued together.

Note that the \( S \)-matrix is unitary, since it is simply a basis transformation between the two sets of wavefunction which both span the vector
space $V(T^2)$ of the torus surface $T^2$ where the two solid tori are glued


together. Note also that the element $S_{00}$, corresponding to the element


of the $S$-matrix where the vacuum particle (no particle at all!) is put


around both handles. (Here we are using 0 to mean the vacuum.) This
tells us that

$$Z(S^3) = S_{00} \leq 1 \quad (7.7)$$


and in fact, should be strictly less than one unless there are no nontrivial


particle types and $S$ is a one-by-one dimensional matrix.


Another way of viewing the $S$ matrix is as a simple link between two


strands, as shown in Fig. 7.13. As with the Kauffman bracket invariant,
we can construct a set of diagrammatic rules to give a value to knots.
Soon, in chapters 8-16 we will construct diagrammatic rules to help us
“evaluate” knots like this. These rules will be somewhat similar to the
rules for the Kauffman bracket invariant, only now we need to keep track
of labels on world lines as well.


### 7.3.2 $S^2 \times S^1$

Fig. 7.14 Assembling two solid tori to make $S^2 \times S^1$. Here the two contractible
disks $D^2$ are sewed together along their boundaries to make $S^2$.


There is another way we can put two solid tori together to make a closed manifold.21 Instead of attaching longitude-to-meridian and
meridian-to-longitude, we instead attach meridian-to-meridian and long-
itude-to-longitude. (This is perhaps a simpler way to put together two
solid tori!) See Figure 7.14. Here we claim that

$$S^2 \times S^1 = (D^2 \times S^1) \cup_{T^2} (D^2 \times S^1)$$


The sewing together is again done along the common boundary $T^2 = S^1 \times S^1$. The $S^1$ factors in both solid tori are the same, and both of the
$D^2$ have the same $S^1$ boundary. Thus we are sewing together two disks
$D^2$ along their $S^1$ boundaries to make a 2-sphere $S^2$ (imagine cutting
a sphere in half along its equator and getting two disks which are the
north and south hemispheres).

As in the previous case, we can put world lines through the handles of the solid tori if we want. If we do so we have

$$\langle Z(D^2 \times S^1; b) | Z(D^2 \times S^1; a) \rangle = \delta_{ab}$$

---

21In fact there are an infinite number of ways two tori can be sewed together to form a closed manifold. These are discussed in detail in the appendix to this chapter, section 7.4.

22One should be warned that $S^2 \times S^1$ cannot be embedded in usual three dimensional space, so visualizing it is very hard!

23It is worth considering how the world lines, in the case where $a = b$, are positioned in the $S^2 \times S^1$. The world line around the handle of one torus enters each $S^2$ sphere through one hemisphere and the world line around the handle of the other torus exits each $S^2$ sphere through the other hemisphere. This fits with the principle that a nonzero amplitude of two particles on the surface of a sphere can only occur if the two particles are a particle-antiparticle pair. This is discussed in section 8.4.
The reason it is a delta function is that both the bra and ket are really the same wavefunctions (we have not switched longitude to meridian). So except for the conjugation we should expect that we are getting the same basis of states for both tori.

In particular, we have the case where we put no particle (the vacuum) around both handles, we have (i.e., \(a = b = I = 0\))

\[(Z(D^2 \times S^1))Z(D^2 \times S^1)) = \delta_{ab} = 1\]

So we have the result

\[Z(S^2 \times S^1) = 1 \quad (7.8)\]

Note that this agrees with two of our prior statements. On the other hand Eq. 7.5 says that \(Z\) for any two dimensional manifold crossed with \(S^1\) should be the dimension of the Hilbert space for that manifold; and on the other hand Eq. 7.4 states that the dimension of the Hilbert space on a sphere is 1.

### 7.3.3 Connected Sums

An important notion in topology is the idea of a connected sum of two manifolds \(M\) and \(M'\), denoted by \(M \# M'\) which is equal to \(M' \# M\). To form the connected sum of two \((D + 1)\)-dimensional closed manifolds, one deletes a small \((D + 1)\)-dimensional ball \(B^{D+1}\) from each of the constituent manifolds, and sews the resulting pieces together along their \(D\)-dimensional sphere boundaries\(^{24}\). Note that a \((D + 1)\)-dimensional sphere acts as the identity for the connected sum

\[S^{D+1} \# M \equiv M \quad (7.9)\]

The reason for Eq. 7.9 is that \(S^{D+1}\) can be assembled from two balls \(B^{D+1}\) attached along their \(S^D\) surfaces\(^{25}\). So the connected sum in Eq. 7.9 is just deleting \(B^{D+1}\) from the manifold \(M\) and replacing it with \(B^{D+1}\) that results from deleting \(B^{D+1}\) from \(S^{D+1}\).

Here will specialize to the case where the manifolds are three dimensional\(^{26}\) and we will derive a simple formula for \(Z(M \# M')\).

First, let us define \(M \setminus B\) to be the manifold \(M\) with a small ball \(B\) removed. Similarly let \(M' \setminus B'\) be the manifold \(M'\) with a small ball \(B'\) removed. Gluing together \(M \setminus B\) with \(M' \setminus B'\) results in

\[M \# M' = (M \setminus B) \cup_{\partial(M \setminus B)} (M' \setminus B')\]

Invoking Eq. 7.1 thus gives us

\[Z(M \# M') = (Z(M \setminus B) \mid Z(M' \setminus B')) \quad (7.10)\]

Now we can reconstruct \(Z\) for each manifold by gluing the pieces back together

\[Z(M) = (Z(M \setminus B) \mid Z(B)) \quad (7.11)\]

For example, the connected sum of two two-dimensional torus surfaces is a two-handled torus surface:

\[T^2 \# T^2 = \text{two handle torus}\]

For example, the usual two-dimensional sphere \(S^2\) sphere can be assembled from two \(B^2\) discs (also known as \(D^2\) disks) sewed together along their \(S^1\) circle boundaries.

In three dimensions there is a notion of a prime decomposition of a manifold. A three-dimensional manifold \(M\) is prime if \(M = N \# P\) implies either \(N = S^3\) or \(P = S^3\). Any orientable compact three-manifold has a decomposition into primes

\[M = P_1 \# P_2 \# \ldots \# P_N\]

which is unique up to reordering the terms and connected sums with \(S^3\) (See Milnor [1962]). In \(D = 4\) such decompositions are not unique. In \(D = 2\) decompositions of orientable manifolds are unique and the only prime manifold is the torus.
\[ Z(\mathcal{M}') = \langle Z(B') \mid Z(\mathcal{M}' \setminus B') \rangle \]  
(7.12)

Now it is crucial that the boundaries of these two manifolds \( \mathcal{M} \setminus B \) and \( \mathcal{M}' \setminus B' \) are spheres \( S^2 \) and the Hilbert space on this sphere is one dimensional, as pointed out in Eq. 7.4. For states \( |a\rangle, |b\rangle, |c\rangle, |d\rangle \) in a one dimensional Hilbert space we have the relationship\(^\text{27}\)

\[ \langle a|b\rangle\langle c|d\rangle = \langle a|d\rangle\langle c|b\rangle \]

So we have

\[ \langle Z(\mathcal{M}\setminus B) \mid Z(\mathcal{M}' \setminus B') \rangle \langle Z(B') \mid Z(\mathcal{M} \setminus B) \rangle \]
\[ = \langle Z(\mathcal{M}\setminus B) \mid Z(B) \rangle \langle Z(B') \mid Z(\mathcal{M}' \setminus B') \rangle \]

Finally we realize that when we sew together two three-dimensional balls \( B \) and \( B' \) along their \( S^2 \) surfaces we obtain the manifold \( S^3 \) (see comment below Eq. 7.9), so that

\[ \langle Z(B') \mid Z(B) \rangle = Z(S^3) \]  
(7.14)

Using Eqs. 7.10, 7.11, 7.12 and 7.14 in Eq. 7.13 we obtain our final result

\[ Z(\mathcal{M} \# \mathcal{M}') = \frac{Z(\mathcal{M})Z(\mathcal{M}')}{Z(S^3)} \]  
(7.15)

We can check that this equation is compatible with Eq. 7.9 by setting \( \mathcal{M}' = S^3 \) for example.

### 7.4 Appendix: Sewing Two Solid Tori Together

While this discussion is a bit outside the main train of thought (being the development of TQFTs) it is interesting to think about the different ways two solid tori may be sewed together to obtain a closed manifold.

A solid torus is written as \( D^2 \times S^1 \). We define the meridian \( m \) to be the \( S^1 \) boundary of any \( D^2 \). I.e., the meridian is a loop on the surface around the contractible direction of the solid torus. We define the longitude \( l \) as being any loop around the surface of the solid torus which intersects a meridian at one point. This definition unfortunately has some (necessary) ambiguity. A line that loops around the meridian \( n \) times as it goes around the noncontractible direction of the torus, is just as good a definition of a longitude (an example of this is Fig. 7.15, which is \( n = 1 \)).

Let us choose a meridian \( m_1 \) on the surface of one solid torus and choose to sew it to the line \(-qm_2 + pl_2\) of the second solid torus (that is, the line that goes \( p \) times around the longitude and \(-q \) times around
meridian, we make $-q$ negative so that the two tori surfaces are op-posite oriented for attaching them together. Once the two lines are glued together this uniquely defines how the rest of the two torus surfaces are glued together. The resulting object is known as the “Lens space” $L(q, p)$. In section 7.3.1 we showed that $L(0, 1) = S^3$ and in section 7.3.2 we showed that $L(1, 0) = S^2 \times S^1$. Note that due to the ambiguity of definition of the longitude of the torus $-qm_2 + pl_2$, under redefinition of the longitude goes to $(-q - np)m_2 + pl_2$. Thus $L(q + np, p) = L(q, p)$, and in particular, $L(1, 1) = S^3$ also.

### 7.5 Appendix: Cobordisms and Category Theory

While I have promised not to resort to the use of category theory, I will briefly break this promise in order to describe how many mathematicians view topological quantum field theories. The reason I think this is worth mentioning here is so the interested reader has some minimal exposure to a few of the key notions.

First, we should mention roughly what a category is: A category is a mathematical structure which has objects and morphisms. A morphism is a maps (or arrow) between objects. Two categories that we need are (1) the category known as Vect whose objects are vector spaces and whose morphisms are linear maps between vector spaces. (2) the category known as Cob$(D+1)$ whose objects are closed $D$ dimensional manifolds (what we called $\Sigma$) and whose morphisms are $(D+1$-dimensional) cobordisms between these manifolds.

A functor is a mapping (or morphism) between two categories that preserves certain structures. In particular it should map objects to objects and morphisms to morphisms.

Mathematicians often define a topological quantum field theory to be a functor

$$F : \text{Cob}(D+1) \rightarrow \text{Vect}$$

Each manifold (object in Cob) is mapped to a vector space (object in Vect). Disjoint unions of manifolds become tensor products of vector spaces. Cobordisms between manifolds are mapped to linear maps between vector spaces.

### Chapter Summary

- The Atiyah Axioms formalize the idea of a topological quantum field theory.
Further Reading

For discussion on the Atiyah Axioms see Atiyah [1988, 1997].

A discussion of gluing together manifolds (as in sections 7.3-7.4) is given by Rolfsen [1976] or Saveliev [2012]. The book Farb and Margalit [2012] may also be useful. We will discuss this type of sewing further in chapter 22 below.

For further reading on category theory, a classic reference the classic reference is MacLane [1971], which was written long before the idea of topological quantum field theory was around. A beautiful masters thesis by Bartlett [2005] discusses TQFTs from the category perspective.
Part II

Anyon Basics
Fusion and Structure of Hilbert Space

As discussed in section 7.1, each two-dimensional surface (a slice of a three-dimensional space-time manifold) has an associated Hilbert space. In the case where there are particles in this surface, the dimension of the Hilbert space will reflect the nature of the particles. We now seek to understand the structure of this Hilbert space and how it depends on the particles. At the same time we will building up a diagrammatic algebra with the goal of constructing a mapping from world-lines of particles to complex numbers (a definitions of a TQFT as given in Fig. 7.1). We briefly introduced graphical notation in section 4.2.1 and we will continue that development here. For those who prefer more mathematical detail, in section 8.6 (as well as in chapter 12) we will introduce tensor description of diagrams and the associated Hilbert spaces.

8.1 Basics of Particles and Fusion — The Abelian Case

Particle types:

There should be a finite set of labels which we call particle types. For now, let us call them \( a, b, c, \) etc.

Fusion

World lines can merge which we call fusion, or do the reverse, which we call splitting. If an \( a \) particle merges with \( b \) to give \( c \), we write \( a \times b = b \times a = c \). Fusion and splitting are shown diagrammatically in Fig. 8.1. Sometimes colloquially we call both diagrams “fusion”.

It should be noted that we can think of two particles as fusing together even if they are not close together. We need only draw a circle around both particles and think about the “total” particle type inside the circle. For example, we sometimes draw pictures like shown in Fig. 8.2.

In our abelian anyon model of charges and fluxes (see section 4.2), if the statistical angle is \( \theta = \pi p/m \) (\( p \) and \( m \) relatively prime and not both odd) then we have \( m \) species \( a = (aq, a\Phi) \) for \( a = 0 \ldots m - 1 \), where \( q\Phi = \pi p/m \). The fusion rules are simply addition modulo \( m \). That is \( a \times b = (a + b) \mod m \).
It is annoying that we have so many different ways to express the identity, but in different contexts different notations seem natural. For example, if our set of particles is fusing by addition (as we discussed in the charge-flux model) the identity should be 0. But if our group fuses by multiplication, identity is more naturally 1. See note 5 in chapter 41.

Identity

Exactly one of the particles should be called the identity or vacuum. We write this as 1 or 0 or $I$ or $\epsilon$. The identity fuses trivially

$$a \times I = a$$

for any particle type $a$. In the charge-flux model (section 4.2) we should think of the identity as being no charge and no flux. Fusion with the identity is depicted schematically in Fig. 8.3. Often we do not draw the identity particle at all, being that it is equivalent to the absence of any (nontrivial) particle.

$$a \times I = a$$

Fig. 8.3 Fusion of a particle with the identity $a \times I = a$. The dotted line indicates the identity. In some of these pictures the $a$ particle appears to move slightly to the left. However, this is not important for topological properties since the path can be deformed continuously to a particle that does not move.

Antiparticles

Each particle $a$ should have a unique antiparticle which we denote as $\bar{a}$. The antiparticle is defined by $a \times \bar{a} = I$. (There should only be one particle which fuses with any $a$ to give the identity!). A particle going forward in time should be equivalent to an antiparticle going backwards in time as shown in Fig. 8.4. Fusion to the identity can be thought of as a particle turning around in space-time as shown in Fig. 8.5.

A particle may be its own antiparticle, in which case we do not need to draw arrows on its world lines. An example of this in our charge-flux model from section 4.2 would be the “2” particle (fusion of 2 elementary anyons, see section 4.3) in the case of $\theta = \pi/4$. Also, the identity particle $I$ is always its own antiparticle.

8.2 Multiple Fusion Channels - the Nonabelian Case

For the nonabelian theories as we have discussed above (for example in Section 3.5), the dimension of the Hilbert space must increase with the number of particles present. How does this occur? In nonabelian models we have multiple possible orthogonal fusion channels

$$a \times b = c + d + \ldots$$

meaning that $a$ and $b$ can come together to form either $c$ or $d$ or $\ldots$, as shown in Fig. 8.6. A theory is nonabelian if any two particles fuse in
such a way that there are multiple possible fusion channels (i.e., there is more than one particle listed on the right hand side of Eq. 8.1).

If there are $s$ possible fusion channels for $a \times b$, then the two particles $a$ and $b$ have an $s$ dimensional Hilbert space (part of what we called $V(\Sigma)$).

What is this Hilbert space associated with multiple fusion channels? A slightly imperfect analogy is that of angular momentum addition. We know the rule for adding spin $1/2$, 

$$\frac{1}{2} \otimes \frac{1}{2} = 0 \oplus 1,$$

which tells us that two spin $1/2$'s can fuse to form a singlet or a triplet. As with the case of spins, we can think about the two particles being in a wavefunction such that they fuse in one particular fusion channel or the other — even if the two particles are not close together. The singlet or $J = 0$ state of angular momentum is the identity here: it has no spin at all. The analogy with spins is not exact though — unlike the case of spins, the individual particles have no internal degrees of freedom (analogous to the 2-states of the spin $1/2$), nor do any results of fusion have an $m_z$ degree of freedom (like a triplet would).

**Locality**

The principle of locality is an predominant theme of anyon physics (if not of physics altogether).

The quantum number (or “charge”) of a particle is locally conserved in space. Consider, for example, Fig. 8.7. On the left, a particle $a$ propagates along and suddenly something complicated happens locally. If only a single particle comes out of this region it must also be a particle of type $a$. (If two particles come out of this region, we could have $a$ split into two other species as in the right of Fig. 8.1). We sometimes call this the **no transmutation** principle. It allows us to conclude that the complicated picture on the left of Fig. 8.7 must be equal to some constant times the simple propagation of an $a$ particle as shown on the right.

If two particles (maybe far away from each other) fuse together to some overall particle type (in a case where multiple fusion channels are available) it is not possible to determine this fusion channel by measuring only one of the initial particles. In order to determine the fusion channel of the two particles, you have to do an experiment that involves both of the initial particles. For example, one can perform an interference measurement that surrounds both of these particles. The fusion channel is **local** to the pair.

Similarly, if we have some particles, $b$ and $c$ and they fuse to $d$ (see Fig. 8.8), no amount of braiding $b$ around $c$ will change this overall fusion channel $d$. The fusion channel is **local** to the pair. If these two then fuse with $a$ to give an overall fusion channel $f$, no amount of braiding $a$, $b$ and $c$ will change the overall fusion channel $f$. However, if $a$ braids

---

![Fig. 8.7](image1.png) **Fig. 8.7** If a particle $a$ goes into a spacetime region, then a net particle charge $a$ must come out. This is also sometimes called the “no-transmutation” principle. From far away, one can ignore any local processes (up to an overall constant).

![Fig. 8.8](image2.png) **Fig. 8.8** In this picture $b$ and $c$ fuse to $d$. Then this $d$ fuses with $a$ to give an overall fusion channel of $f$. No amount of braiding $b$ around $c$ will change the fact that the two of them fuse to $d$. However, if we braid $a$ with $b$ and $c$, this can change the fusion of $b$ with $c$ subject to the constraint that the fusion of all three particles will give $f$. 
Fusion and Structure of Hilbert Space

with \( b \) and \( c \), then the fusion of \( b \) and \( c \) might change, subject to the constraint that the overall channel of all three particles remains \( f \).

Locality gives another important way in which of anyons differs from the fusion of spins. With spins, if you can measure two spins individually you can (at least sometimes) determine the fusion channel of the spins. For anyons you must be able to measure a loop that surrounds both anyons in order to determine their collective fusion channel — measuring each anyon individually does not tell you the fusion of the two!

**Antiparticles in the Case of Multiple Fusion Channels**

When we have multiple fusion channels (i.e., for nonabelian theories) we define antiparticles via the principle that a particle can fuse with its antiparticle to give the identity, although other fusion channels may be possible.

\[
a \times \bar{a} = I + \text{other fusion channels}
\]

As in the abelian case we use the overbar notation to indicate an antiparticle. It should be the case that for each particle \( a \) there is a unique particle that can fuse with it to give the identity, and we call this particle \( \bar{a} \). As in the abelian case, a particle may be its own antiparticle if \( a \times a = I + \text{other fusion channels} \), in which case we do not put an arrow on the line corresponding to the particle.

**8.2.1 Example: Fibonacci Anyons**

Perhaps the simplest nonabelian example is the anyon system known as Fibonacci\(^2\) anyons. Something very close to this is thought to occur in the so-called \( \nu = 12/5 \) quantum Hall state which we will study in more depth in section 39. Fibonacci anyons are closely related to the \( SU(2)_3 \) Chern-Simons theory\(^3\).

In this example the particle set includes only two particles, the identity \( I \) and a nontrivial particle which is often called \( \tau \).

\[
\text{Particle types} = \{ I, \tau \}
\]

The fusion rules are

\[
\begin{align*}
I \times I &= I \\
I \times \tau &= \tau \\
\tau \times \tau &= I + \tau
\end{align*}
\]

The first two of these rules hardly need to be written down (they are implied by the required properties of the identity). It is the final rule that is nontrivial. This final rule also implies that \( \tau \) is its own antiparticle \( \tau = \bar{\tau} \) which means we do not need to put arrows on world lines.

With two Fibonacci anyons the Hilbert space is two dimensional, since the two particles can fuse to \( I \) or \( \tau \), as shown in Fig. 8.9.

With three Fibonacci anyons the Hilbert space is 3 dimensional, as
shown in Fig. 8.10. The key thing to notice is that if the first two particles fuse to \( \tau \), then this combination acts as being a single particle of overall charge \( \tau \) — it can fuse with the third \( \tau \) in two ways.

There is a single state in the Hilbert space of three anyons with overall fusion channel \( I \). This state is labeled as \(^4|N\rangle\). As mentioned above by Fig. 8.7 due to locality, no amount of braiding amongst the three particles will change this overall fusion channel (although braiding may introduce an overall phase).

There are two states in the Hilbert space of three anyons with overall fusion channel \( \tau \). These are labeled \(|1\rangle\) and \(|0\rangle\) in Fig. 8.10. Again, as mentioned above by Fig. 8.7 due to locality, no amount of braiding amongst the three particles will change this overall fusion channel. Further, since in these two basis states the first two particles furthest left are in an eigenstate (either \( I \) in state \(|0\rangle\) or \( \tau \) in state \(|1\rangle\)) no amount of braiding of the first two particles will change that eigenstate from \(|0\rangle\) to \(|1\rangle\) or from \(|1\rangle\) to \(|0\rangle\). However, as we will see below in section 10.1 if we braid the second particle with the third, we can then change the quantum number of the first two particles and rotate between \(|0\rangle\) and \(|1\rangle\).

\(^4\)Here \(|N\rangle\) stands for “noncomputational”, since it is not used in many quantum computing protocols that use Fibonacci anyons.

\[ \tau \tau \tau \]
\[ I \]
\[ \tau \tau \tau \]
\[ \tau \tau \tau \]
\[ = \]
\[ = \]
\[ = \]

\( |N\rangle \)
\( |0\rangle \)
\( |1\rangle \)

Fig. 8.10  Notations for the three different orthogonal fusion channels of three Fibonacci anyons. The notation \(|N\rangle, |1\rangle\) and \(|0\rangle\) are common notations for those interested in topological quantum computing with Fibonacci anyons!

For our Fibonacci system, with 2 particles the Hilbert space is 2 dimensional. With 3 particles the Hilbert space is 3 dimensional. It is easy to see that with 4 particles the Hilbert space is 5 dimensional (fusing a fourth anyon with \(|0\rangle\) or \(|1\rangle\) in figure 8.10 can give either \( I \) or \( \tau \), whereas fusing a fourth anyon with \(|N\rangle\) can only give \( \tau \), thus giving a space of
Fusion and Structure of Hilbert Space

dimension 2+2+1). With five particles the space is 8 dimensional and so forth. This pattern continues following the Fibonacci sequence (Try to show this!), hence the name.

Since the \( N^{th} \) element of the Fibonacci sequence for large \( N \) is approximately

\[
\text{Dim of } N \text{ Anyons} = \text{Fib}_N \sim \left( \frac{1 + \sqrt{5}}{2} \right)^N
\]

(8.2)

We say that the *quantum dimension* of this particle is \( d = \frac{1 + \sqrt{5}}{2} \), the golden mean (See Eq. 3.8).

### 8.2.2 Example: Ising Anyons

The Ising\(^5\) anyon system is extremely closely related to \( SU(2)_2 \) Chern-Simons theory\(^6\), and this general class of anyon is believed to be realized in the \( \nu = 5/2 \) quantum Hall state (see section 39), topological superconductors, and other so-called Majorana systems (see section ***).

The Ising theory has three particle types\(^7\):

\[
\text{Particle types} = \{ I, \sigma, \psi \}
\]

The nontrivial fusion rules are

\[
\begin{align*}
\psi \times \psi &= I \\
\psi \times \sigma &= \sigma \\
\sigma \times \sigma &= I + \psi
\end{align*}
\]

where we have not written the outcome of any fusion with the identity, since the outcome is obvious. Again, each particle is its own antiparticle \( \psi = \bar{\psi} \) and \( \sigma = \bar{\sigma} \) so we need not put arrows on any world-lines.

Fusion of anything with the \( \psi \) particle always gives a unique result on the right hand side. We thus call \( \psi \) an abelian particle (despite the fact that the full theory is nonabelian), or we say that \( \psi \) is a *simple current* (see margin note 8 below). Fusion of many \( \psi \) particles is therefore fairly trivial, since each pair fuses to the identity in only one way.

Fusion of many \( \sigma \) particles, however, is nontrivial. The first two \( \sigma \)’s can either fuse to \( I \) or \( \psi \), but then when the third is included the overall fusion channel must be \( \sigma \) (since fusing \( \sigma \) with either \( \psi \) or \( I \) gives \( \sigma \)). Then adding a fourth \( \sigma \) to this cluster whose overall quantum number is \( \sigma \) again gives two possible outcomes. Such a fusion tree is shown in Fig 8.11. By counting possible trees, we find that the total number of different fusion channels for \( N \) particles of type \( \sigma \) is \( 2^{N/2} \) (rounding down if \( N/2 \) is not an integer). To see this in another way, we can group \( \sigma \) particles together in pairs where each pair gives \( \psi \) or \( I \), so two \( \sigma \) particles comprises a two state system, or a qubit. Then the \( I \)’s and \( \psi \)’s fuse together in a unique way. Since the Hilbert space dimension is \( (\sqrt{2})^N \) the quantum dimension of the \( \sigma \) particle is \( d = \sqrt{2} \) (See Eq. 3.8).

---

\(^5\)The name Ising is used here due to the relationship with the Ising conformal field theory which describes the Ising model in 2D at its critical point.

\(^6\)The fusion rules of Ising and \( SU(2)_2 \) are the same, but there are some spin factors which differ, as well as a Frobenius-Schur indicator — see sections \( 14.2 \) and \( 18.3 \).

\(^7\)Another common notation is to use \( \epsilon \) instead of \( \psi \) in the Ising theory. In \( SU(2)_2 \) the particles \( I, \sigma, \psi \) may be called 0, 1/2, 1 or 0, 1, 2.

---

![Fig. 8.11](image-url) The fusion tree for many \( \sigma \) particles in the Ising anyon theory.
8.3 Fusion and the $N$ matrices

We are well on our way to fully defining an anyon theory. A theory must have a finite set of particles, including a unique identity $I$, with each particle having a unique antiparticle.

The general fusion rules can be written as

$$a \times b = \sum_{c} N_{ab}^{c} c$$

(8.3)

where the $N_{ab}^{c}$ are nonnegative integers known as the fusion multiplicities. $N_{ab}^{c}$ is zero if $a$ and $b$ cannot fuse to $c$. $N_{ab}^{c}$ is one if we have $a \times b = \ldots + c + \ldots$, and $c$ only occurs once on the right hand side. If $c$ occurs more than once on the right hand side, then $N_{ab}^{c}$ simply counts the number of times it occurs.

A particle $a$ is a simple current if $\sum_{c} N_{ab}^{c} = 1$ for each particle $b$.

What does it mean that a particle type can occur more than once in the list of fusion outcomes? It simply means that the fusion result can occur in multiple orthogonal ways in which a diagram with a vertex showing $a$ and $b$ fusing to $c$ should also contain an index ($\mu \in 1 \ldots N_{ab}^{c}$) at the vertex indicating which of the possible $c$ fusion channels occurs, as shown in Fig. 8.12. For most simple anyon theories $N_{ab}^{c}$ is either 0 or 1, and we will not usually consider the more complicated case in examples for simplicity, but they are discussed in the chapter appendices for completeness (See section 9.5.3. See also section ***). It is good to keep in mind that such more complicated cases exist.

Elementary properties of the fusion multiplicity matrices

- Commutativity of fusion $a \times b = b \times a$.

$$N_{ab}^{c} = N_{ba}^{c}$$

(8.4)

- Time reversal

$$N_{ab}^{c} = N_{ba}^{c}$$

(8.5)

- Trivial fusion with the identity

$$N_{aI}^{b} = \delta_{ab}$$

(8.6)

- Uniqueness of inverse

$$N_{ab}^{I} = \delta_{ba}$$

(8.7)

It is sometimes convenient to define

$$N_{abc} = N_{ab}^{c}$$

(8.8)

which is the number of different ways that $a$, $b$, and $c$ can fuse to the identity. An example of this equivalence is shown graphically in Fig. 8.13. The advantage of this representation is that $N_{abc}$ is fully symmetric in all of its indices. For example, using this notation Eq. 8.5 and Eq. 8.7 are actually the same. Further, using Eq. 8.8 along with the symmetry

8A particle $a$ is a simple current if $\sum_{c} N_{ab}^{c} = 1$ for each particle $b$.

9While this does not occur for angular momentum addition of $SU(2)$ (and also will not occur in Chern-Simons theory $SU(2)_k$ correspondingly) it is well known among high energy theorists who consider the fusion of representations of $SU(3)$. Recall that $8 \otimes 8 = 1 \oplus 8 \oplus 10 \oplus \bar{10} \oplus 27$ and the 8 occurs twice on the right.

Fig. 8.12 Multiple fusion channels. In nonabelian theory fusion of $a$ and $b$ to $c$ can occur in multiple orthogonal ways when $N_{ab}^{c} > 1$. To specify which way they fuse, we add an additional index $\mu \in 1 \ldots N_{ab}^{c}$ at the vertex as shown.

Fig. 8.13 An equivalence of $N_{ab}^{c}$ with $N_{abc}$. Both types of vertices have the equivalent fusion multiplicity. Note that the left half of the right picture is exactly equivalent to the right — $c$ is entering the vertex from below (then this $c$ turns over to become a $\bar{c}$ going up on the far right).
of $N_{abc}$ we can derive identities such as

$$N_{ab} = N_{ac} = N_{bc}.$$  

where in the last step we used Eq. (8.5).

A set of particle types with a unique identity and a set of $N_{ab}$’s satisfying the axioms listed here is known as a commutative fusion ring. All TQFTs are described by commutative fusion rings. However, commutative fusion rings exist that are not consistent with any TQFT.

Fusing Multiple Anyons

If we are to fuse, say, five anyons of type $a$ together into a final result of $e$, we can do so via a tree as shown in Fig. 8.14.

![Fig. 8.14 Fusing five $a$ type anyons together into a final result $e$.](image)

It is convenient to think of the tensor $N_{ab}$ as a matrix $N_a$ with indices $b$ and $c$, i.e., we write $N_a^c$, such that we have

$$\text{Dim of fusing five } a \text{ anyons to final result } e = \left[ \left( N_a^c \right) \right]_e$$

Similarly were we to have a larger number $p$ of anyons of type $a$ we would need to calculate $\left[ N_a^c \right]_e^{p-1}$. We recall (See Eq. (3.8)) that the quantum dimension $d_a$ of the anyon $a$ is defined via the fact that the Hilbert space dimension should scale as $d_a N$, where $N$ is the number of $a$ particles fused together. We thus have that

$$d_a = \text{largest eigenvalue of } [N_a]$$

Note that this implies $d_a = d_a$ given the symmetries of $N$.

Example of Fibonacci Anyons

The fusion matrix for the $\tau$ particle in the Fibonacci theory is

$$N_\tau = \begin{pmatrix} 1 & \tau \\ 0 & 1 \end{pmatrix} I$$

where, as indicated here, the first row and first column represent the identity and the second row and second column represent $\tau$. The first row of this matrix says that fusing $\tau$ with the identity gives back $\tau$ and the second row says that fusing $\tau$ with $\tau$ gives $I$ and $\tau$. It is an easy exercise to check that the largest eigenvalue of this matrix is indeed $d_a = \text{largest eigenvalue of } [N_a]$.
\[ d = \frac{(1 + \sqrt{5})}{2}, \text{ in agreement with Eq.}\; 8.2 \]

**Example of Ising Anyons**

The fusion matrix for the \( \sigma \) particle in the Fibonacci theory is

\[
N_{\sigma} = \begin{pmatrix}
I & \sigma & \psi \\
0 & 1 & 0 \\
1 & 0 & 1 \\
0 & 1 & 0
\end{pmatrix}
\]

where the first row and column represent the identity, the second row and column represent \( \sigma \) and the third row and column represent \( \psi \). So, for example, the second row here indicates that \( \sigma \times \sigma = I + \psi \). Again, it is an easy exercise to check that the largest eigenvalue of this matrix is \( d = \sqrt{2} \) as described in section 8.2.2.

### 8.3.1 Associativity

It should be noted that the fusion multiplicity matrices \( N \) are very special matrices since the outcome of a fusion should not depend on the order of fusion. I.e., \( (a \times b) \times c = a \times (b \times c) \).

For example, let us try to calculate how many ways \( a \times b \times c \) can give an outcome of \( e \). We can either try fusing \( a \times b \) first as on the left of Fig. 8.15 or we can try fusing \( b \) and \( c \) first as on the right. Whichever we choose, we are describing the same Hilbert space and we should find the same overall dimension either way. In other words, we should have the same total number of fusion channels. Thus, corresponding to these two possibilities we have the equality

\[
\sum_d N_{ab}^d N_{cd}^e = \sum_f N_{cb}^f N_{af}^e
\]

Again, thinking of \( N_{ab}^e \) as a matrix labeled \( N_a \) with indices \( b \) and \( c \), this tells us that

\[
[N_a, N_c] = 0 \quad (8.12)
\]

Therefore all of the \( N \) matrices commute with each other. In addition the \( N \)'s are normal matrices, meaning that they commute with their own transpose (Since \( [N_a, N_a] = 0 \) and \( N_a = N_a^T \) by Eq. 8.9). A set of normal matrices that all commute can be simultaneously diagonalized, thus

\[
[U^\dagger N_a U]_{xy} = \delta_{xy} \lambda_x^{(a)}
\]

and all \( N_a \)'s get diagonalized with the same unitary matrix \( U \). Surprisingly (as we will see below in section 17.3.1) for well behaved (so-called “modular”12 anyon theories) the matrix \( U \) is precisely the modular S-matrix we discussed above in Eq. 7.6!

---

12For nonmodular theories, or even fusion rings which don’t even correspond to anyon theories of any type, we can still diagonalize \( N \) in the form of Eq. 8.13 and the resulting unitary matrix \( U \) is sometimes known as the mock S-matrix.
8.4 Application of Fusion: Dimension of Hilbert Space on 2-Manifolds

The structure of fusion rules can be used to calculate the ground state degeneracy of wavefunctions on 2-dimensional manifolds. Here we will again be examining the Hilbert space $V(\Sigma)$ where $\Sigma$ is our 2-manifold which may or may not have particles in it.

Let us start by considering the Hilbert space $V(S^2)$ where $S^2$ is the sphere. As mentioned previously in Eq. 7.5, there is a unique ground state in this situation because there are no non-contractible loops (See sections 4.1 and 4.3.1). The dimension of the Hilbert space is just 1,

$$\text{Dim } V(S^2) = 1.$$

This will be the starting point for our understanding. All other configurations (change of topology, adding particles etc) will be related back to this reference configuration.

Now let us consider the possibility of having a single (nontrivial) anyon on the sphere. In fact such a thing is not possible because you can only create particles in a way that conserves that overall quantum number. If we start with no particles on the sphere, the total anyon charge must be conserved — i.e., everything on the sphere must fuse together to total quantum number of the identity. Thus, we have

$$\text{Dim } V(S^2 \text{ with one (nontrivial) anyon}) = 0 \quad (8.14)$$

Another way to explain this is to realize that, since particle-antiparticles are made in pairs, there is no space-time history that could prepare the state with just a single (non-vacuum) particle on the sphere.

We can however consider the possibility of two anyons on a sphere. We can create an $a$ particle with an $\bar{a}$ particle, and since these two particles must fuse back to the identity in a unique way we have

$$\text{Dim } V(S^2 \text{ with one } a \text{ and one } \bar{a}) = 1$$

The two particles must be antiparticles of each other, otherwise no state is allowed and the dimension of the Hilbert space is zero. This is a general principle: the fusion of all the particles on the sphere must be the vacuum, since these particles must have (at some point in history) been pulled from the vacuum.

Now we could also imagine puncturing the sphere to make a hole where the particles were. In the spirit of what we did in section 7.2.1 we could re-fill the hole with any particle type. However, if we refill one hole with a particular particle type $a$, then the other hole can only get filled in with the anti-particle type $\bar{a}$. Nonetheless, we can conclude that

$$\text{Dim } V(S^2 \text{ with two unlabeled punctures}) = \text{Number of particle types}$$
8.4 Application of Fusion: Dimension of Hilbert Space on 2-Manifolds

Now consider the procedure shown in Fig. 8.16. We start with the twice punctured sphere. The two punctures can be labeled with any particle-antiparticle pair labels. We can then deform the sphere to sew the two punctures together in a procedure that is sometimes called surgery (We will discuss surgery in more detail in chapter 22). The result of this surgery is the torus surface $T^2$ and we conclude that

$$\dim V(T^2) = \text{Number of particle types}$$

as we have already discussed. The general rule of surgery is that two punctures can be sewed together when they have opposing particle types (i.e., a particle and its antiparticle). This is exactly the gluing property of the TQFT. Although we are gluing together pieces along a 1-dimensional boundary (the edge of the punctures), we should realize that there is also a time direction, which we have implicitly assumed is compactified into $S^1_{\text{time}}$. Thus we are actually sewing together the 2-surface $(S^1_{\text{puncture}} \times S^1_{\text{time}})$ with another 2-surface $(S^1_{\text{puncture}} \times S^1_{\text{time}})$, and the inner product between the two wavefunctions on these two-surfaces ensures that the quantum number on these two punctures are conjugate to each other$^{18}$. $^{18}$In Eq. 7.3 we had a torus surface which we crossed with an interval of time and we closed up the interval to form a circle, thus giving $\text{Tr}[Z(T^2 \times I_{\text{time}})] = Z(T^2 \times S^1_{\text{time}}) = \dim V(T^2)$. In contrast, in Fig. 8.16 we have a cylinder $S^1 \times I$ (topologically the same as a sphere with two holes) crossed with $S^1_{\text{time}}$ and we close the cylinder to get $\text{Tr}[Z((S^1 \times I) \times S^1_{\text{time}})] = Z(T^2 \times S^1_{\text{time}})$.

We can continue on to consider a sphere with three particles. Similarly we should expect that the three particle should fuse to the identity as shown in Fig. 8.17. We can then think of the sphere with three particles as being a sphere with three labeled punctures which is known as a “pair of pants”, for reasons that are obvious in Figure 8.18. It turns out that any orientable 2-dimensional manifold (except $S^2$ or $T^2$ which we have already considered) can be constructed by sewing together the punctures of pants — this is known as a “pants decomposition”. For example, in Fig. 8.19 we sew together two pair of pants to obtain a two handled torus.

In Fig. 8.18 we had a torus surface which we crossed with an interval of time and we closed up the interval to form a circle, thus giving $\text{Tr}[Z(T^2 \times I_{\text{time}})] = Z(T^2 \times S^1_{\text{time}}) = \dim V(T^2)$.
To find the ground state degeneracy of the two handled torus, we assemble the manifold using two pair of pants as shown in Fig. 8.19 and then we simply need to figure out the number of possible fusion channels where we could satisfy \( a \times b \times c \rightarrow I \) (for the bottom pair of pants) and \( \bar{a} \times \bar{b} \times \bar{c} \rightarrow I \) (for the top pair of pants). This number of possible fusion channels is given in terms of the fusion multiplicities \( N_{abc} \) as shown in Fig. 8.17. Essentially we are just looking at the number of ways we can assign labels to the punctures when we glue the objects together. Thus we have

\[
\dim V(\text{Two handled torus}) = \sum_{abc} N_{abc} N_{\bar{a}\bar{b}\bar{c}}
\]

Fig. 8.19 Sewing together two pair of pants to form a two-handled torus.

Another interesting use of the pants diagram is to determine the degeneracy of a torus \( T^2 \) with a single anyon on it labelled \( a \). Unlike the sphere, where one cannot have a single anyon on the surface (see Eq. 8.14) one can have a single anyon on a torus (see note 15 of this chapter). To see how this is possible, take a pants diagram with the holes labelled \( b, \bar{b}, \) and \( a \). Connect up the \( b \) to the \( \bar{b} \) to give a torus with a single puncture remaining labeled \( a \). Thus we conclude that

\[
\dim V(T^2 \text{ with one } a) = \sum_{b} N_{\bar{b}ba} \equiv L_a
\]

where we have defined this quantity to be called \( L_a \).

One final example is to determine the ground state degeneracy of a three handled torus. There are many ways we might cut a three handled torus into pieces, but a convenient decomposition is the one shown in
Here there are three tori each with a puncture in it (marked as a red collar), and a single pants in the middle connecting the three. Each torus with a puncture has a Hilbert space dimension $L_a$ where $a$ is the quantum number assigned to the puncture. Thus the total dimension of the Hilbert space is conveniently written as

$$\text{Dim } V(\text{Three handled torus}) = \sum_{abc} L_a L_b L_c N_{abc} \tag{8.16}$$

**Example: Fibonacci Anyons**

With the Fibonacci fusion rules, there are five ways we can fuse three particles and get the identity.

$$N_{III} = 1$$
$$N_{\tau\tau\tau} = N_{\tau I\tau} = N_{I\tau\tau} = 1$$
$$N_{\tau\tau\tau} = 1$$

and all other $N_{abc} = 0$. Thus there are five possible labelings of the punctures in a pants diagram that allow overall fusion to the identity. If we match these together on both top and bottom of the diagram on the left of Fig. 8.19 we conclude that in the Fibonacci theory we have

$$Z(\text{Two handled torus } \times S^1) = \text{Dim } V(\text{Two handled torus}) = 5.$$

Similarly, we can consider the degeneracy of states for a torus with a single $\tau$ particle on its surface

$$\text{Dim } V(T^2 \text{ with one } \tau \text{ particle on it}) = 1$$

coming from the allowed fusion $N_{\tau\tau\tau} = 1$. Thus we have $L_I = 2$ and $L_\tau = 1$. It is then easy to plug into Eq. 8.16 to obtain

$$\text{Dim } V(\text{Three handled torus}) = 15.$$

### 8.5 Product Theories

A very common construction is to consider the product of two anyon theories. Given two anyon theories (let us call them $T$ and $t$) with particle types

$$a, b, c, \ldots \in t$$
$$A, B, C, \ldots \in T$$

we consider the product theory $T \times t$. The Hilbert space of the product theory is just the product of the Hilbert spaces of the constituent theories. So any arbitrary particle $\alpha$ in the product theory is composed of
one particle from each of the constituent theories

\[ \alpha \in T \times t \implies \alpha = (Y, x) \text{ with } Y \in T \text{ and } x \in t \]

Roughly we can think of this as putting both theories in the same place at the same time — particles from \( T \) can exist (combined with \( I \) from \( t \)) and particles from \( t \) can exist (combined with \( I \) from \( T \)) and any combination of particles from both theories can also exist.

For example, in the theory \((\text{Ising} \times \text{Fibonacci})\), there are 6 particle types which we can label as

\((I, I)\) \((I, \tau)\) \((\sigma, I)\) \((\sigma, \tau)\) \((\psi, I)\) \((\psi, \tau)\)

The fusion multiplicity matrices \( N \) for the product theories are just the product of the \( N \) matrices for the constituent theories

\[ N_{(A,a),(B,b)}^{C,c} = N_{A,B}^{C} N_{a,b}^{c} \]

### 8.6 Appendix: Tensor Description of Fusion and Splitting Spaces

Let us now try to give a bit more precise mathematical meaning to idea of fusion as well as to some of the diagrams we have been drawing\(^\text{19}\). For each fusion \( N_{ab}^{c} \) we define a space \( V_{ab}^{c} \) known as a fusion space and a space \( V_{ab}^{c} \) known as a splitting space. Both of these spaces have dimension

\[ \dim V_{ab}^{c} = \dim V_{ab}^{c} = N_{ab}^{c} \]

Each of these spaces can be given an orthonormal basis, which we label with an index \( \mu \). We can write states in this space as kets which we draw as diagrams. For example

\[
\begin{array}{c}
\mu \\
\downarrow \\
\begin{array}{c}
\ a \\
\c \\
\ b
\end{array}
\end{array}
\]

\[ = |a, b; c, \mu \rangle \in V_{ab}^{c} \quad (8.17) \]

describes the splitting space. The Hermitian conjugate, the corresponding bras, are drawn as fusion diagrams

\[
\begin{array}{c}
\mu \\
\downarrow \\
\begin{array}{c}
\ a \\
\c \\
\ b
\end{array}
\end{array}
\]

\[ = \langle a, b; c, \mu | \in V_{ab}^{c} \quad (8.18) \]

which describes the fusion space. In the most commonly considered case, \( N_{ab}^{c} = 1 \) in which case there is a unique state and we do not need to specify \( \mu \) since it has only one possible value. Cases where \( N_{ab}^{c} = 0 \) are non-allowed fusions meaning that the space \( V_{ab}^{c} \) and \( V_{ab}^{c} \) are zero dimensional.

In Eq. 8.17 we have described states in the space associated with

---

\(^{19}\)Those who feel they have a good understanding of the physics without needing the mathematics may be able to skip this section.
a single anyon splitting into two, and in Eq. 8.18 we have described
states in the space associated with two anyons fusing into one. It is
possible to also describe the splitting or fusion space for a single anyon
splitting or fusing into more pieces. For example, when splitting/fusing
into three pieces the relevant spaces are often denoted as
\[ V_{abc} \]
We do not mean the identity here. See note 11 of this chapter. We use this notation to match
that of the next chapter.

Hermitian conjugate \( V^e_{abc} \). A basis of states in this space can be written
diagrammatically with the above described splitting vertices as

\[ |(a, b); d, \mu \rangle \otimes |d, c; e, \nu \rangle \in V_d^a \otimes V_e^d \in V_{abc}^e \quad (8.19) \]

The full splitting space \( V_{abc}^e \) can thus be described as
\[ V_{abc}^e \cong \bigoplus_d V_d^a \otimes V_e^d \] (8.20)

with a corresponding dimension of this space
\[ \dim V_{abc}^e = \sum_d N_d^a N_e^d . \quad (8.21) \]

On the other hand, we could just as well have described a state in this
space as

\[ |a, f; e, \lambda \rangle \otimes |b, c; f, \eta \rangle \in V_e^a \otimes V_f^b \in V_{abc}^e \quad (8.22) \]

In this language the full splitting space \( V_{abc}^e \) can be described as
\[ V_{abc}^e \cong \bigoplus_f V_e^a \otimes V_f^b \] (8.23)

with a corresponding dimension of this space
\[ \dim V_{abc}^e = \sum_d N_d^a N_f^b . \quad (8.24) \]

We thus have
\[ V_{abc}^e \cong \bigoplus_d V_d^a \otimes V_e^d \cong \bigoplus_f V_e^a \otimes V_f^b \] (8.25)

where “\( \cong \)” means “isomorphic to”. In other words, these are two iso-
morphic descriptions of the same space. Equating the two different
expressions (Eq. 8.21 and 8.24) for the dimension of this space recovers
the equality Eq. 8.11. The isomorphism between these two descriptions

\[ 20 \] We do not mean \( e \) to necessarily be the identity here. See note 11 of this chapter. We use this notation to match
that of the next chapter.

\[ 21 \] The insertion of the parenthesis \((a, b)\)
in Eq. 8.19 and similarly the parenthe-
sis \((b, c)\) in Eq. 8.22 are crucial to in-
dicate which splitting is closest to the
leaves of the tree (furthest from the
root). Without the parenthesis one can
have ambiguous notation, such as in the
Fibonacci theory, where
\[ |\tau \tau; \tau\rangle \otimes |\tau \tau; \tau\rangle \]
could mean either the state in Eq. 8.19
or Eq. 8.22. The notation is telling us
something important: that the kets in
Eq. 8.19 and Eq. 8.22 are living in dif-
ferent, albeit isomorphic, spaces.
of the same space will be explored in detail in the next chapter.

One can describe more complicated splitting and fusion spaces in an analogous way. For example, the space $V_{e}^{aaaaa}$ can be described as

$$V_{e}^{aaaaa} \cong \bigoplus_{b,c,d} V_{b}^{aa} \otimes V_{c}^{ba} \otimes V_{d}^{ca} \otimes V_{e}^{da}$$

where each term in the direct sum (i.e., each term with fixed $b,c,d$) is drawn diagrammatically as in Fig. 8.14.

**Chapter Summary**

- This is

**Further Reading**

This is some reading.

**Exercises**

**Exercise 8.1 Quantum Dimension**

Let $N_{ab}^{c}$ be the fusion multiplicity matrices of a TQFT

$$a \times b = \sum_{c} N_{ab}^{c} c$$

meaning that $N_{ab}^{c}$ is the number of distinct ways that $a$ and $b$ can fuse to $c$. (In many, or even most, theories of interest all $N$’s are either 0 or 1).

The quantum dimension $d_{a}$ of a particle $a$ is defined as the largest eigenvalue of the matrix $[N_{a}]_{b}$ where this is now thought of as a two dimensional matrix with $a$ fixed and $b,c$ the indices.

Show that

$$d_{a}d_{b} = \sum_{c} N_{ab}^{c} d_{c}$$

We will prove this formula algebraically in Chapter 17. However there is a simple and much more physical way to get to the the result: Imagine fusing together $M$ anyons of type $a$ and $M$ anyons of type $b$ where $M$ gets very large and determine the dimension of space that results. Then imagine fusing together $a \times b$ and do this $M$ times and then fuse together all the results.

**Exercise 8.2 Fusion and Ground State Degeneracy**

To determine the ground state degeneracy of a 2-manifold in a 2+1 dimensional TQFT one can cut the manifold into pieces and sew back together. One can think of the open “edges” or connecting tube-ends as each having a label given by one of the particle types (i.e., one of the anyons) of the theory. Really we are labeling each edge with a basis element of a possible Hilbert space. The labels on two tubes that have been connected together must match (label $a$ on one tube fits into label $\bar{a}$ on another tube.) To calculate the ground state degeneracy we must keep track of all possible ways that these assembled
tubes could have been labeled. For example, when we assemble a torus as in Fig. 8.16 we must match the quantum number on one open end to the (opposite) quantum number on the opposite open end. The ground state degeneracy is then just the number of different possible labels, or equivalently the number of different particle types.

For more complicated 2-d manifolds, we can decompose the manifold into so-called pants diagrams that look like Fig. 8.18. When we sew together pants diagrams, we should include a factor of the fusions multiplicity $N_{ab}^c$ for each pants which has its three tube edges labeled with $a$, $b$ and $c$.

(a) Write a general formula for the ground state degeneracy of an $M$-handled torus in terms of the $N$ matrices.

(b) For the Fibonacci anyon model, find the ground state degeneracy of a 4-handled torus.

(c) Show that in the limit of large number of handles $M$ the ground state degeneracy scales as $\sim D^{2M}$ where $D^2 = \sum_a d_a^2$.

Exercise 8.3 Consistency of Fusion Rules

Show by using commutativity and associativity of fusion along with identity $\mathbf{5.5}$ that no anyon theory can have a particle $a$ such that $a \times a = a$ meaning $a$ fuses to $a$ to form only $a$ and nothing else.
Change of Basis and $F^\dagger$-Matrices

Let us consider the case of three anyons $a$, $b$ and $c$ that fuse together to form an anyon $e$. As mentioned several times previously (See Fig. 8.15 or Eqs. 8.19 and 8.22) one can describe the state of these three particles in two different ways. We can describe the space by describing how $a$ fuses with $b$ (the value of $d$ on the left of Fig. 9.1), or by how $b$ fuses with $c$ (the value of $f$ on the right of Fig. 9.1). Either of these two descriptions should be able to describe any state of the three anyons $a$, $b$ and $c$ fusing to $e$. However, in the two different cases these states are described in different bases. We define the change of basis as a set of unitary matrices $2, 3, 4$ called $F$, as shown in Fig. 9.1.

Fig. 9.1 The $F$-matrix makes a change of basis between the two different ways of describing the space spanned by the fusion of three anyon charges $a$, $b$, and $c$ when they all fuse to a total quantum number of $e$. For fixed $a, b, c$ and $e$, the matrix $F$ is unitary in its subscripts $d, f$. Here $F$ is defined to be zero if the fusion diagram is not allowed, i.e., if any of the fusion multiplicities $N_{ab}^d$, $N_{dc}^e$, $N_{bc}^f$, $N_{af}^e$ are zero.

Several brief comments are in order. First, as noted in the caption of Fig. 9.1 the $F$-matrix is considered to be zero if any of the vertices on either side of the diagram are not allowed vertices of the fusion algebra. Secondly $F$ moves involving the identity particle (i.e., with $a$, $b$ or $c$ being the identity in the figure) are chosen to have value of unity $5$. In particular this means

$$[F_{e}^b]_{de} = [F_e^a]_{ac} = [F_e^b]_{eb} = 1$$

(9.1)

$4$ In the notation of section 8.5 the $F$-matrix describes the isomorphism $V_{d}^{ab} \otimes V_{e}^{dc} \cong V_{f}^{e} \otimes V_{f}^{bc}$. We can write the basis change more algebraically as

$$|(a, b); d) \otimes |d, c; e) = \sum_{f} [F_{e}^{abc}]_{df} |a, f; e) \otimes |b, c; f)$$

which represents Fig. 9.1 (where we have again suppressed indices $\mu, \ldots$ at the vertices for simplicity).

$5$ This involves a gauge choice, see section 9.5.

$6$ For simplicity we are assuming no fusion multiplicities $R_{ab}^e$ greater than 1. In cases where $N_{ab}^b > 1$ (as in Fig. 9.2), each vertex gets an additional index which ranges from 1 to its multiplicity so that the $F$-matrix gets additional indices as well. This case is discussed in section 9.5.5.

$3$ The conventions for writing $F$-matrices used in this chapter match that of Refs. Kitaev [2006] and Bonderson [2007].
Finally, being a change of basis, the $F$-matrix (for fixed $a, b, c, e$) is unitary viewed as a matrix with indices $d$ and $f$.

This idea of change of basis is familiar from angular momentum addition where the $F$-matrix is known as a $6j$ symbol (note it has 6 indices). One can combine three objects with $L^2$ angular momenta values $a, b$ and $c$ in order to get $L^2$ angular momentum $e$, and quite similarly you can describe this space in terms of $a$ combined with $b$ to get $d$ (as in the left of Fig. 9.1) or in terms of $b$ combined with $c$ to get $f$ (as in the right of Fig. 9.1). In fact, even when studying TQFTs, sometimes people refer to $F$-matrices as $6j$ symbols.

It is important to emphasize that an $F$-matrix can act on a portion of a diagram, as shown in Fig. 9.2. This allows us to convert any tree structure in a fusion diagram to any other tree structure.

### 9.1 Example: Fibonacci Anyons

Again we turn to the example of Fibonacci anyons for clarification. We imagine fusing together three $\tau$ particles. As shown in Fig. 8.10, there is a single state $|N\rangle$ in which the three fuse to the identity $I$. It should not matter if we choose to fuse the leftmost two anyons first, or the rightmost two. In either case there is only one possible state for the outcome. We can thus draw the simple identity shown in Fig. 9.3. Mathematically we would write that $F_{\tau\tau\tau} = 1$. (And as noted in Eq. 9.1 if any of the three upper indices are the identity, we also have $F = 1$). The more interesting situation is the case where the three Fibonacci anyons fuse to $\tau$. In this case, there is a two dimensional space of states, and this two dimensional space can be described in two ways. We can fuse the left two particles first to get either $I$ (yielding overall state $|0\rangle$) or to get $\tau$ (yielding overall state $|1\rangle$). See the top of Fig. 9.4. On the other hand, we could fuse the right two particles first to get either $I$ (yielding overall state $|0'\rangle$) or to get $\tau$ (yielding overall state $|1'\rangle$). See the bottom of Fig. 9.4.

The space of states spanned by the three anyons is the same in either
9.1 Example: Fibonacci Anyons

Fusing the two particles on the left first

\[ \tau \tau \tau = |0\rangle \]

\[ \tau \tau \tau = |1\rangle \]

Fusing the two particles on the right first

\[ \tau \tau \tau = |0'\rangle \]

\[ \tau \tau \tau = |1'\rangle \]

Fig. 9.4 Two ways to describe the same two dimensional space in the case of Fibonacci anyons. The basis \{\ket{0}, \ket{1}\} fuses the left two particles first, whereas the basis \{\ket{0'}, \ket{1'}\} fuses the right two particles first.

description. Thus, there must be a unitary basis transform given by

\[
\begin{pmatrix}
|0\rangle \\
|1\rangle
\end{pmatrix} =
\begin{pmatrix}
F_{00} & F_{01'} \\
F_{10'} & F_{11'}
\end{pmatrix}
\begin{pmatrix}
|0'\rangle \\
|1'\rangle
\end{pmatrix}
\]

(9.2)

Here \( F \) is a two by two matrix, and in the notation of the \( F \)-matrix defined in Fig. 9.1 this two by two matrix is \([F_{\tau \tau \tau}]_{ab}\) and the indices \( a, b \) should take the values \( I \) and \( \tau \) instead of 0 and 1, but we have used abbreviated notation here for more clarity.

For the Fibonacci theory the \( F \)-matrix is given explicitly by

\[
F_{\tau \tau \tau} = F = \begin{pmatrix}
\phi^{-1} & \phi^{-1/2} \\
\phi^{-1/2} & -\phi^{-1}
\end{pmatrix}
\]

(9.3)

where \( \phi^{-1} = (\sqrt{5} - 1)/2 \), so \( \phi \) is the golden mean. As one should expect for a change of basis, this matrix is unitary. In Section 9.3 we will discuss how this matrix is derived (See also section 18.2).

\[6\] We can redefine kets with different gauge choices (see section [13]) and this will insert some phases into the off-diagonal of this matrix, but the simplest gauge choice gives the matrix as shown.
9.2 Example: Ising Anyons

The situation with Ising anyons is quite similar, so we will be rather brief. Let us fuse three $\sigma$ particles to an overall fusion channel of $\sigma$. There is no other choice, three $\sigma$ particles can only fuse to $\sigma$ (i.e., there is no $|N\rangle$ state. See section 8.2.2). There are two possible states in the Hilbert space which we can write in either of two bases as shown in Fig. 9.5 — either fusing the left two particles first or fusing the right two particles first. Analogous to the Fibonacci case we can write an $F$-matrix which relates the two basis descriptions as in Eq. 9.2. However, here the $F$-matrix is instead given by

$$F_{\sigma\sigma\sigma} = F = \frac{1}{\sqrt{2}} \begin{pmatrix} 1 & 1 \\ 1 & -1 \end{pmatrix} \quad (9.4)$$

which is sometimes known as a Hadamard matrix. Deriving this form of the $F$-matrix will be described roughly in section 9.3 below, and is done in detail in sections 18.3 and 19.4 below (See also exercise 9.7).

In the Ising theory we can also look at situations where we have both $\sigma$ and $\psi$ particles. In this case we have

$$[F_{\sigma\sigma\sigma}]_{\sigma\sigma\sigma} = -1 \quad (9.5)$$
$$[F_{\psi\psi\sigma}]_{\sigma\sigma\sigma} = -1 \quad (9.6)$$

Eq. 9.5 is shown diagrammatically in Fig. 9.6. The other elements of $F$ in the Ising theory which we have not mentioned so far (i.e., those not described by Eqs. 9.4-9.6) are either 1 if all the fusion vertices are allowed, or are zero if any of the fusion vertices are not allowed (See Fig. 8.1 caption).

The presence of the minus signs in Eq. 9.5 (for example) may seem a bit puzzling being that the diagrams on the left and right of Fig. 9.6 are describing the same state in the Hilbert space. However, we will see in the next section why this sign is required in order to have a consistent $F$-matrix (See exercise 9.3 for a more detailed calculation).

9.3 Pentagon

It is possible to describe the same Hilbert space in many ways. For example, with three anyons, as in Fig. 8.15, one can describe the state in terms of the fusion channel of the two anyons on the left, or in terms of the two on the right. I.e., we can describe $(a \times b) \times c$ or $a \times (b \times c)$, and as in Fig. 9.1, these two descriptions can be related via an $F$-matrix.

When there are four anyons, there are still more options of how we group particles to describe the states of the Hilbert space, and these can also be related to each other via $F$-matrices similarly (analogous to that shown in Fig. 8.2). The fact that we can change the connectivity of these tree diagrams then allows one to make multiple changes in the trees as shown in Fig. 9.7. Indeed, in this figure one sees that one can
Fig. 9.7  Pentagon Diagram. Each step in the diagram is a new description of the same basis of states via and $F$-matrix.

go from the far left to the far right of the diagram via two completely different paths (the top and the bottom path) and the end result on the far right should be the same either way. This diagram, known as the pentagon diagram\(^8\), puts a very strong constraint on the $F$-matrices, which written out algebraically would be

$$
[F_{cd}^{f}][F_{e}^{a}]_{f}^{h} = \sum_{k} [F_{b}^{a}]^{h}^{g} [F_{d}^{h}][F_{e}^{a}]^{k}^{l} [F_{c}^{l}][F_{d}^{k}]^{h}^{l}
$$

(9.7)

where the left hand side represents the top route of the figure and the right hand side represents the bottom route.\(^9\)

For very simple theories, such as the Fibonacci anyon theory, the fusion rules and the Pentagon diagram are sufficient to completely define the $F$-matrices (up to some gauge convention choices as in section 9.4). See exercise \[^5\] Further, for any given set of fusion rules there are a finite set of possible solutions of the pentagon equation\(^10\) — a property that goes by the name “Ocneanu rigidity”\(^11\).

One might think that one could write down more complicated trees and more complicated paths through the trees analogous to Fig. 9.7 and somehow derive additional constraints on the $F$-matrices. A theorem by MacLane \[^{[1971]}\], known as the “coherence theorem”, guarantees that no more complicated trees generate new identities beyond the pentagon diagram.

8 An analogous relation holds for $6j$ symbols of angular momentum addition, known often as the Elliott-Biedenharn identity.

9 It is very worth working through this to make sure you understand how this equation matches up with the figure! Note that in the equation the $F$-matrices are written in an order such that those furthest right in Fig. 9.7 are furthest right in the equations.

10 A finite set of gauge inequivalent solutions. I.e., a gauge transform of a given solution does not count as a new solution.

11 Ocneanu did not manage to ever publish this important result. See for example Etingof et al [2003].

9.4  Gauge Transforms

We have the freedom to make gauge transformations on our diagrams and these will be reflected in the $F$-matrix. While this is a bit of a technical point, we make frequent use gauge transformation in some later chapters so it is worth discussing it briefly here.
A gauge choice is a choice of a phase associated with the vertices in a diagram. If we chance this gauge choice, diagrams are then multiplied by phases.

In particular a gauge transformation multiplies the vertices in a diagram by a phase as shown in Fig. 9.8. The tilde over the vertex on the right notates that we have made a gauge transform to a tilde gauge.

\[ |a, b; c \rangle = u_{ab}^c |a, b; c \rangle \]

As we shall see in section 14.2, some gauge choices are much more natural than others, but we should always keep in mind that we have this freedom.

Note that if one of the upper legs is the identity \((a = I)\) or \((b = I)\) in Fig. 9.8 we typically do not allow a gauge transform of this type of vertex, since the presence of a vertex with the vacuum is the same as the absence of a vertex with the vacuum (i.e., we can add or remove lines labeled by \(I\) for free).

9.5 Appendix: \(F\)-matrix Odds and Ends

9.5.1 Product Theories

Given two anyon theories \(T\) and \(t\), we can construct the product theory \(T \times t\) as in section 8.5. If the theory \(T\) has consistent \(F\)-matrices \([F_{ABC}]_{DF}\) and the theory \(t\) has consistent \(F\)-matrices \([F_{abc}]_{df}\) ("consistent" here means satisfying the pentagon relation), then the product theory has consistent \(F\)-matrices

\[ [F_{(A,a)(B,b)(C,c)}]_{(D,d),(F,f)} = [F_{E}^{ABC}]_{DF} [F_{e}^{abc}]_{df} \]

The point here is that in a product theory, the two constituent theories don’t “see” each other at all.
9.5.2 Unitarity of $F$

The $F$-matrix relation we defined as

$$a \, b \, c \, d \, e = \sum_{f} [F_{e}^{abc}]_{df}$$

The fact that $F$ is unitary in its indices $d$ and $f$ means we can also write

$$a \, b \, c \, d \, e = \sum_{d} [F_{e}^{abc}]^{*} \, (f_{d})$$

9.5.3 $F$-matrix with higher fusion multiplicities

In cases where there are fusion multiplicities $N_{ab}^{c}$ greater than 1, each vertex gets an additional index as shown in Fig. 8.12. The $F$-matrix must also describe what happens to these indices under basis transform. We thus have a more general basis-change equation given in Fig. 9.9.

$$a \, b \, c \, d \, e, \mu, \nu = \sum_{f, \alpha, \beta} [F_{e}^{abc}]_{(d_{\mu \nu})} \, (f_{\alpha \beta})$$

Fig. 9.9 The $F$-matrix equation with fusion multiplicities greater than one. Here the vertex indices are $\mu \in 1 \ldots N_{ab}^{c}$ and $\nu \in 1 \ldots N_{bc}^{e}$, and $\alpha \in 1 \ldots N_{af}^{e}$ and $\beta \in 1 \ldots N_{ef}^{c}$. The subscripts $(d_{\mu \nu})$ and $(f_{\alpha \beta})$ are "super-indices", of the matrix $F_{e}^{abc}$. I.e., $d_{\mu}$ and $\nu$ are joined together to make a single index.

In the language of section 8.6 this $F$-transform is written as

$$|a, b; d, \mu \rangle \otimes |d, c; e, \nu \rangle = \sum_{f, \alpha, \beta} [F_{e}^{abc}]_{(d_{\mu \nu})(f_{\alpha \beta})} |a, f; e, \beta \rangle \otimes |b, c; f, \alpha \rangle$$

Gauge Transforms with higher fusion multiplicities

With higher fusion multiplicities $N_{ab}^{c} > 1$, our diagrams have indices at the vertices. Gauge transforms are generally a unitary matrix within this index space and then take the form shown in Fig. 8.10.

$$|a, b; c, \mu \rangle = \sum_{\mu'} |u_{\mu}^{ab} \rangle_{\mu' \mu} |a, b; c, \mu \rangle$$

Again, this is much more easily expressed using the notation of section 8.6, where we can just write

$$|a, b; c, \mu \rangle = \sum_{\mu'} [u_{\mu}^{ab}]_{\mu' \mu} |a, b; c, \mu \rangle$$
Fig. 9.10 We have the freedom to make a gauge transform of a vertex by multiplying by a unitary matrix $[u_{c}^{ab}]_{\mu\mu'}$. The tilde on the right notates that the vertex is in the tilde gauge.

Chapter Summary

- This is the summary

Further Reading

Exercises

Exercise 9.1 F-gauge choice

(a) Explain why in the Fibonacci theory, $[F_{\tau\tau\tau}]_{\tau\tau}$ is gauge independent but $[F_{\tau\tau\tau}]_{\tau\tau}$ is gauge dependent.

(b) Explain why in the Ising theory is $[F_{\sigma\psi\sigma}]_{\sigma\sigma}$ is gauge independent, but $[F_{\sigma\psi\sigma}]_{\sigma\sigma}$ is gauge dependent.

Exercise 9.2 F’s with the vacuum field $I$

Explain why $[F_{e}^{aIc}]_{ac} = [F_{d}^{aI}]_{ab} = [F_{f}^{I}be]_{be} = 1$.

Exercise 9.3 Ising Pentagon

Consider a system of Ising anyons. Given the fusion rules, $F_{\sigma\psi\sigma\psi}$ will be a 2 by 2 matrix in the case of $x = y = z = w = \sigma$ (given by Eq. 9.4.4) and is a simply a scalar otherwise. One might hope that these scalars can all be taken to be unity. Unfortunately this is not the case. By examining the pentagon equation, Eq. 9.7.1 in the case of $a = b = c = \sigma$ and $d = f = \psi$ show that taking the scalar to always be unity is not consistent. Show further that choosing $[F_{\psi\psi\psi}]_{\sigma\sigma} = -1$ (and leaving the other scalars to be unity) allows a consistent solution of the pentagon for $a = b = c = \sigma$ and $d = f = \psi$.

Exercise 9.4 Fibonacci Pentagon

In the Fibonacci anyon model, there are two particle types which are usually called $I$ and $\tau$. The only nontrivial fusion rule is $\tau \times \tau = I + \tau$. With these fusion rules, the $F$-matrix is completely fixed up to a gauge freedom (corresponding to adding a phase to some of the kets). If we choose all elements of the $F$-matrix to be real, then the $F$-matrix is completely determined by
the pentagon up to one sign (gauge) choice. Using the pentagon equation determine the $F$-matrix. (To get you started, note that in Fig. 9.7 the variables $a, b, c, d, e, f, g, h$ can only take values $I$ and $\tau$. You only need to consider the cases where $a, b, c, d$ are all $\tau$).

If you are stuck as to how to start, part of the calculation is given in Nayak et al. [2008].

**Exercise 9.5 Pentagon and Fusion Multiplicities**

Consider the case of Appendix 9.5.3 where there are fusion multiplicities $N_{ab}^c > 1$. Write the generalization of the pentagon equation Eq. 9.7.

**Exercise 9.6 Gauge Change**

(a.i) Confirm that the $F$-matrix transforms under gauge change as indicated in Eq. 9.8. (a.ii) Show that a solution of the pentagon equation remains a solution under any gauge transformation.

(b.i) Analogous to (a.i) Confirm Eq. 9.9. (b.ii) Analogous to (a.ii) show that a solution of the pentagon equation remains a solution under any gauge transformation. (You will need to solve problem 9.5 first!)

**Exercise 9.7 Ising F-matrix**

[Hard] As discussed in the earlier problem, “Ising Anyons and Majorana Fermions” (Ex. 4.3), one can express Ising anyons in terms of Majorana fermions which are operators $\gamma$, with anticommutations $\{\gamma_i, \gamma_j\} = 2\delta_{ij}$. As discussed there we can choose any two Majoranas and construct a fermion operator

$$c_{12}^\dagger = \frac{1}{2}(\gamma_1 + i\gamma_2)$$

then the corresponding fermion orbital can be either filled or empty. We might write this as $|0_{12}\rangle = c_{12}^\dagger |1_{12}\rangle$ and $|1_{12}\rangle = c_{12}^\dagger |0_{12}\rangle$. The subscript 12 here meaning that we have made the orbital out of Majoranas number 1 and 2. Note however, that we have to be careful that $|0_{12}\rangle = e^{i\phi} |1_{12}\rangle$ where $\phi$ is a gauge choice which is arbitrary (think about this if it is not obvious already).

Let us consider a system of 4 Majoranas, $\gamma_1, \gamma_2, \gamma_3, \gamma_4$. Consider the basis of states

$$|a\rangle = |0_{12}0_{34}\rangle$$

$$|b\rangle = |0_{12}1_{34}\rangle$$

$$|c\rangle = |1_{12}0_{34}\rangle$$

$$|d\rangle = |1_{12}1_{34}\rangle$$

rewrite these states in terms of basis of states

$$|a'\rangle = |0_{41}0_{23}\rangle$$

$$|b'\rangle = |0_{41}1_{23}\rangle$$

$$|c'\rangle = |1_{41}0_{23}\rangle$$

$$|d'\rangle = |1_{41}1_{23}\rangle$$

Hence determine the $F$-matrix for Ising anyons. Be cautious about fermionic anticommutations: $c_{ij}^\dagger c_{ij}^\dagger = -c_{ij}^\dagger c_{ij}^\dagger$ so if we define $|1_x1_y\rangle = c_{ij}^\dagger |0_x0_y\rangle$ with the convention that $|0_x0_y\rangle = |0_y0_x\rangle$ then we will have $|1_x1_y\rangle = -|1_y1_x\rangle$. Note also that you have to make a gauge choice of some phases (analogous to the mentioned gauge choice above). You can choose $F$ to be always real.
Exchanging Identical Particles

We would now like to determine what happens when two particles are exchanged with each other. As one might expect for anyons, phases are accumulated from such exchanges. However, one must be cautious because the phase accumulated will generally depend on the fusion channel of the particles being exchanged.

10.1 Introducing the $R$-matrix

Let us begin with a simple case where two identical particles of type $a$ are braided around each other. Let us specify that the two particles fuse together in an overall channel $c$. Let us call this quantum mechanical state $|\text{state}\rangle$ as shown in two different notations in Fig. 10.1.

\[
\begin{array}{c}
\begin{array}{c}
\text{a} \\
\text{c}
\end{array}
&
\begin{array}{c}
\begin{array}{c}
\text{a} \\
\text{a}
\end{array}
\end{array}
\end{array}
= \begin{array}{c}
\begin{array}{c}
\text{a} \quad \text{a} \\
\text{c}
\end{array}
\end{array}
= |\text{state}\rangle
\]

Fig. 10.1 Two $a$ particles fusing to a $c$ particle.

We then (half)-braid the two particles around each other (counterclockwise observing from above\(^1\)). The final fusion channel of the two $a$ particles is still $c$ (by the locality principle of section 8.2). However, a phase will be accumulated in the process which call $R_{ac}^{aa}$ as shown in Fig. 10.2. The inverse phase would be accumulated for an exchange in the opposite direction.

\[
\begin{array}{c}
\begin{array}{c}
\text{a} \\
\text{c}
\end{array}
&
\begin{array}{c}
\begin{array}{c}
\text{a} \\
\text{a}
\end{array}
\end{array}
\end{array}
= \begin{array}{c}
\begin{array}{c}
\text{a} \quad \text{a} \\
\text{c}
\end{array}
\end{array}
= R_{ac}^{aa} |\text{state}\rangle
\]

Fig. 10.2 The phase accumulated by exchanging two $a$ particles that fuse to $c$ is called $R_{ac}^{aa}$.

\(^1\)In the language of the braid group we would call this exchange $\sigma$. See section 3.3.1.

This so-called $R$-matrix along with the corresponding $F$-matrices will allow us to compute the result of braiding any number of $a$ particles around each other in arbitrary ways.

Let us consider the case of three anyons of type $a$. We can write a basis for the possible states of three anyons as shown in Fig. 10.3.
Exchanging Identical Particles

Fig. 10.3 A basis of states for three \( \alpha \) type anyons fusing to an overall quantum number \( f \).

\[
\alpha \alpha \alpha \overset{\alpha \alpha \alpha}{\chi} f = |\alpha; f\rangle
\]

Fig. 10.4 Exchanging the two left particles incurs a phase \( R^{\alpha\alpha}_c \).

\[
\hat{\sigma}_1 |\alpha; f\rangle = \alpha \overset{\alpha \alpha \alpha}{\chi} f = \sum_{c}^\alpha \alpha \alpha \overset{\alpha \alpha \alpha}{\chi} f = R^{\alpha\alpha}_c |\alpha; f\rangle
\]

We now consider exchanging the left two particles as shown in Fig. 10.4 (We call the operator that performs this exchange \( \hat{\sigma}_1 \) in analogy with the braid group discussed in section 3.3.1.) Since we know the fusion channel of these two particles (\( \alpha \)) we know that the phase accumulated in this exchange is just \( R^{\alpha\alpha}_c \). This seems fairly simple as it is precisely the type of exchange we defined in Fig. 10.2 above.

As with all operators in quantum mechanics that can be implemented as a time evolution, the exchange operator is linear, meaning that it acts on superpositions by acting on each term individually:

\[
\hat{\sigma}_1 \sum_c^\alpha \alpha c |\alpha; f\rangle = \sum_c^\alpha \alpha c R^{\alpha\alpha}_c |\alpha; f\rangle
\]

Let us now instead consider exchanging the right two particles, an operation we call \( \hat{\sigma}_2 \). Since the right two particles are not in a definite fusion channel we cannot directly apply the \( R \)-matrix. However, we can use the \( F \)-matrix to rewrite our state as a superposition of states where the right two particles are in a definite fusion channel as shown in Fig. 10.5.

\[
\alpha \overset{\alpha \alpha \alpha}{\chi} f = \sum_g [F^\alpha_{\alpha\alpha} f]_{cg} \alpha \overset{\alpha \alpha \alpha}{\chi} f
\]

Fig. 10.5 Using an \( F \)-move to work in the basis with a known fusion channel of the right two particles.

One we have transformed to this new basis, then we can exchange the right two particles and apply the \( R \)-matrix directly to the right two particles as shown in Fig. 10.6. Once we have established the effect of the exchange we can (if desired) convert back into the original basis which describes the fusion of the left two particles using \( F^{-1} \).
Introducing the $R$-matrix

$$\hat{\sigma}_2|c; f\rangle = \sum_g [F_f^{aaa}]_{cg}$$

$$= \sum_g R_g^{aa} [F_f^{aaa}]_{cg}$$

Fig. 10.6 In order to describe exchange of the right two particles, we first change
to a basis where the fusion channel of those two particles is explicit. We can then
apply the $R$-matrix directly.

The result of this procedure in terms of the original basis, is given by

$$\hat{\sigma}_2|c; f\rangle = \sum_{g,z} [F_f^{aaa}]_{cg} R_g^{aa} [(F_f^{aaa})^{-1}]_{gz} |z; f\rangle$$

The general principle is that to evaluate any exchange of identical
particles, we can always use $F$-matrices to convert to a basis where the
fusion channel of the two particles to be braided is known. Once we are
working in this basis, we can then apply the $R$-matrix directly. At
the end we can transform back to the original basis if we so desire. This
scheme works for any set of identical particles given appropriate $F$-
and $R$-matrices.

10.1.1 Locality

An important principle which we will often use is that result of braiding
a group of particles with a given total quantum number $c$ is the same as if
that entire group were replaced with just a single particle with quantum
number $c$. For example, in Fig. 10.7 when we braid a cluster of $a, b$ with
overall quantum number $c$ around a cluster $x, y, z$ with overall quantum
number $w$, the phase accumulated should be the same as if we simply
braided $c$ around $w$.

Fig. 10.7 Braiding a cluster of particles with overall quantum number $c$ around a
cluster of particles with overall quantum number $w$ should have the same effect as
braiding a single $c$ particle, around a single $w$ particle. The result should just be a
phase dependent on the quantum numbers $c, w$ and $f$. In chapter 13 we will refer to
this phase as $R_f^{cw} R_{cw}$. Note that $a$ does not wrap around $b$. If it did that would
accumulate an additional phase.
10.2 Some Examples

Since the idea of using the $R$-matrix is quite important, it is worth working through a few examples explicitly.

10.2.1 Fibonacci Anyons

Recall the properties of Fibonacci anyons (see section 8.2.1): There is only one nontrivial particle type which we call $\tau$ and the only nontrivial fusion rule is $\tau \times \tau = I + \tau$. As we saw in section 8.2.1, the fusion rule implies that there are two possible states of two Fibonacci anyons: The state where they fuse together to form $I$ and the state where they fuse together to $\tau$. (See Fig. 10.8. See also Fig. 8.9 where we previously introduced these two states). We call these states $|I\rangle$ and $|\tau\rangle$ respectively.

Now consider the operator $\hat{\sigma}$ that exchanges the two Fibonacci anyons counterclockwise as viewed from above, as shown in Fig. 10.9. This operator yields the phase $R_{\tau\tau}^I$ if the fusion channel of the two particles is $I$ or $R_{\tau\tau}^\tau$ if the fusion channel of the two particles is $\tau$.

$$\hat{\sigma}|I\rangle = \begin{array}{c} \tau \tau \\ I \end{array} = R_{\tau\tau}^I |I\rangle$$
$$\hat{\sigma}|\tau\rangle = \begin{array}{c} \tau \tau \\ \tau \end{array} = R_{\tau\tau}^\tau |\tau\rangle$$

In section 13.3 below (see also exercise 10.6) we will explain how we actually compute the phases $R_{\tau\tau}^I$ and $R_{\tau\tau}^\tau$. For now, it suffices to give the answers that for right-handed Fibonacci anyons

$$R_{\tau\tau}^I = e^{i3\pi/5} \quad R_{\tau\tau}^\tau = e^{-4\pi i/5}$$

There also exists a left-handed type of Fibonacci anyons for which the phases are complex conjugate of these.

As with all operators in quantum mechanics we can act on superpositions by acting on each term individually:

$$\hat{\sigma} \left( \alpha|I\rangle + \beta|\tau\rangle \right) = \alpha R_{\tau\tau}^I |I\rangle + \beta R_{\tau\tau}^\tau |\tau\rangle$$

If we think of our two states $|I\rangle$ and $|\tau\rangle$ as the two states of a qubit, the $\hat{\sigma}$ operator is what is known as a controlled phase gate in quantum information processing — the phase accumulated depends on the state of the qubit.

Next let us consider the possible states of three Fibonacci anyon. As
described in section 8.2.1 the space of such states is three-dimensional, and we can choose as a basis the two states shown in Fig. 10.10 (we already introduced these states in Fig. 8.10 above). Now consider an operator $\hat{\sigma}_1$ that braids the two leftmost particles around each other as shown in Fig. 10.11. Here the phase accumulated depends on the fusion channel of the leftmost two particles, entirely analogous to Fig. 10.9.

$$\hat{\sigma}_1|N\rangle = \begin{array}{c}
\begin{array}{c}
\tau \\
\tau \\
\tau \\
\tau
\end{array}
\end{array} = R^\tau_\tau |N\rangle$$

$$\hat{\sigma}_1|0\rangle = \begin{array}{c}
\begin{array}{c}
\tau \\
\tau \\
\tau \\
\tau
\end{array}
\end{array} = R^\tau_\tau |0\rangle$$

$$\hat{\sigma}_1|1\rangle = \begin{array}{c}
\begin{array}{c}
\tau \\
\tau \\
\tau \\
\tau
\end{array}
\end{array} = R^\tau_\tau |1\rangle$$

Fig. 10.11 Exchanging the left two particles.

More interesting is the question of what happens if we exchange the right two particles as shown in Fig. 10.12. As discussed in Section 10.1, the trick here is to use the $F$-matrix to change the basis such that we know the fusion channel of the right two particles, and then once we know the fusion channel we can use the $R$-matrix. If we want, we can then use the $F$-matrix to transform back to the original basis.

$$\hat{\sigma}_2|0\rangle = \begin{array}{c}
\begin{array}{c}
\tau \\
\tau \\
\tau \\
\tau
\end{array}
\end{array} = \begin{array}{c}
\begin{array}{c}
\tau \\
\tau \\
\tau \\
\tau
\end{array}
\end{array} = |0'\rangle$$

Fig. 10.12 Exchanging the two particles on the right for the $|0\rangle$ state where these two particles on the right are not in a definite fusion channel. Note that in the tree diagram on the left the state below the dashed red line is exactly $|0\rangle$.

To see how this works, Recall that we can use the $F$-matrix to write the $|0\rangle$ state in the basis of the $|0'\rangle$ and $|1'\rangle$ as in Eq. 9.2 which we reproduce the relevant parts of here:

$$|0\rangle = F_{00'}|0'\rangle + F_{01'}|1'\rangle$$

where $|0'\rangle$ and $|1'\rangle$ are shown in Fig. 10.13. Note that here $F_{ab}$ is shorthand for $[F^TT^T]_{ab}$.

Fig. 10.13 In the prime basis the two particles on the right are in a definite fusion channel.
On the right hand side of Fig. 10.13 (i.e., in the prime basis) we know the fusion channel of the rightmost two particles, so we can braid them around each other and use the $R$-matrix to compute the corresponding phase as shown in Fig. 10.14.

\[
\begin{align*}
\tau \tau \tau &= F_{00'} \tau \tau \tau + F_{01'} \tau \tau \tau \\
\tau \tau \tau &= F_{00'} R_1^{\tau \tau} \tau \tau \tau + F_{01'} R_1^{\tau \tau} \tau \tau \tau \\
\tau \tau \tau &= F_{00'} R_1^{\tau \tau} |0'\rangle + F_{01'} R_1^{\tau \tau} |1'\rangle \\
\tau \tau \tau &= F_{00'} R_1^{\tau \tau} (\langle F^{-1} |0'0|0\rangle + \langle F^{-1} |0'1|1\rangle) \\
\tau \tau \tau &= F_{00'} R_1^{\tau \tau} \left(\langle F^{-1} |1'0|0\rangle + \langle F^{-1} |1'1|1\rangle \right) \\
\tau \tau \tau &= (F_{00'} R_1^{\tau \tau} [F^{-1}]_{0'0} + F_{01'} R_1^{\tau \tau} [F^{-1}]_{1'0}) |0\rangle \\
\tau \tau \tau &= (F_{00'} R_1^{\tau \tau} [F^{-1}]_{0'1} + F_{01'} R_1^{\tau \tau} [F^{-1}]_{1'1}) |1\rangle
\end{align*}
\]

Fig. 10.14 To exchange the right two particles we first use an $F$-move so that we know the fusion channel of these two particles, then we can apply $R$ and then $F^{-1}$ to transform back into the original basis.

Where between Eq. 10.4 and 10.5 we have used the inverse $F$ transform to put the result back in the original [0] and [1] basis. The final result, Eq. 10.6 is precisely the same as Eq. 10.1 just written out in all of its detail.  

We can summarize the results of the two possible braiding operations on the three dimensional Hilbert space. Assuming right-handed Fibonacci anyons and using a basis $|N\rangle, |0\rangle, |1\rangle$ (also notated as $|\tau; I\rangle, |I; \tau\rangle, |\tau; \tau\rangle$) we have

\[
\begin{align*}
\delta_1 &= \begin{pmatrix}
e^{3\pi i/5} & e^{-4\pi i/5} \\
e^{3\pi i/5} & e^{3\pi i/5}
\end{pmatrix} \\
\delta_2 &= \begin{pmatrix}
\phi^{-1} e^{4\pi i/5} & \phi^{-1/2} e^{-3\pi i/5} \\
\phi^{-1/2} e^{-3\pi i/5} & -\phi^{-1}
\end{pmatrix}
\end{align*}
\]

where $\phi = (\sqrt{5} + 1)/2$ is the golden mean.

### 10.2.2 Ising Anyons

For Ising anyons the situation is perhaps even simpler since three $\sigma$ particles have only two fusion channels (See section 8.2.2). The appropriate $F$-matrices are given by Eq. 9.3 and the $R$-matrices for a right-handed...
Exercises

Ising theory are given by

\[ R^\sigma_\sigma = e^{-i\pi/8} \]  
\[ R^\sigma_\psi = e^{i3\pi/8} = iR^\sigma_\sigma \]  

with the \( R \)-matrices for a left-handed theory being the complex conjugates of these expressions. From the \( R \)-matrix, we immediately obtain the form of the exchange operator \( \hat{\sigma}_1 \), that counterclockwise exchanges the leftmost two Ising anyons

\[ \hat{\sigma}_1 = e^{-i\pi/8} \begin{pmatrix} 1 & 0 \\ 0 & i \end{pmatrix} \]

(10.10)

Then using Eq. 10.11 we can evaluate the exchange operator \( \hat{\sigma}_2 \) which counterclockwise exchanges the rightmost two anyons of the three, giving

\[ \hat{\sigma}_2 = \frac{e^{i\pi/8}}{\sqrt{2}} \begin{pmatrix} 1 & -i \\ -i & 1 \end{pmatrix} \]

(10.11)

This is some reading

Exercises

Exercise 10.1 Calculating Exchanges
(a) Use Eq. 10.1 to confirm Eq. 10.11
(b) Use Eq. 10.1 to confirm Eq. 10.7
(c) Confirm the braiding relation \( \hat{\sigma}_1 \hat{\sigma}_2 \hat{\sigma}_1 = \hat{\sigma}_2 \hat{\sigma}_1 \hat{\sigma}_2 \) in both cases. What does this identity mean geometrically. See exercise 3.1.

Exercise 10.2 Ising Anyons Redux
In exercise 3.3 we introduced a representation for the exchange matrices for Ising anyons which, for three anyons, would be of the form

\[ \hat{\sigma}_1 = \frac{e^{i\alpha}}{\sqrt{2}}(1 + \gamma_1 \gamma_2) \]

(10.12)

\[ \hat{\sigma}_2 = \frac{e^{i\alpha}}{\sqrt{2}}(1 + \gamma_2 \gamma_3) \]

(10.13)

where the \( \gamma \)'s are Majorana operators defined by

\[ \{\gamma_i, \gamma_j\} \equiv \gamma_i \gamma_j + \gamma_j \gamma_i = 2\delta_{ij} \]

with \( \gamma_i = \gamma_i^\dagger \).

Show that the exchange matrices in Eq. 10.11 are equivalent to this representation. How does one represent the \( |0\rangle \) and \( |1\rangle \) state of the Hilbert space
120 Exercises

in this language? The answer may not be unique.

**Exercise 10.3 Exchanging More Particles**

(a) Consider a system of 4 identical Ising anyons. Use the $F$- and $R$-matrices to calculate the braid matrices $\hat{\sigma}_1, \hat{\sigma}_2,$ and $\hat{\sigma}_3$. (You should be able to check your answer using the Majorana representation of exercise 10.3)

(b) [Harder] Consider a system of 4 identical Fibonacci anyons. Use the $F$- and $R$-matrices to calculate the braid matrices $\hat{\sigma}_1, \hat{\sigma}_2,$ and $\hat{\sigma}_3$.

**Exercise 10.4 Determinant and Trace of Braid Matrices**

Consider a system of $N$-identical anyons with a total Hilbert space dimension $D$. The braid matrices $\hat{\sigma}_1, \hat{\sigma}_2,...,\hat{\sigma}_{N-1}$ are all $D$-dimensional. Show that each of these matrices has the same determinant, and each of these matrices has the same trace. Hint: This is easy if you think about it right!

**Exercise 10.5 Checking the locality constraint**

[a] Easy Consider Fig. 10.16. The braid on the left can be written as $\hat{b}_3 = \hat{\sigma}_2 \hat{\sigma}_1 \hat{\sigma}_2$.

(a) For the Fibonacci theory with $a = \tau$ check that the matrix $\hat{b}_3$ gives just a phase, which is dependent on the fusion channel $c$. I.e., show the matrix $\hat{b}_3$ is a diagonal matrix of complex phases. Show further that these phases are the same as the phase that would be accumulated for taking a single $\tau$ particle around the particle $c$.

(b) Consider the same braid for the Ising theory with $a = \sigma$. Show again that the result is a $c$-dependent phase.

[Hard] Consider the braid shown on the left of Fig. 10.17. The braid can be written as $\hat{b}_4 = \hat{\sigma}_3 \hat{\sigma}_2 \hat{\sigma}_1 \hat{\sigma}_2$.

(c) Consider Ising anyons where $a = \sigma$. Use the $F$ and $R$-matrices to calculate $\hat{\sigma}_3$ (See exercise 10.3 a). Since the fusion of three $\sigma$ anyons always gives $c = \sigma$, calculate $b_4$, show this is a phase times the identity matrix, and show that the phase matches the phase of taking a single $\sigma$ all the way around another $\sigma$.

(d) Consider Fibonacci anyons with $a = \tau$. Use the $F$ and $R$-matrices to calculate $\hat{\sigma}_3$. (See exercise 10.3 b). Check that $\hat{b}_4$ is a diagonal matrix of phases. Check the phases match the two possible phases accumulated by wrapping a single $\tau$ all the way around a single particle $c$ which can be $I$ or $\tau$.

**Exercise 10.6 Enforcing the locality constraint**

The locality constraint shown in Fig. 10.16 turns out to be extremely powerful. In this exercise we will use this constraint to (almost) derive the possible values for the $R$-matrix for Fibonacci anyons given the known $F$-matrix.

Consider an anyon theory with Fibonacci fusion rules and Fibonacci $F$-matrix as in Eq. 10.2.

(a) [Easy] Confirm the locality constraint shown in Fig. 10.16 (see also Fig. 10.7) given the values of $R$ given in Eq. 10.2. Make sure to confirm the equality for all three cases $f = I, c = \tau$ and $f = \tau, c = I$ and $f = \tau, c = \tau$.

Note that on the left of Fig. 10.16 is the braiding operation $\hat{O} = \hat{\sigma}_2 \hat{\sigma}_1 \hat{\sigma}_2$, whereas the operation on the right is $\sigma^\tau$.

(b) Show that the locality constraint of Fig. 10.16 would also be satisfied by

$$R_I^\tau \rightarrow -R_I^\tau \quad R_\sigma^\tau \rightarrow -R_\sigma^\tau$$ (10.14)

It will turn out (See *** below) that this additional solution is spurious, as there are other consistency conditions it does not satisfy.
(c) In addition to right and left handed Fibonacci anyons and the two additional spurious solutions provided by Eq. 10.14, there are four additional possible sets of $R$-matrices that are consistent with the $F$-matrices of the Fibonacci theory given the locality constraint of Fig. 10.16. These additional solutions are all fairly trivial. Can you guess any of them?

If we cannot guess the additional possible $R$-matrices, we can derive them explicitly (and show that no others exist). Let us suppose that we do not know the values of the $R$-matrix elements $R^\tau_\tau$ and $R^\tau_I$.

(d) For the case of $f = I$ and $c = \tau$ show that Fig. 10.16 implies

$$[R^\tau_\tau]^4 = [R^\tau_I]^2 \quad (10.15)$$

(e) [Harder] For the case of $f = \tau$ we have a two-dimensional Hilbert space spanned by the two values of $c = I$ or $c = \tau$. Any linear operator on this Hilbert space should be a 2 by 2 matrix. Thus the locality constraint Eq. 10.16 is actually an equality of 2 by 2 matrices. Derive this equality.

(f) Use this result, in combination with Eq. 10.15 to find all possible $R$-matrices that satisfy the locality constraint. You should find a total of eight solutions. Six of these are spurious as we will see in section 13.3.

The calculation you have just done is equivalent to enforcing the so-called hexagon condition which we will discuss in section 13.3 below.
Computing with Anyons

Having discussed the basics of anyon theories, we are now in a position to discuss how one might perform quantum computations with braids.

In chapter 2 we briefly introduced some ideas of topological quantum computation. In chapter 8 we discussed how we might define a qubit in several simple anyon theories. In the current chapter we will briefly discuss how anyons can be used to fulfill the requirements for quantum computation.

11.1 Quantum Computing

To have a quantum computer, we must first have a Hilbert space, and we usually think of this Hilbert space as being built from small pieces, such as qubits or qutrits. This Hilbert space will be the quantum memory that the computer acts on.

Once we have our Hilbert space, our model of a quantum computer has three key steps for quantum computation:

1. Find a Hilbert space to work with.
2. Initialize the Hilbert space in some known state.
3. Perform a controlled unitary operation on the Hilbert space.
4. Measure some degree of freedom in the Hilbert space.

If the controlled unitary (step 2) is implemented as a series of unitary operations each of which acts on only small parts of the Hilbert space (such as acting on just a few qubits at a time), we call this scheme for quantum computation the quantum circuit model.

We will discuss each of the above steps (0)-(3) for our anyon systems in section 11.2 below. First, however, we will introduce the idea of what it means for a quantum computer to be “universal” in the quantum circuit model.

There are other models of quantum computation. We mention in particular the measurement schemes (See Raussendorf and Briegel [2001] [Cross et al. [2002]], where no unitary is explicitly performed, but rather the computation is implemented as a series of measurements on an initial highly entangled state. In the context of topological quantum computation an important variant is a computation that is implemented by a combination of unitary operations and projective measurements. The earliest proposal for quantum computing with anyons, by Kitaev in 1997, was of this type (See Kitaev [1997]). See also footnote [15] below.

1 For more of the basics of quantum computation, a classic reference is Nielsen and Chuang [2000]. We also provide a bit more basic information in section 24.1.

2 Qubits are two state systems (such as a spin-½), qutrits are three state systems etc. The general case is known as a qudit. See the introduction to quantum information in chapter 24.

3 There are variants on this theme. For example, it might be sufficient to initialize into a state that is only partially known, or it might be sufficient to have a somewhat noisy measurement. Most interesting is the issue of whether one can tolerate some amount of imperfection in the system (noise in the system, uncontrolled operations on the Hilbert space, etc). We will discuss this issue further in chapter 24.

4 There are other models of quantum computation. We mention in particular the measurement schemes (See Raussendorf and Briegel [2001] [Cross et al. [2002]], where no unitary is explicitly performed, but rather the computation is implemented as a series of measurements on an initial highly entangled state. In the context of topological quantum computation an important variant is a computation that is implemented by a combination of unitary operations and projective measurements. The earliest proposal for quantum computing with anyons, by Kitaev in 1997, was of this type (See Kitaev [1997]). See also footnote [15] below.
5Recall a matrix $U$ is unitary if and only if $U^*U = UU^* = 1$.

6Quantum mechanical time evolution is always unitary. This is simply the statement that a normalized ket remains normalized. It is worth noting that we are excluding the possibility of making measurements (which are often considered to be nonunitary) on the outcomes of the end of the computation. This would be outside of the quantum circuit model.

7All of quantum mechanics can be viewed as unitary time evolution. Measurements may look like they are non-unitary, but one can always include the measuring apparatus within the system being considered and then the full system (including the measuring apparatus) then obeys unitary evolution. The idea of including measurement within your system in order to maintain unitarity is sometimes known as “the church of the larger Hilbert space”.

8Recall that the last step of a quantum computation, after applying a unitary $U$ (via some sequence of gates as in Eq. 11.1) to our Hilbert space, we obtain an output “answer” by measuring whether some particular qubits are in the $|0\rangle$ state or the $|1\rangle$ state. The probabilities of these outcomes is completely independent of the overall phase of the $U$. I.e., if we changed $U \rightarrow e^{i\phi}U$ we would have the same probabilities of outcomes.

9I believe this distance measure was introduced by Fowler [2011]. Other definitions of distance can also be used. Relationships between this distance measure and more conventional operator norms are given by Field and Simula [2012] and Amy [2013].

11.1.1 Universal Quantum Computing in the Quantum Circuit Model

Let us suppose our Hilbert space consists of $N$ qubits (each qubit being a two state system). The Hilbert space dimension is then $D = 2^N$. The space of possible unitary operations on these qubits, is just the group of $D$ dimensional unitary matrices --- a group known as $U(D)$.

Let us now suppose our quantum computer can implement any one of $p$ different elementary operations (usually called “gates”) in a single time step (each gates will act only on a small number of qubits). Each gate corresponds to a particular unitary operation $U_n \in U(D)$ with $n \in 1, \ldots, p$ that is applied to the Hilbert space. A sequence of such gates constructs a particular unitary operation which is just the product of the successive gates (the time order runs from right to left)

$$U = U_{i_1} \ldots U_{i_2} U_{i_1}$$ (11.1)

where the number of gates $t$, can be thought of as the “run time” of the computation.

Suppose there is some particular computation we would like to perform, and this computation corresponds to a unitary $U$ which we hope to construct via a series of gates as in Eq. (11.1). Note, however, that in quantum computation we are never worried about the overall phase of our result. As such if we want to construct some particular unitary $U$, for the purpose of quantum computation, it just as good to construct $e^{i\phi}U$ for any value of $\phi$.

Unfortunately, even with this freedom of phase most unitary operations (except for a set of measure zero) are actually impossible to construct exactly from a finite set of elementary gates as in Eq. (11.1). Fortunately, for computational purposes it is good enough to approximate the desired unitary operation to some (potentially high) accuracy. Sets of gates that can always make such an accurate approximation are called universal. We will be more precise about the definition of this word in a moment.

Since we will be discussing approximations of desired operations, it is useful to define a distance between two unitary matrices so we can measure the accuracy of our approximation. Given two $D$ dimensional unitary matrices $U$ and $V$, we define a phase invariant distance measure between them as

$$\text{dist}(U;V) = \sqrt{1 - \frac{|\text{Tr}[U^*V]|}{D}}$$ (11.2)

Note that multiplying either matrix by an overall phase leaves dist unchanged, and if $U$ and $V$ are the same up to a phase, then dist is zero. We say that $V$ is a good approximation of $U$ up to a phase if $\text{dist}(U;V)$ is small.

Having defined this distance measure, we can be more precise about what we mean that a set of gates is universal. A gate set $U_n \in U(D)$
with \( n \in 1, \ldots, p \) is universal if for any any desired target operation we would like to perform \( U_{\text{target}} \in U(D) \) we can find a sequence of gates \( U_{i_2} \ldots U_{i_1} U_{i_3} \) such that the phase invariant distance to the target is less than any desired error tolerance \( \epsilon \)

\[
\text{dist}(U_{\text{target}}; U_{i_2} \ldots U_{i_1} U_{i_3}) < \epsilon
\] (11.3)

no matter how small an \( \epsilon \) we choose. In other words, our gate set can approximate any target unitary as precisely as we want.

We might wonder how long a run time (how many gates) will we typically need to have? A beautiful theorem by Kitaev and Solovay assures us that the run time is not too long. In particular,

\[
t \sim O \left( \log \left( \frac{1}{\epsilon} \right) \right)
\] (11.4)

We are thus guaranteed that if we have a universal gate set, then the run time of the computer gets at most logarithmically longer as we try to increase the quality of our approximation of the target operation \( U_{\text{target}} \).

The essence of this theorem is as follows. If we consider a sequence of \( t \) gates, (i.e., a run time of \( t \)), if there are \( p \) different elementary gates, we can construct roughly \( p^t \) different sequences of gates. Thus as \( t \) gets larger, there are exponentially more possible unitaries we can construct and these roughly cover the space \( U(D) \) evenly. With the number of points we can construct in this space growing exponentially with \( t \), the distance \( \epsilon \) of an arbitrary target unitary to the nearest unitary we can construct must drop exponentially with \( t \), hence justifying Eq. 11.3.

It is a nontrivial calculation to determine which set of elementary gates is sufficient to have a quantum computer which is universal. However, an important result is that if one can perform arbitrary rotations on a single qubit and in addition if one can perform any entangling two-qubit operation between any of these two bits (or even between just neighboring bits), then one has a universal quantum computer.

10. The Kitaev-Solovay theorem, often viewed as one of the most fundamental results of quantum computation, is discussed nicely in Dawson and Nielsen [2006] and Harrow [2001].

11. The usual proof of the Kitaev-Solovay theorem assumes that the gate set must contain inverses. In other words, if \( U_n \) is one of the elementary gates, then \( U_n^{-1} \) should also be one of the elementary gates.

12. We will not get exactly \( p^t \) different unitaries, since more than one sequences might generate the same unitary operation.

13. This important theorem is sometimes known as the Brylinski theorem after its discoverers, Brylinski and Brylinski [2002]. The authors are married. A simpler version of the proof is given by Bremner et al. [2002].

### 11.2 Topological Quantum Computing

#### 11.2.1 Hilbert space

With a topological quantum computer, a qubit (or qutrit, etc.) can be formed from multiple anyons which can be put into multiple fusion channels (See chapter 8). For example, with Fibonacci anyons a qubit might be formed from three Fibonacci anyons fusing to \( \tau \) as shown in Fig. 9.4. With the Ising theory, one might use a cluster of three Ising anyons fusing to \( \sigma \) as as qubit as shown in Fig. 9.5. There are, of course, many more options of how one encodes a qubit in any given theory. For example, in the Ising theory it may be more convenient to work with clusters of four Ising anyons fusing to \( I \) as shown in Fig. 11.1.

![Fig. 11.1 A qubit made from four Ising anyons in an overall fusion channel of \( I \). The two states of the qubit are \( \psi \) and \( \phi \). Note that due to the fusion rules of the Ising theory, if the overall state of the four qubits is \( I \), then if the left two anyons are in state \( \psi \), the right two must also be in state \( \psi \). Using a qubit made of 4-anyons has advantages for other topological theories such as \( SU(2)_k \) with \( k > 4 \). See, for example, Hormozi et al. [2009].](image-url)
11.2.2 Measurement (in brief) and initialization

A topological qubit could be measured in several ways, depending on the particular physical system in question. The general principle of locality which we introduced in section 8.2 (See in particular Fig. 8.8) gives a good idea how such measurements can (or can’t) be done.

Let us suppose, for example, we have two anyons of type \(a\) and we would like to measure their fusion channel. Given the principle of locality, to measure this fusion channel we must perform an operation which is local to both particles, i.e., a measurement that surrounds both.

One way to measure the fusion channel of two anyons is to bring them together to the same point, or at least bring them physically close on a microscopic scale. When two anyons are microscopically close to each other, in essence their wavefunctions mix with each other and in this case measurement of almost any nontrivial operator near that location will suffice to distinguish between the different possible fusion channels. For example, one could measure the energy of the two anyons, or the force between them, which would generally distinguish the fusion channels. Note, however, when the anyons are moved macroscopically far apart no local operators should be able to distinguish the fusion channels. (We will discuss precisely why this is the case in chapter *** below).

Another way to measure the fusion channel of two anyons would be to leave the two anyons far apart from each other but implement a measurement that surrounds them both — such as Aharanov-Bohm-type interference as shown in Fig. 11.2. Here a test particle wave is split into two partial waves which travel on opposite sides of the anyons to be measured and then reinterfere with each other. This is entirely analogous to the regular Aharanov-Bohm effect (See section 4.1 and Fig. 4.2), where the partial waves travel on opposite sides of a flux and then reinterfere. In the usual Aharanov-Bohm effect, the net phase we measure is the phase of wrapping a single test particle all the way around the central region (See Eq. 4.3). Analogously here we measure the phase of wrapping the probe anyon all the way around the anyons in the central region to measure their fusion channel as shown in Fig. 11.3. As we will discuss in chapter ***, experiments of this sort have been attempted in quantum Hall systems.

Once we know how to measure the state of the anyons in our Hilbert space (and assuming we know how to manipulate our qubits) it is then fairly trivial to initialize the Hilbert space. We simply measure the state of a qubit: If it is in the state we want, we are done. If it is in some other state, we apply the appropriate unitary operation to put it into the desired initial state. We will discuss unitary operations next.

11.2.3 Universal Braiding

The most interesting part of a topological quantum computation is the idea that we can apply a controlled unitary operation on our Hilbert space by braiding anyons around each other. The elementary gates of
the system (or elementary unitary operations) are the (counterclockwise)
exchanges of two identical anyons, which, in braid group notation, we
call $\hat{\sigma}_n$, as well as the inverse (clockwise) exchanges $\hat{\sigma}_n^{-1}$, where $n \in 1, 2, \ldots, (N - 1)$ for a system of $N$ identical anyons. Each of these
braid operators corresponds to a unitary matrix operating on the Hilbert
space.

It turns out that for many types of nonabelian anyon theories, the
gate set made up of elementary braiding exchanges is universal in the
sense defined in section 11.1.1\textsuperscript{14}. For example, braiding is universal for
Fibonacci anyons. Similarly $SU(2)_k$ Chern-Simons theory is universal
for $k = 3$ and $k > 4$. In fact, among nonabelian anyon theories, theories
where braiding is not universal are somewhat of an exception. Ising
anyons and the closely related $SU(2)_2$ Chern-Simons anyons are two of
these nonuniversal exception\textsuperscript{15}.

It turns out that any system of $N$ identical anyons that is capable of
universal quantum computation by braiding, is also capable of universal
quantum computation by weaving\textsuperscript{16}. Here, what we mean by “weave” is
that we fix the positions of $N - 1$ of the anyons and only move the one
remaining anyon around all the other stationary anyons. An example of
a weave is shown in Fig. 11.4. The weaves are a very restricted subset
of the possible braids, but still the weaves form a universal set of gates
for these anyon systems. This result will be important below in section
11.4.1 (See also exercise 11.5).

In fact, if one is able to measure fusion channels easily\textsuperscript{17}, it is also
possible to implement universal quantum computation just by making
many measurements of fusion channels, without physically braiding any
particles around any others\textsuperscript{18}.

11.3 Fibonacci Example

As an example, we will focus on the case of Fibonacci anyons, which is
potentially the simplest anyon system which is universal for quantum
computation.

11.3.1 A Single Fibonacci Qubit

Let us consider a single qubit made of three Fibonacci anyons. We have
discussed this several times before in sections 8.2.1 and 10.2.1.

To remind the reader, there are three possible states of three Fibonacci
anyons which we label $|N\rangle$, $|0\rangle$, $|1\rangle$ (See Fig. 8.10 — which represents
a qubit (the states $|0\rangle$ and $|1\rangle$) and one additional “noncomputational”
state $|N\rangle$ which we will not use for storing quantum information.

\textsuperscript{14}This result was shown by Freedman et al. 2002a [4]. These papers are not particularly easy to read for physicists.

\textsuperscript{15}SU(2)\textsubscript{4} is an interesting case where braiding alone is not universal. However, if we are allowed to go outside of the quantum
circuit model and combine braiding with many projective measurements (i.e., not just making one measurement at the end
of the computation), then $SU(2)\textsubscript{4}$ anyons can implement universal quantum computation (See Refs. Levaillant et al. 2013;
Cui and Wang 2013). In fact, the first proposal of a topological quantum computer, by Kitaev in 1997 (published as Kitaev
1999), described a computation scheme which involved both braiding and projective measurement. A simple discussion of
this scheme is given by Preskill 2004 with early extensions of the scheme given by Mochon 2003 2004.

\textsuperscript{16}Note that measurement schemes of the Aharonov-Bohm type, as in
Fig. 11.3 involve braiding a test particle around other particles.

\textsuperscript{17}See Ref. Bonderson et al. 2008a.

\textsuperscript{18}See Fig. 11.4 A weave is a braid where
only one particle moves and all the
other particles remain stationary. All
the particles in this figure are supposed
to be of the same type. The single particle
that moves is colored red just for clarity.

\textsuperscript{19}This is proven by Simon et al. 2006.
We now think about braiding our three anyons. In the braid group on three strands, $B_3$ (See section 3.3.1), there are two generators, $\sigma_1$, exchanging the first two strands counterclockwise, and $\sigma_2$, exchanging the second two strands counterclockwise. Any braid of three particles can be constructed as some product of $\sigma_1, \sigma_2, \sigma_1^{-1},$ and $\sigma_2^{-1}$ in some order as shown, for example, in Fig. 11.5.

The action of these braid operations on the three-dimensional Hilbert space is shown in Eqs. 10.6 and 10.7 which we calculated in section 10.2.1. By multiplying these matrices together, we can figure out how any complicated braid acts on our Hilbert space. In fact for now we are only interested in how the matrices act on the space of the qubit states $|0\rangle$ and $|1\rangle$ and we will return to worry about the $|N\rangle$ state below in section 11.4.1.

**Example of X Gate**

We are now interested in the following simple quantum computation problem: Given a particular target unitary operation which we might want to perform on our qubit, how should we move the anyons? I.e., what braid should we do to implement the target operation?

For example, suppose we want to design a braid that implements an X-gate (just a Pauli $\sigma_x$)

$$U_{\text{target}} = X = \begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix} \quad (11.5)$$

With a very short braid (Shown in Fig. 11.5), we can make a fairly poor approximation to this gate (this braid is the best we can do with only five braid operations) given by

$$U_{\text{approx}} = \hat{\sigma}_2^{-1} \hat{\sigma}_1^3 \hat{\sigma}_2^{-1} \approx e^{-3\pi i/5} \begin{pmatrix} 0.073 - 0.225i & 0.972 \\ 0.972 & -0.073 - 0.225i \end{pmatrix} \quad (11.6)$$

For the approximation given in Eq. 11.6 the phase invariant distance from the target is

$$\text{dist}(U_{\text{target}} : U_{\text{approx}}) \approx 0.17$$

which is not a great approximation. However, with a longer braid having nine braid operations, shown in Fig. 11.6, one can make a better approximation with a trace distance $\text{dist} \approx 0.08$. If we consider braids that are longer and longer, we can get successively better approximations to the desired target as would be expected from the Kitaev-Solovay theorem discussed in section 11.1.1.

As mentioned in section 11.2.3 it is possible to find braids that are weaves, meaning that only a single anyon moves. For completeness, we show a weave in Fig. 11.7 that implements an X-gate to precision $\text{dist} \approx 0.18$. Note that due to the restricted weave form of this braid, a slightly larger number of elementary exchanges are required to reach...
roughly the same precision as in Fig. 11.5. As with braids, at least in principle, by using longer weaves one can get as close to the target as we like.

11.3.2 Topological Quantum Compiling: Single Qubit

Even if there exists a braid that performs a unitary operation that approximates some target operation within some small error distance \( \epsilon \), it is a nontrivial task to figure out what that braid is. In other words, how do you know what braid you should implement on your computer in order to perform the desired operation?

The general task of determining which elementary gates should be performed, and in what order, to implement some desired target unitary is known as quantum compiling\(^{21}\). For a topological quantum computer, the task of designing a braid is therefore known as topological quantum compiling. Here we will discuss several approaches to topological quantum compiling in order of their complexity, and their effectiveness.\(^{22}\)

We continue to focus only on compiling braids for a single Fibonacci qubit. Multi-qubit braids will be discussed in section 11.4 below.

Brute Force Search

If we are willing to accept a fairly poor approximation of our target unitary (a fairly large \( \text{dist} \) between our approximation and the target) we will be able to use a fairly short sequence of our elementary gates (i.e., a short braid). In this case we can consider some maximum gate sequence length \( t \) (maximum run time) and search all possible gate sequences of length less than \( t \), choosing the one that best approximates our target. We should expect to achieve a distance to the target that drops exponentially with \( t \), as discussed near Eq. 11.4.

If we are considering a single qubit made of three Fibonacci anyons, our elementary gates are the braid generators \( \hat{\sigma}_1, \hat{\sigma}_2, \hat{\sigma}_1^{-1}, \hat{\sigma}_2^{-1} \). This means that if we want to search through all braids of length \( t \) we have to search roughly \( 4^t \) braids. While there are some tricks that allow us to reduce this number somewhat\(^{23}\), the computational effort\(^{24}\) will always grow exponentially with the length \( t \). If one wants to make highly accurate approximation of a target unitary with a very small error distance, run times \( t \) can become large enough that brute force searching becomes unfeasible.

Kitaev-Solovay Algorithm

Kitaev and Solovay\(^{25}\) provide us an explicit algorithm to construct very accurate approximations of any desired unitary given a universal set of elementary gates with reasonable (not exponentially growing!) computational effort\(^{23}\). The essence of this algorithm is as follows. Let us suppose that by brute-force search we can approximate any unitary op-

![Fig. 11.7](image-url)
uration to within a distance $\text{dist} \sim \epsilon_0$ with a sequence of elementary 
gates (elementary braids in the topological case) of length $t_0$. Let us say 
that the classical computational time to achieve this is $T_0$. Now given 
a target unitary $U_{\text{target}}^{(0)}$ that would like to approximate, we start with 
this brute-force search, and construct our approximation $U_{\text{approx}}^{(0)}$ which 
is accurate to within $\text{dist} \sim \epsilon_0$. This is our 0th level of approximation 
of the target. We would next like to repair this approximation with 
another series of gates to make it more accurate. We thus define 

$$U_{\text{target}}^{(1)} = [U_{\text{approx}}^{(0)}]^{-1} U_{\text{target}}^{(0)}.$$

If we could find a series of gates that would exactly give us $U_{\text{target}}^{(1)}$ we 
could exactly construct the original objective $U_{\text{target}}^{(0)}$ as 

$$U_{\text{target}}^{(0)} = U_{\text{approx}}^{(0)} U_{\text{target}}^{(1)}.$$

However, it is not obvious that we have any better way to approximate 
$U_{\text{target}}^{(1)}$ than we had to approximate $U_{\text{target}}^{(0)}$, so why does this help? The 
key here is that $U_{\text{target}}^{(1)}$ is necessarily close ($\text{dist} \sim \epsilon_0$) to the identity. 
We then decompose 

$$U_{\text{target}}^{(1)} = VWV^{-1}W^{-1}$$

with $W$ and $V$ being unitary operations close to the identity ($\text{dist} \sim \sqrt{\epsilon_0}$). We then have an amazing result, that if we are able to approximate 
$V$ and $W$ to an accuracy $\epsilon_0$ (which we can do here by brute-force search) 
we will get $U_{\text{target}}^{(1)}$ accurate to $\text{dist} \sim \epsilon_0^{3/2}$. Thus we obtain 

$$U_{\text{target}}^{(0)} = U_{\text{approx}}^{(0)} VWV^{-1}W^{-1}$$

accurate to order $\epsilon_0^{3/2}$. The total sequence of gates is now of length $5t_0$ 
since each of other factors on the right hand side of Eq. 11.7 is of length $t_0$. The classical computational effort to achieve this is roughly $3T_0$ since 
we must search for $U_{\text{approx}}^{(0)}$ and $V$ and $W$.

This scheme can then be iterated to make our approximation even 
better. The only change it that the next level of approximation, instead 
of using brute force search to make approximations good to $\text{dist} \sim \epsilon_0$ 
we use the entire above described algorithm to make all of our approximations 
good to $\text{dist} \sim \epsilon_0^{3/2}$. When $U_{\text{approx}}^{(0)}$ and $V$ and $W$ are calculated 
to order $\epsilon_0^{3/2}$ our new approximation for $U_{\text{target}}^{(0)}$ will be an accurate to 
$\text{dist} \sim (\epsilon_0^{3/2})^{3/2}$.

The entire scheme can be iterated recursively to any level of accuracy. 
At the $n^{th}$ level of this approximation, we have a gate sequence of length 
$5^n t_0$ and an accuracy $\text{dist} \sim \epsilon_0^{(3/2)^n}$ and the computational effort to 
achieve this scales as $3^n T_0$.

Thus if we want to achieve some overall accuracy $\epsilon$ of our operation,
the gate sequence will be of length
\[ t \sim \mathcal{O}\left(\log(1/\epsilon)^{\left[\log(\ln(5)/\ln(3/2))\right]}\right) = \mathcal{O}\left(\log(1/\epsilon)^{3.969}\right) \]
and this requires classical computation time
\[ T \sim \mathcal{O}\left(\log(1/\epsilon)^{\left[\log(\ln(3)/\ln(3/2))\right]}\right) = \mathcal{O}\left(\log(1/\epsilon)^{2.710}\right). \]
While this algorithm produces gate sequences that are longer than one obtains with brute force searching (which produces gate sequence lengths as in Eq. 11.4) for the same desired accuracy \( \epsilon \), it has the advantage that it is computationally feasible for much smaller values of \( \epsilon \) and can therefore produce more accurate results.

**Galois Theory Optimal Compiling:**

A rather remarkable scheme for quantum compiling was developed in Kliuchnikov et al. [2014] based on ideas from Galois theory. While we cannot review Galois theory here, nor can we even do justice to the details of the algorithm, we can nonetheless discuss some of structure of the problem that makes this approach possible.

It turns out that any unitary that can be constructed by braiding three Fibonacci anyons can be written (up to a phase) in the form
\[ U(u, v, k) = \begin{pmatrix} u & v^* \omega^k \phi^{-1/2} \\ v \phi^{-1/2} & -u^* \omega^k \end{pmatrix} \]
where \( \phi = (1 + \sqrt{5})/2 \) is the golden mean, \( k \) is an integer, \( \omega = e^{2\pi i/10} \), and
\[ |u|^2 + \phi |v|^2 = 1 \]
where \( u \) and \( v \) come from the so-called ring of cyclotomic integers \( \mathbb{Z}[\omega] \), which means that
\[ u = \sum_{i=0}^{3} a_i \omega^i \quad \quad v = \sum_{i=0}^{3} b_i \omega^i \]
with coefficients \( a_i \) and \( b_i \) all being integers. The fact that the unitaries that can be generated by braiding take a very restricted mathematical form is, in fact, a generic property of all anyon theories, although the particular form taken depends on the particular anyon theory.

Further, given values of \( u, v, \) and \( k \) a relatively simple algorithm is provided that finds a braid that results exactly in this unitary, where length of the braid is no longer than
\[ t \sim \log \left( \sum_{i=0}^{3} |a_i \omega^i|^2 + \sum_{i=0}^{3} |a_i \omega^{3i}|^2 \right) \]
This procedure is known as *exact synthesis* as it constructs exactly the
The braid shown here between two Fibonacci qubits entangles the two qubits but also results in leakage error. When we use three Fibonacci anyons as a qubit, we set the overall fusion channel of the three to be $\tau$, so $c = d = \tau$ in this figure. The quantum information is stored in the quantum numbers $a$ and $b$. The shown braid results in some amplitude ending up in the noncomputational space where either $c$ or $d$ is $I$ rather than $\tau$.

The end result of this approach is an algorithm that, although it does not find the absolute optimal braid\(^{29}\), it nonetheless is asymptotically optimal in the sense that it produces braids of length

$$t \sim O(\log(1/\epsilon))$$

as in Eq. 11.4. Further, the computational time to achieve this scales only as $T \sim O(\log(1/\epsilon)^2)$. Using this type of approach, it is easily possible to generate braids with error distances of order $10^{-100}$ or even better, and these braids are longer than the absolute optimal braid by only a factor of order unity.

### 11.4 Two-Qubit Gates

Having studied single qubit operations, we now turn to a brief discussion of two-qubit gates\(^{30}\). As mentioned in section 11.1.1 the Brylinski theorem tells us that to have a universal quantum computer, we need only have single qubit rotations along with any entangling two qubit gate. To construct such an entangling two qubit gate we will need to have a braid that physically entangles the world lines of the anyons comprising the two qubits such as the example shown in Fig. 11.8.

However there is a crucial complication with braiding anyons between qubits. If we perform a braid such as that shown in Fig. 11.8 the fusion channel of the anyons comprising each of the qubits (quantum numbers $c$ and $d$ in the figure) are not preserved (see the discussion of locality in section 8.6) and this means that amplitude can leak into the noncomputational space.

To be more explicit for the Fibonacci case, recall that we encode our qubits ($|0\rangle$ or $|1\rangle$) by using three Fibonacci anyons in overall fusion

\(^{29}\)The “optimal braid” is the one that would be found by brute force search if one had the exponentially enormous computational power necessary to find it.

\(^{30}\)Here we are constructing a two qubit unitary operation, which we will call a two-qubit gate, from our elementary gates — the elementary braid operations.
channel $\tau$ (See Fig. 8.10). The fusion channel of the three anyons to $I$ is termed non-computational $|N\rangle$ and is not used for computation. If some of the amplitude of the wavefunction ends up in this noncomputational space, it is called leakage error, and only very small quantities of such leakage errors can be tolerated for any realistic computation. Braids like the one shown in Fig. 11.8 always produce some amplitude of noncomputational states. The problem of leakage error in two-qubit gates is not special to Fibonacci anyons, but is in fact a generic property of all anyon theories that have universal braiding.

While we cannot completely eliminate leakage, we can in principle design entangling gates with arbitrarily small (albeit non-zero) leakage. Such braids with low leakage error do exist, but finding them is highly nontrivial. Inconveniently, the Hilbert space of six Fibonacci anyons, as in Fig. 11.8 is 13 dimensional. Searching such a large space for particular unitaries with low leakage is numerically unfeasible. We thus need a more clever way to design braids with low leakage.

In designing any computation, it is almost always advantageous to simplify the desired task into smaller tasks that can be addressed one at a time. This “divide and conquer” approach will allow us to tackle the job of designing two-qubit gates. In the next section we will give an example of how entangling gates with negligible leakage can be designed.

### 11.4.1 Controlled Gates

In quantum computation it is often very convenient to use entangling gates which are so-called controlled $U$-gates, or $C(U)$ where $U$ is a 2-dimensional unitary matrix. A controlled $U$-gate acts on two qubits such that one qubit (the “target” qubit) is acted on with a 2-dimensional unitary operator $U$ if and only if the other qubit (the “control” qubit) is in the $|1\rangle$ state, whereas the control qubit remains unchanged:

$$C(U):egin{align*}
|0\rangle \otimes |0\rangle &\rightarrow |0\rangle \otimes |0\rangle \\
|1\rangle \otimes |0\rangle &\rightarrow |1\rangle \otimes |0\rangle \\
|0\rangle \otimes |1\rangle &\rightarrow (U|0\rangle) \otimes |1\rangle \\
|1\rangle \otimes |1\rangle &\rightarrow (U|1\rangle) \otimes |1\rangle
\end{align*}$$

Thus the first qubit here is being controlled by the second qubit. A very commonly used example of a controlled gate is the case of $U = X$ (See Eq. 11.5) which we call a controlled-X, or more often a controlled-NOT (or CNOT) gate.

The key to our construction of controlled gates is the locality principle of section 10.1.1. If we are given a cluster of 2 anyons which are in the $\tau$ fusion channel (for example, set $c = \tau$ in Fig. 10.7) and we braid it around some other anyons, this will have the same effect as if we just braided a single $\tau$ around the other anyons. However, if the cluster of 2 anyons is in the trivial (or $I$) fusion channel, then braiding this cluster never does anything, as braiding the vacuum particle always is trivial. Thus we can see that the effect of the braid is “controlled” by the fusion channel of the two anyons.

---

31 Ainsworth and Slingerland [2011] show that it is not possible to design completely leakage free gates for any universal anyon theory, and leakage can only be made approximately zero.

32 This space is subdivided into an 8 dimensional subspace with $f = \tau$ and a 5 dimensional subspace with $f = I$. No braiding of these six anyons will change the $f$ quantum number. Note, however, that gates must have low leakage independent of the value of $f$.

33 As always in quantum mechanics the operation acts linearly on superpositions.

34 The constructions discussed here were introduced in Bonesteel et al. [2005].
Controlled $\hat{\sigma}_2^2$ gate

Consider the construction shown in Fig. 11.10. On the far left of this figure, we have shown a weave, meaning only a single anyon, the one drawn in red, moves and the other two anyons remain stationary (see the discussion by Fig. 11.4). This weave has been designed to have approximately the same effect as if the two blue anyons are wrapped around each other (exchanged twice counterclockwise), i.e., $\hat{\sigma}_2^2$ as shown in the figure. For the particular weave shown, the distance to the target $\hat{\sigma}_2^2$ is $\text{dist} \approx 0.12$. We could make the approximation of $\hat{\sigma}_2^2$ more accurate by using a longer weave using any of the compiling methods discussed in section 11.3.2 above. Note that the equivalence between the weave on the far left and $\sigma_2^2$ is true as a $3 \times 3$ matrix acting on the full three dimensional Hilbert space spanned by three Fibonacci anyons (i.e., on the space spanned by $|0\rangle$, $|1\rangle$, $|N\rangle$, not just $|0\rangle, |1\rangle$).

![Fig. 11.10](image.png)

Fig. 11.10 Construction of a controlled gate using Fibonacci anyons. The weave on the far left is designed to have almost the same effect on the Hilbert space as the braiding (two counterclockwise exchanges) of the two blue particles as shown. Using a longer weave one can more closely approximate the braiding of the two blue particles. On the right, we have a system of 6 anyons representing two qubits. The right (red) we take the cluster of two red anyons, labeled $b$.

Now consider the braid of six anyons on the right of Fig. 11.10 representing two qubits — the right (red) anyons are the control qubit and the left (blue) anyons are the target qubits. We group the two red anyons in fusion channel $b$ and we move them around as a single unit to form the same weave as shown on the far left (here using the two red anyons and the right two anyons of the blue qubit). If these two red anyons are in the vacuum fusion channel $b = I$, then this braiding has no effect on the Hilbert space (braiding of the vacuum particle is always trivial). On the other hand, if the two anyons are in fusion channel $b = \tau$ then this braid is equivalent to moving a single $\tau$ particle through exactly the same weave as on the far left, thus having the same effect as exchanging the two right-most blue anyons twice counterclockwise. We have thus
constructed a controlled operation, \( C(\hat{\sigma}_2) \) which is notated on the far right of the figure in a manner analogous to Fig. 11.9. The operation implemented on the blue qubit is (approximately) a full braiding of the right two blue strands, if and only if the right qubit \((b)\) is in the \(\tau\) or \(|1\rangle\) state.

A crucial feature of this construction is that, to the extent that the weave we use accurately approximates \( \hat{\sigma}_2 \), the resulting construction leads to no leakage error. The right hand qubit \((b)\) is completely unchanged (hence not creating leakage of this qubit), and the effect on the left hand qubit \((a)\) is designed to be equivalent to just braiding two of the blue anyons — which does not create leakage either.

### Controlled \( U \)-gate

With a bit more work, we can in fact make any controlled \( C(U) \) gate for an arbitrary two dimensional unitary \( U \), as in Fig. 11.9 (up to an overall phase as discussed in section 11.1.1).

First let us discuss the so-called injection weave described in Fig. 11.11. An ideal injection weave is meant to leave the Hilbert space unchanged (it only applies an identity matrix). However, it has the nontrivial effect of rearranging the three strands comprising a qubit. As shown in the diagram, the injection weave moves the red strand from the far right at the bottom to the far left at the top. As discussed in section 11.3.2 we can more precisely approximate the ideal injection by using a longer weave.

We now construct the braid shown on the left of Fig. 11.12. As in Fig. 11.10 we group together the two red anyons in fusion channel \( b \) and we move them around as a group. These two anyons are first put through an injection weave with the right most two blue anyons. This moves the group of two anyons into the middle position of the right qubit. A weave to implement an arbitrary unitary \( U \) is then implemented on the three strands furthest left, treating the two red strands grouped together as a single strand. Finally, this injection weave is inverted to bring the two red particles back to their original position. The braid constructed in this way will implement a controlled \( U \) gate \( \text{C}(U) \), as shown using the notation of Fig. 11.9 on the right of Fig. 11.12. The left (blue) qubit \((a)\) has the unitary \( U \) applied to it, if and only if the right qubit \((b)\) is in the \(|1\rangle\) or \(\tau\) state. If we choose \( U \) to be an \( X \) gate, such that the necessary weave in the middle step is a weave like that shown in Fig. 11.7, we obtain a \( \text{C}(X) \) or controlled NOT gate (CNOT).

To understand this procedure we realize that the only two anyons that are moved in this procedure are the two red anyons in state \( b \), and these two are moved as a group. As in Fig. 11.10 if these two anyons are in fusion channel \( b = I \) (or \(|0\rangle\)) then the Hilbert space is left unchanged. However, if \( b = \tau \) (or \(|1\rangle\)) then there will be an effect on the blue qubit — hence we have a controlled rotation. Let us now consider this case when \( b = \tau \), so that we should think of the two red strands as being a single \( \tau \) strand. The injection weaves are designed to have no effect on

---

**Fig. 11.11** An approximate injection weave is shown on the left. A perfect injection has no effect on the Hilbert space (it applies the identity matrix to the Hilbert space) but moves the red strand from the right to the left of the three anyons. The distance to the target for this particular weave is \( \text{dist} = 0.09 \). With a longer weave one can more accurately approximate a perfect injection.
the Hilbert space — their only effect is to move the red double strand inside of the blue qubit. The unitary rotation $\hat{U}$ is thus only nontrivial effect on the Hilbert space.

Chapter Summary

- Summary Item 1
- Summary Item 2 etc

Further Reading

Most of the same ideas can be generalized for other anyon systems.

- Freedman et al. [2002a] Freedman et al. [2002b] for the initial work showing SU(2)k is universal
- Mochon [2003] Mochon [2004] for universal quantum computing with topological superconductor. (This was slightly after freedman)
- Bonderson et al. [2008] for interferometry
- Bonderson [2009] for splitting of topological degeneracy
- Bonderson et al. [2008a] for measurement only tqc.
- Field and Simula [2018] Simula review
- Bonev et al. [2005] Original Compiling Fib anyons
- Hormozi et al. [2007] Quant compiling PRB
- Hormozi et al. [2009] Compiling RR states
- Simon et al. [2006] One mobile particle
- Nielsen and Chuang [2000] Nielsen and Chuang
- Brylinski and Brylinski [2002] Brylinski theorem
- Klinchkinov et al. [2014] Galois theory
- Maybe cite original Solovay and Kitaev article.
Exercises

Exercise 11.1 Ising Nonuniversality
The braiding matrices for Ising anyons are given by Eqs. 10.10 and 10.11. Demonstrate that any multiplication of these matrices, and their inverses, will only produce a finite number of possible results. Thus conclude that Ising anyons are not universal for quantum computation. Hint: write the braiding matrices as \( e^{i\alpha}U \), where \( U \) is unitary with unit determinant, i.e., is an element of SU(2). Then note that any SU(2) matrix can be thought of as a rotation \( \exp(i\hat{n}\cdot\sigma \theta/2) \) where here \( \theta \) is an angle of rotation \( \hat{n} \) is the axis of rotation and \( \sigma \) is the vector of Pauli spin matrices.

Exercise 11.2 Brute Force Search
Given the braid matrices for Fibonacci anyons in Eq. 10.6 and 10.7, write a computer program for brute-force searching braids up to length 10. Ignoring the noncomputational state \( |N\rangle \), and ignoring the overall phase as usual, determine the closest approximation to the Hadamard gate

\[
H = \frac{1}{\sqrt{2}} \begin{pmatrix} 1 & 1 \\ 1 & -1 \end{pmatrix}
\]

Partial Answer: A braid of length 10 exists with phase-invariant distance to target \( \text{dist} \approx 0.084 \).

Exercise 11.3 Scaling of Kitaev-Solovay Algorithm
Given the discussion just above Eq. 11.8, prove Eqs. 11.8 and 11.9.

Exercise 11.4 About the Injection Weave
One might wonder why we choose to work with an injection weave in Fig. 11.11 which moves the red strand from the far right at the bottom all the way to the far left on the top. Show that for three Fibonacci anyons, there does not exist any injection weave that moves the (red) strand from the far right on the bottom to the middle on the top, even up to an overall phase. I.e., show that no weave exists starting on the bottom far left ending in the middle on the top whose effect on the three dimensional Hilbert space is \( e^{i\phi}1_{3 \times 3} \) for any phase \( \phi \).

Exercise 11.5 Universal Weaving and the Injection Weave
Consider injection weaves as described in Fig. 11.11. Let us assume that we can construct an injection weave of arbitrary precision. Given such an (approximately) perfect injection weave show that for any number of anyons \( N > 3 \), a weave can be constructed that performs the same unitary operation on the Hilbert space as any given braid. A more general mathematical proof of the universality of weaving is also given in Simon et al. 2006.
Part III

Anyon Diagrammatics (in detail)
One of our objectives is to come up with some diagrammatic rules (somewhat analogous to those of the Kauffman bracket invariant) which will allow us to evaluate any diagram of world-lines (i.e., a labeled link, possibly now including diagrams where particles come together and fuse, or split apart) and get an output which is a complex number as desired in Fig. 12.1. In chapters 9-10 we have been putting together some of the necessary pieces for these diagrammatic rules. Here we will begin to formalize our diagrammatic algebra a bit more precisely$^2$. While we try to physically motivate all of our steps, in essence the rules of this chapter can be taken to be axioms of the diagrammatic algebra.

In this chapter we will focus only on planar diagrams — i.e., we do not allow lines to cross over and under each other forming braids. We can roughly think of such planar diagrams as being particles moving in 1+1 dimension. Since there are no over and under-crossings the only nontrivial possibility is that particles come together to fuse, or they split apart. An example of a planar fusion diagram is shown in Fig. 12.1. It is convenient to draw diagrams so that no lines are drawn exactly horizontally. The reader should be cautioned that there are several different normalizations of diagrams — two in particular that we will discuss. These two normalization conventions are useful in different contexts. We will start with a more “physics” oriented normalization in this chapter but we switch to a more topologically oriented normalization in chapter 14 and in later chapters.

We start by briefly reviewing some of the notions introduced in chapters 8-9. We assume a set of particle types $a, b, c, \ldots$ which we will draw as labeled lines with arrows in our diagrammatic algebra. This set of particles includes a unique identity or vacuum particle $I$, which may be drawn as a dotted line, or may not be drawn at all since it corresponds to the absence of any particles. Each particle type has a unique antiparticle denoted with an overbar (e.g., $\bar{a}$ for the antiparticle of $a$). As we discussed in section 8.1, if we reverse the arrow on a line we turn a particle into its antiparticle. If a particle is its own antiparticle we do not draw an arrow on its line.

Fusion rules are given by the matrices $N_{ab}^{cd}$, having the properties discussed in section 8.3. We will also assume a consistent$^3$ set of $F$-matrices as discussed in chapter 9.

---

$^1$This chapter through chapter 15 develop the diagrammatic algebra in some detail. Those who would like a brief and easier (albeit not as general) introduction to diagrammatic algebra should go straight to chapter 16.

$^2$Formally, some of the mathematical structure of planar diagrams was introduced in section 8.6. The rules we are defining in this chapter is known as a unitary fusion category to mathematicians. If various additional properties are satisfied other names may be used (spherical category, braided category, modular category, etc.).

$^3$The word “consistent” here means that the $F$-matrices satisfy the pentagon Eq. 9.3.

---

Fig. 12.1 A planar fusion diagram starting and ending at the vacuum.
12.1 Diagrams as Operators

If, like Fig. 12.1, a diagram starts at the bottom from the vacuum and ends at the top with the vacuum, we interpret that diagram to represent a complex number, or an amplitude. However, we will also consider diagrams that have “loose ends” (lines sticking off the top or bottom of the page) meaning that they may not begin or end with the vacuum. We can view these diagrams with loose ends as being a sub-diagram of a larger diagram that begins and ends in the vacuum. However, it is also useful to give such diagrams quantum mechanical meaning in their own right.

Our convention is that when we draw a diagram with world-lines that end pointing upwards we should view these particles as kets (independent of the direction of any arrow drawn on the world-line). If world-lines end pointing downwards, we mean them to be bras. Many diagrams will have world-lines that point both up and down, in which case we mean that the diagram has some particles that live in the vector space of kets and some in the dual (bra) space. Such diagrams can be interpreted as operators that take as input the lines coming in from the bottom and give as output the lines going out the top. The lines coming in from the bottom are thus in the bra part of the operator and the lines pointing out the top are the ket part of the operator. If we consider, for example, diagrams with \( M_{in} \) incoming lines from the bottom and \( M_{out} \) lines going out the top, we can write a general operator as

\[
\text{Operator} = \sum_{n,m,q} C_{n,m,q} |n, M_{out}; q\rangle \langle m, M_{in}; q| \quad (12.1)
\]

An example of such an operator is shown diagrammatically in Fig. 12.2 with two incoming and three outgoing lines. In Eq. (12.1) the states \( |n, M_{out}; q\rangle \) are an orthonormal complete set of states of \( M_{out} \) particles where all the particles together fuse to the quantum number \( q \); and similarly the states \( |m, M_{in}; q\rangle \) are an orthonormal complete set of states of \( M_{in} \) particles where all the particles together fuse to the quantum number \( q \). The value of the coefficients \( C_{n,m,q} \) depend on the details of the diagram being considered. The fact that the operator is necessarily diagonal in the variable \( q \) means that the total quantum number of all of the incoming particles must be the same as the total quantum number of all the outgoing particles (i.e., they fuse to the same overall charge). This conservation of overall quantum number is a reflection of the locality principle of section 8.2.

Generally in a diagram, lines will be labeled with particle types and (if the particle is not self-dual) arrows. We have not labeled the incoming and outgoing lines in Fig. 12.2 with the assumption that these labels and arrows occur inside the hidden box. However, it is sometimes useful to reinstate these labels as in Fig. 12.3. As we will discuss in more detail in section 12.2.1 a label restricts the quantum number of the corresponding line...
Diagrams as Operators

\[ \sum_{n,m,q} C^{(1)}_{n,m,q} |n, 3; q\rangle\langle m, 2; q| \]

**Fig. 12.2** Diagram 1, representing an arbitrary diagram (or linear combination, i.e., weighted sum, of diagrams), is understood as part of a larger diagram, and is interpreted as an operator. Incoming lines from the bottom correspond to bras and outgoing lines towards the top correspond to kets. The states \(|n, 3; q\rangle\) are a complete set of states for 3 particles where all the particles together fuse to the quantum number \(q\). Similarly, the states \(|m, 2; q\rangle\) are a complete set of states for 2 particles where all the particles together fuse to the quantum number \(q\). The superscript on \(C^{(1)}_{n,m,q}\) indicates that these constants correspond to the particular “diagram 1” in the box. The total quantum number \(q\) of all the particles is conserved by the operator due to the locality principle from section 8.2.

\[ \sum_{n,m,q} C^{(2)}_{n,m,q} |n, 3; q\rangle\langle m, 2; q| \]

**Fig. 12.3** In a figure with labeled incoming and outgoing lines, the quantum numbers on these lines are fixed, as compared to Fig. 12.2 where the diagram may have a superposition of quantum numbers on the external lines.

We now introduce an important diagrammatic principle:

**Hermitian Conjugation:** Reflecting a diagram around a horizontal axis and then reversing the direction of all arrows implements Hermitian conjugation.\(^8\)

For example, reflecting Fig. 12.3 and then reversing the arrows on all lines results in the Hermitian conjugate diagram

\[ \sum_{n,m,q} [C^{(2)}_{n,m,q}]^* |m, 2; q\rangle\langle n, 3; q| \]

**Fig. 12.4** Flipping the diagram in Fig. 12.3 results in the Hermitian conjugate. The coefficients \(C^{(2)}_{n,m,q}\) in Fig. 12.3 are complex conjugated to obtain \([C^{(2)}_{n,m,q}]^*\) here.

It is crucial that when we turn a bra into a ket (reflecting the diagram and then reversing the arrows), down-pointing arrows remain down-pointing and up-pointing arrows remain up-pointing (Note, for example, that the arrow on \(a\) is pointing up both in Fig. 12.3 and 12.4).
Diagrams which start from the vacuum at the bottom are an important special case. When there are no incoming lines at the bottom of a diagram the expression become

\[ |\text{ket} \rangle = \sum_{n,q} C_{n,\emptyset,q} |n, M_{out}; q \rangle \]  \hspace{2cm} (12.2)

which we can also interpret as an “operator” which accepts the vacuum as an input at the bottom and gives a ket as an output at the top. The symbol \( \emptyset \) here means that the \( m \) index used in Fig. 12.2 and Eq. 12.1 is just the empty set (nothing summed over), or equivalently that the diagram starts from the vacuum. An example of such a diagram is shown in Fig. 12.5.

\[ \sum_{n,q} C_{n,\emptyset,q} (3) |n, 3; q \rangle \]  \hspace{2cm} (diagram 3)

Fig. 12.5 A diagram with no incoming lines at the bottom is interpreted as a ket.

Similarly we can consider diagrams which end in the vacuum at the top. When there are no outgoing lines at the top of a diagram we have

\[ \langle \text{bra} | = \sum_{m,q} C_{\emptyset,m,q} \langle m, M_{in}; q | \]  \hspace{2cm} (12.3)

which is an operator that accepts a ket as an input and gives a complex number as an output. An example of such a diagram is shown in Fig. 12.6.

\[ \sum_{m,q} C_{\emptyset,m,q} (4) \langle m, 3; q | \]  \hspace{2cm} (diagram 4)

Fig. 12.6 A diagram with no outgoing lines at the top is interpreted as a bra.

If diagram 3 happens to be the reflection of diagram 4 around a horizontal axis with all arrows reversed, then these two diagrams are hermitian conjugates of each other and \( C_{n,\emptyset,q}^{(4)} = [C_{\emptyset,n,q}^{(3)}]^* \).

12.1.1 Stacking operators

Stacking operators on top of each other contracts bras with kets in the natural way\(^9\). For example, if we define the operator, diagram 5, as in Fig. 12.7 we can then stack diagram 5 (Fig. 12.7) on top of diagram 1 (Fig. 12.2) to obtain Fig. 12.8 The resultant operator, diagram 6, on the right is given by

\[ \sum_{n,q} C_{n,\emptyset,q}^{(3)} \langle n, 3; q | \]

---

\(^9\)The observant reader will see similarities between this stacking procedure and the stacking of manifolds with boundary discussed in chapter 7. These similarities are not a coincidence!
12.2 Basis of States

In our definition of an operator (Eq. 12.1) we invoked the existence of a complete orthonormal basis of states $|n, M; q\rangle$ for $M$ particles having total quantum number $q$. We now would like to specify some details of this basis.
Planar Diagrams

\[
\begin{align*}
\text{diagram 4} & = \left( \sum_{m,q} C^{(4)}_{m,q} |m,3; q\rangle \langle m,\emptyset,3; q| \right) \left( \sum_{n,q} C^{(3)}_{n,q} |n,\emptyset,3; q\rangle \langle n,\emptyset,3; q| \right) \\
& = \sum_{m,q} C^{(4)}_{m,q} C^{(3)}_{m,q}
\end{align*}
\]

Fig. 12.9 Stacking a bra operator on top of a ket operator generates a scalar. We have used orthonormality of the kets \(|n,3; q\rangle\) on the right hand side.

12.2.1 One Particle

We begin by considering a single particle at a time. For a single particle, an orthogonal complete basis is given by the different particle types \(1010\). In keeping with the notation of Fig. 12.2 the state \(|a\rangle\) should be notated \(|a,1; a\rangle\) to indicate a single line, but here we use just \(|a\rangle\) for simplicity.

\(|a\rangle\) (including the vacuum \(|\emptyset\rangle\)). We denote a projector onto a particular particle type as a simple labeled straight line as shown in Fig. 12.10. The vacuum can be drawn as a dotted line, or may not be drawn at all.

\[|a\rangle = |a\rangle \langle a|\]

Fig. 12.10 A labeled straight line is just an projector onto the particle type.

Since the different particle types are assumed orthonormal \(|a|b\rangle = \delta_{ab}\), applying two such projectors in a row diagrammatically gives the identity shown in Fig. 12.11.

\[
\begin{align*}
& b \\
\text{diagram 11} \quad a \\
& = \langle b|a\rangle \langle a| = \delta_{ab} |a\rangle \langle a| = \delta_{ab} \quad a
\end{align*}
\]

Fig. 12.11 Orthogonality of projection operators.

This identity exemplifies the more general rule shown in Fig. 12.12 which also agrees with the fact that the operators in Eq. 12.1 are diagonal in the overall quantum number \(q\). Again this is simply a reflection of the locality, or no-transmutation, principle\(^\text{11}\) of section 8.2 (See in particular Fig. 8.7).

\[
\begin{align*}
& b \\
\text{anything} \\
& = 0 \quad \text{unless } a = b \\
& a
\end{align*}
\]

Fig. 12.12 The locality, or no-transmutation, principle as in Fig. 8.7.

Since we assume the set of particle types is complete, the identity operator is given by the sum over all particle types as in Fig. 12.13 where the sum includes the vacuum particle. We represent the identity operator on the right in Fig. 12.13 as a straight unlabeled line. This is convenient since it allows us to extend labeled lines by appending unlabeled lines.
\[ \sum_a a = \sum_a |a \rangle \langle a| = \text{identity} = \]

**Fig. 12.13** The completeness relation for single lines.

### 12.2 Two Particles

Let us now move on to the case of two particles. As discussed in chapter 5 to fully describe the state of two particles, we need to give the quantum number (particle type) of each particle and the fusion channel between the two particles. We thus draw the state of two anyons with a vertex diagram as shown in Fig. 12.14

![Diagram](image)

**Fig. 12.14** Particles \( a \) and \( b \) have fusion channel \( c \).

The notation of the ket \( |a, b; c\rangle \) means that the total quantum number of particles \( a \) and \( b \) is \( c \) (or \( a \) and \( b \) fuse to \( c \)). If \( N_{ab}^c = 0 \), i.e., if the diagram is a disallowed fusion, then the value of the diagram is zero. The set of states \( |a, b; c\rangle \) for all possible \( a, b, c \) is assumed to form an orthonormal complete set of states for two anyons. Note in particular that for \( a \neq b \) the ket \( |a, b; c\rangle \) is orthogonal to \( |b, a; c\rangle \) — i.e., in our planar diagram algebra, it matters which particle is to the left and which is to the right.

The Hermitian conjugate of the vertex ket Fig. 12.14, the corresponding bra, is shown in Fig. 12.15

![Diagram](image)

**Fig. 12.15** This is the bra which is the Hermitian conjugate of the ket in Fig. 12.14.

The fact that the diagram for the bra looks like the ket upside-down is in accordance with our general principle of Hermitian conjugation introduced in section 12.1 (See the discussion near Fig. 12.4).

To take inner products between a bra (like Fig. 12.14) and a ket (like Fig. 12.15) we simply stack the bra on top of the ket, in accordance with section 12.1.1 to produce the diagram shown in Fig. 12.16

The fact that we obtain delta functions on the right is equivalent to the statement that the kets \( |a, b; c\rangle \) form an orthonormal set. The normalization of Fig. 12.16 (i.e, that one gets Kronecker deltas on the right and no numerical constants) is our *physics normalization*. This normalization will be changed in chapter 14.

Note that the first two delta functions \( \delta_{aa'} \) and \( \delta_{bb'} \) in Fig. 12.16 can be interpreted as a result of Fig. 12.14 (the lines are angled instead of vertical).
Planar Diagrams

\[ \delta_{aa'} \delta_{bb'} \delta_{cc'} \]

Fig. 12.16 The inner product between the bra in Fig. 12.14 and a ket in Fig. 12.15. This gives Kronecker deltas on the right given the physics normalization we are using in this chapter. The normalization will be changed in chapter 14.

of vertical, but this does not change their meaning). As a result, the diagram of Fig. 12.16 is often written in the simplified form shown in Fig. 14.7

\[ \delta_{cc'} \]

Fig. 12.17 A simplified version of the inner product in Fig. 12.16. This gives a Kronecker delta on the right in the physics normalization we are using in this chapter. As always if the vertex is disallowed by fusion (if \( a \times b \) does not include \( c \)) then the value of the diagram is zero. The normalization will be changed in chapter 14.

The fact that \( c \) must equal \( c' \) in Figs. 12.16 and 12.17 is consistent with the no-transmutation principle Fig. 12.12.

The principle of orthonormality of vertices implies the useful result that a loop, as shown in Fig. 12.18 is given the value of unity (This is just Fig. 12.17 where we have set \( c = c' = I \) and not drawn the identity line). At the risk of being repetitive we once again note that we will change this normalization in chapter 14 below and in later chapters, although it is correct for this section.

\[ \langle \text{state} | \text{state} \rangle = 1 \quad \text{Physics Normalization} \]

Fig. 12.18 The orthonormality of trees implies a particle loop gets a value of 1 if we are using physics normalization.

Since the vertex diagrams \( |a, b, c\rangle \) from Fig. 12.14 form a complete set of states for the two particles, we can construct an identity operator for two strands as shown in Fig. 12.19.

We can derive a useful identity from Fig. 12.19 by applying projectors \( |x\rangle \langle x| \) and \( |y\rangle \langle y| \) to left and right strings respectively to obtain Fig. 12.20.
\[ \sum_{a,b,c} |a, b; c \rangle \langle a, b; c| = \sum_{a, b, c} |a, b; c \rangle \langle a, b; c| \]

**Fig. 12.19** Insertion of a complete set of states. This figure uses physics normalized diagrams. The normalization will be changed in chapter 14.

\[ \sum_{c} |x, y; c \rangle \langle x, y; c| = \sum_{c} |x, y; c \rangle \langle x, y; c| \]

**Fig. 12.20** Insertion of a complete set of states, with fixed quantum numbers \( x \) and \( y \) on both ends. This figure uses physics normalized diagrams. The normalization will be changed in chapter 14.

An arbitrary operator with two incoming and two outgoing lines can be written as in Fig. 12.21 where the coefficients \( C(x, y), (a, b), c \) are arbitrary (depending on what operator we want to define).

\[ \sum_{a, b, c, x, y} C(x, y), (a, b), c = \sum_{a, b, c, x, y} C(x, y), (a, b), c |x, y; c \rangle \langle a, b; c| \]

**Fig. 12.21** An arbitrary operator with two incoming and two outgoing lines. The coefficients \( C \) are arbitrary.

### 12.2.3 Three Particles

We can continue on and consider states of three particles. All the same principles apply here. As discussed in chapter 8, we can write an orthonormal complete set of states for three particles as a fusion tree\(^{18,19}\) as in Fig. 12.22.

\[ |(a, b), c, d; e \rangle = |(a, b); d \rangle \otimes |d, c; e \rangle \]

**Fig. 12.22** An orthonormal set of states for three particles can be described as a fusion tree. We have used two different notations on the right. The latter notation matches that of section 8.6.

If either \( N_{ab}^d = 0 \) or \( N_{dc}^e = 0 \) then the corresponding fusion is disallowed and the value of the diagram is zero\(^{20}\). The corresponding kets are obtained using the Hermitian conjugation rule of flipping the diagram and reversing arrows as shown in Fig. 12.23.

\[ \text{In cases where there are fusion multiplicities } N_{ab}^d > 1 \text{ or } N_{dc}^e > 1 \text{ then we must place an additional index } \mu \text{ or } \lambda \text{ at the corresponding index. See for example, section 8.5.3.} \]

\[ \text{As mentioned in the above note 13, although we write this as a ket, it is really an operator, and to emphasize this we might write something like } |(a, b), c, d; e \rangle |c \rangle \text{ instead.} \]

\[ \text{This is already implied by looking at the individual vertices and considering the rules of a single vertex as in Fig. 12.13.} \]
The inner product of such states is given by stacking the bra on top of the ket as in Fig. 12.24.

Note that in Fig. 12.24 one can focus one's attention on the left branches (with \(d', a, b, a', b', d'\)), which look exactly like Fig. 12.16 thus immediately obtaining \(\delta_{aa'}\delta_{bb'}\delta_{cc'}\delta_{dd'}\delta_{ee'}\) and replacing the small diamond on the left branch with a single \(d\)-line. The remaining figure then looks exactly like the figure 12.16 and gives us the delta functions \(\delta_{cc'}\delta_{ee'}\) as well. Thus the orthonormality of these tree states is not a separate assumption but can be derived from the orthonormality of two particle states that we used in section 12.2.2.

The completeness of this set of states similarly can be expressed with diagrams as shown in Fig. 12.25.

Once again we can derive this completeness relation from what we know about the two-particle case. We can start in the very center of Fig. 12.25, considering the lines \(d, c, e, d', c'\), and apply the completeness relation Fig. 12.20 This splits off the \(c\)-line to the right which, summed over its index gives a single unlabeled line on the right as in Fig. 12.13.
The remaining diagram on the left (with lines \(a, b, d, a', b'\)) is of the form of Fig. 12.19 which summed over gives two unlabeled lines. Thus the completeness relation for three particles is not an independent assumption but follows from the completeness of the one and two particle cases.

One can use these basis states to build arbitrary operators with three particle states. Just as an example, in Fig. 12.26 we show the most general form of an operator that takes two particles as an input and gives three particles as an output.

\[
\sum_{a, c, e, x, y, z, w} C_{x,y,z,w}^{a,c,e} |(x, y) z; w; e\rangle \langle a, b; e|
\]

The coefficients \(C\) are arbitrary.

### 12.2.4 \(F\)-Matrices Again

In defining our three particle states in Fig. 12.22 we have fused the two particles \(a\) and \(b\) on the left first to form \(d\) and then fused \(d\) with \(c\) to form \(e\). (Our notation to indicate this is \(|(a, b) c; d; e\rangle\). However, we could have chosen to fuse the particles in a different order to form a different tree as shown in Fig. 12.27. Here \(b\) and \(c\) fuse together to form \(f\) and then \(a\) and \(f\) fuse together to form \(e\). We notate this state as \(|a, (b, c) f; e\rangle\).

\[
\sum_{a, c, e, x, y, z, w} C_{x,y,z,w}^{a,c,e} |(x, y) z; w; e\rangle \langle a, f; e| = |a, (b, c) f; e\rangle = |a, f; e\rangle \otimes |(b, c); f\rangle
\]

The set of states defined by the fusion trees in Fig. 12.27 also form a perfectly good (but different) complete orthonormal basis of states for three particles. For example, we have the orthogonality relation shown in Fig. 12.28 (compare Fig. 12.24). As described in detail in chapter 9, if we draw trees with different branching structure, we are describing the same Hilbert space, but in a different basis — the basis change being given by a unitary \(F\)-matrix transform as shown in Fig. 12.29 (see also discussion of Fig. 9.1). Similarly we have the relation between the Hermitian conjugate states as shown in Fig. 12.30. Note that because the \(F\)-matrix is unitary in its
two outside indices (d and f in Fig. 12.29) we have

\[ [F_{abc}^{e}]_{df}^* = ([F_{abc}^{e}]^*)_{fd} = ([F_{abc}^{e}]^{-1})_{fd} \]

12.2.5 More Particles

The principles we have developed for one-, two-, and three-particle states are easily extended to greater numbers of particles. Each shape of fusion tree defines a different orthonormal complete set of states. For example, with four particles, we might choose the tree shape shown in the left of Fig. 12.31 or we might choose the tree shape shown in the right of Fig. 12.31. Either one of these makes a perfectly good orthonormal basis for four particles — and these two bases are related to each other by \( F \)-matrices as discussed in chapter 9.

The left-hand tree structure in Fig. 12.31, with all of the particles on top branching from a single line going from top left to bottom right, is sometimes known as the “standard basis.”

One can use \( F \)-moves to evaluate more complicated diagrams. An example of this is shown in Fig. 12.32.
12.3 Causal Isotopy

Keeping with the idea of diagrams that are planar (no over- and under-crossings), we now consider how we may deform these diagrams. When we discussed the Kauffman bracket invariant we were allowed to freely deform any diagram as long as we did not cut any strands. This property is known as isotopy invariance. Analogously, if a planar diagram retains the same value for any deformation that does not involve cutting strands or crossing them over each other, we say the theory has full planar isotopy invariance. Examples of this are shown in Fig. 12.33.

We need to ask how much topological invariance we should really expect from our physical theories. In the mathematical world of TQFTs and knot invariants, it is fine to assume that all directions are equivalent, and we can freely distort a line travelling in the $x$ direction (horizontally) on the page to a line travelling in the $t$ direction (vertically). However, in that case we had regular isotopy invariance meaning that we can deform knots freely in 3D as long as we treated the strands as ribbons. See sections 2.2.1 and 2.6.1.
For a theory with full planar isotopy invariance, these two diagrams should evaluate to the same result. Full planar isotopy invariance allows us to distort the diagram in any way as long as we do not cut any strands or cross lines through each other.

in real physical systems, generically the time direction might need to be treated differently from the space directions. In this section we will discuss topologically theories that allow deformation in space, but without allowing one to freely exchange the time and space directions. In particular some amount of causality might be demanded.

In chapter 16 we will consider a subset of theories which have a much higher level of topological invariance, known as regular isotopy invariance, which allows us to freely distort diagrams in either the space or time direction and further allow us to interchange the two.

In this chapter through chapter 15 we do not assume regular isotopy invariance (or full planar isotopy in the case of planar diagrams) but rather assume only what we call causal isotopy. Here we allow deformation of space-time diagrams so long as we do not change the time-direction orientation of any lines or vertices. In other words, the path of a particle that is moving forward in time should not be distorted such that it is moving backwards in time (and vice-versa, a particle moving backwards should not be distorted so that it is moving forwards) — but other than this constraint, any smooth deformation is allowed. Two examples of deformations that are allowed under causal isotopy are shown in Fig. 12.34.

Certain deformation of diagrams are not allowed by causal isotopy.
Two examples of such disallowed deformations are given in Fig. 12.35. On the left of the figure we see a particle which turns around in time. This need not be the same as the particle moving straight in time as it involves a particle creation event and a particle annihilation event. On the right of Fig. 12.35 a vertex is altered so instead of an $a$ particle going out of the vertex, a $\bar{a}$ particle goes in. In this case we must have a $a$ with a annihilation event in the far right diagram that does not exist in the simpler diagram where $a$ and $c$ directly fuse to $b$. Thus these two diagrams do not necessarily evaluate to the same result. (Although in some cases, such as in chapter 16, one may have a simple theory for which the transformations showed in Fig. 12.35 are allowed, such theories are not generic.)

12.4 Summary of Planar Diagram Rules in Physics Normalization

With the principles we have now discussed we should be able to evaluate any planar diagram — taking a space-time process which starts and ends in the vacuum and turning it into an amplitude (i.e, a complex number). The same principles can be used to simplify operators such as Eq. 12.2.

Here are a summary of the important rules we have learned for diagram evaluation:

1. One is free to continuously deform a diagram consistent with causal isotopy as described in section 12.3. That is, particles must not change their direction in time due to the deformation.
2. One is free to add or remove lines from a diagram if they are labeled with the identity or vacuum ($I$). See the example in Fig. 12.36.
3. Reversing the arrow on a line turns a particle into its antiparticle (See Fig. 8.4).
4. Regions must maintain their quantum number locally as in Fig. 12.12.
   In particular this means that a line must maintain its quantum number unless it fuses with another line, or splits.
5. Splitting and fusion vertices are allowed for fusion multiplicities $N_{ab}^c > 0$ (See section 5.3). This includes particle creation and
annihilation as a special case where a particle-antiparticle pair fuse to the vacuum or split from the vacuum (An example is shown in Fig. 12.36).

(6) Hermitian conjugation is given by reflection of a diagram around a horizontal line along with flipping the direction of arrows (See Fig. 12.4 or for example, Fig. 12.23)

(7) One can use $F$-moves to change the structure of fusion trees in order to simplify. For example, in Fig. 12.32

(8) Once one reduces a diagram into tree structures that have the same branching in the upper and lower half (as on the right of Fig. 12.32) we can use the orthonormality of trees to complete the evaluation. In cases where the diagram starts and ends in the vacuum this reduces the diagram to a complex number (See, for example, in Fig. 12.32). More generally operator diagrams can be reduced to simple forms analogous to Fig. 12.26.

With these principles (and given an $F$-matrix as input information — which will depend on the particular physical system we are considering) it is possible to fully evaluate any planar diagram, starting and ending in the vacuum, into a complex number. While there may be many strategies to use these rules to reduce a complicated diagram to a single complex number, the final result is independent of the order in which we apply the rules\textsuperscript{24}.

The mathematical structure we have defined thus far (our Hilbert space and $F$-matrices) is known as a “unitary fusion category”. There is more structure to be uncovered in further chapters that follows from what we have defined so far, and there are many special cases to be discussed. In addition note that here we have only described planar diagrams, so we have not yet described 2+1 dimensional theories — in order to describe these, we will have to include braiding rules for our diagrams we will add in chapter 13.

12.4.1 A Simple Example

As a simple example, let us try to evaluate the diagram shown on the far left of Fig. 12.38. We first work on a small part of the diagram as shown in Fig. 12.37 (Note that this is the same as the far right of Fig. 12.35).

\[ a \xrightarrow{\ b \quad \ c \quad} [F^a_{\ b \ c}] I_c \]

\[ a \xrightarrow{\ b \quad \ c \quad} [F^a_{\ b \ c}] I_c \]

**Fig. 12.37** To evaluate the diagram on the left, the vacuum line is inserted and an $F$-move is made. The bubble is then removed with Fig. 12.16. These diagrams use physics normalization. We will re-examine this diagram using a different normalization in section 14.3.
12.5 Appendix: Higher Fusion Multiplicities

When we have a theory with higher fusion multiplicities (i.e., $N_{ab}^c > 1$ for at least one fusion channel), then the vertices must be given indices as well as lines having indices, and tree states are orthogonal in these indices as well. For example, we would need to modify Figs. 12.17 and 12.20 to the form shown in Figs. 12.39 and 12.40. See also the discussion of the $F$-matrix with higher fusion multiplicities in section 9.5.3.

![Fig. 12.39](image)

The bubble diagram when there are fusion multiplicities. This diagram is a result of the orthonormality of tree diagrams. The variables at the vertices must match in order for the result to be nonzero. This diagram is drawn in the physics normalization. We will change the normalization in chapter 16.

![Fig. 12.40](image)

Insertion of a complete set of states. When there are fusion multiplicities, these must be summed over as well $\mu \in N_{ab}^c$. This diagram is drawn in the physics normalization. We will change the normalization in chapter 16.

**Chapter Summary**

- This is an item
Further Reading

This is some reading.

Exercises

Exercise 12.1 Evaluating diagrams with $F$-matrices
Evaluate the following diagram, writing the result in terms of $F$'s.

Exercise 12.2 Locality Principle
Show that the locality principle (Fig. 12.12) is derivable from our other rules for evaluating diagrams, and is not therefore an independent assumption.
Braiding Diagrams

In chapters 8, 9, and 12 we focused on planar diagrams. These diagrams can be thought of as describing the physics of objects that live in 1+1 dimension. More to the point, the nontrivial physics we discovered is really just a reflection of the nontrivial structure of the Hilbert spaces we are working with.

Here we extend our diagrammatic rules to the 2+1 dimensional world. In particular we want to describe what happens when we braid world lines. In chapter 10 we started to discuss braiding of identical particles and we continue that discussion here.

13.1 Three Dimensional Diagrams

We begin by generalizing the concept of a diagram that we developed in chapters 9, 10. The diagrams we want to consider now allow over- and under-crossings of lines as in Fig. 13.1. We will end up with a set of rules that are conceptually similar to the knot-invariants we discussed way back in chapter 2 — starting with a picture of a generalized knot (like Fig. 13.1), we reduce it to an output number. The generalization here is that the lines have labels \((a, b, c, \ldots)\) and lines can fuse with each other in addition to crossing over and under each other.

We should be somewhat cautious here that whereas when we considered the Kauffman bracket invariant, we had regular isotopy invariance — meaning that, treating strands as ribbons, any deformation of the diagram was allowed as long as we did not cut any strands. In contrast here (while we should still treat strands as ribbons) not all deformations are allowed. In general we will only have the same type of causal isotopy as described in section 12.3 (that is, we cannot freely deform a particle line going forward in time to one that goes backwards in time). Of course there do exist anyon theories with a higher level of isotopy invariance (regular isotopy), which we will discuss in chapter 16 but we should realize that these are not generic.

Our rules for evaluating diagrams with over- and under-crossings will be a consistent extension of a set of rules for evaluating planar diagrams. Our next task is to consider how we handle over- and under-crossings. With this information, used in conjunction with the rules we have already developed for planar algebras, we will be able to evaluate any diagram in 2+1 dimensions.

1This chapter continues the development of the diagrammatic algebra in some detail. Those who would like a brief and easier (albeit not as general) introduction to diagrammatic algebra should go straight to chapter 16.

Fig. 13.1 A diagram with over- and under-crossings representing a process in 2+1 dimensions.

2In mathematical language, the rules introduced in this chapter give additional structure to a unitary fusion category to make it a unitary braided fusion category, or unitary ribbon fusion category (these notions are equivalent).
13.2 Braiding Non-Identical Particles

We now turn to ask what happens if we exchange two different particle species, say $a$ and $b$. We might be tempted to do something similar to Figs. 10.1 and 10.2 — that is, we define a state with two particles in a given fusion channel then we exchange the two particles and determine the phase accumulated in this process. However, such a scheme cannot work in the case of non-identical particles. The reason this fails is that when the two particles are not identical the initial and final states are fundamentally different and cannot be related to each other by just a phase — for example, the initial state for Fig. 13.2 has $a$ to the left of $b$ whereas the final state has $a$ to the right of $b$.

Nonetheless, the $R$-matrix can still be precisely defined even when we are braiding non-identical particles. Diagrammatically we define the $R$-matrix as shown in Fig. 13.3. On the right of this figure, the particles $b$ and $a$ come from $c$, with $a$ going off to the left and $b$ to the right. In the left of the figure, the two particles are moved away from each other, $b$ to the right and $a$ to the left, before they are braided around each other. The key here is that in both cases, the final state of the system has $b$ on the left and $a$ on the right, and in both cases the two particles fuse to a quantum number $c$, so that the two processes can be compared to each other and differ from each other only by a phase, which we define\(^3\) to be $R^{ab}_{c}$.

![Fig. 13.3](image)

In a unitary theory the $R$-matrix is always just a complex phase. Note that $R^{ab}_{c}$ is not defined if $a$ and $b$ are not allowed to fuse to $c$ (i.e., if $N^{c}_{ab} = 0$). Further, note that braiding anything with the identity (vacuum) particle should be trivial,

$$R^{Ia}_{a} = R^{aI}_{a} = 1.$$

A full braid (two exchanges in the same direction) of two particles $a$ and $b$ fusing to $c$ is given by $R^{ba}_{c} R^{ab}_{c}$ as shown in Fig. 13.4. Note that in the representation on the far left of the figure, in both the initial and final configurations of particles, the $b$ particle is to the left of the $a$ particle meaning that we can understand this process as simply incurring an Aharonov-Bohm-like phase dependent on the fusion channel.

![Fig. 13.4](image)

Similarly we have the inverse braid shown in Fig. 13.5. Note carefully that the upper legs are labeled in the opposite order comparing Eq. 13.3 to 13.5. The reason for this is that after a single exchange the order of the two legs at the vertex is switched, so in the

\(^3\)The notation we use matches that of Bonderson [2007].
13.2 Braiding Non-Identical Particles

13.2.1 Summary of Rules for Evaluating any 2+1 D Diagram with Physics Normalization

The rules for evaluating any diagram in 2+1 dimensions (working with physics normalization of diagrams) are thus a very simple extension of the rules presented in section 12.4. We simply add two more rules...
Fig. 13.7 A generic crossing can be reduced to a planar diagram using the $R$-matrix. In the second step a complete set of particles $c$ is inserted as in Fig. 12.20. Note this figure uses physics normalization.

Fig. 13.8 The inverse crossing. This figure uses physics normalization.

(1) We are allowed to use $R$-moves as in Fig. 13.3 and 13.5. In particular, this allows resolving of crossings by using Fig. 13.7 and 13.8.

(2) Once any diagram is reduced to a planar diagram, we can use the rules of section 12.4.

As with the case of planar diagram, there is some degree of deformation of diagrams (causal isotopy, see section 12.3) which is freely allowed. Here again the rules are similar: any deformation that does not involve cutting lines, or changing the time-direction of motion is allowed. Without introducing new assumptions, natural moves such as those shown in Fig. 13.9 and 13.10 can be derived (See exercise 13.4). These are nothing more than the Reidermeister Type II and III moves introduced in section 2.6.1 although here the strands carry labels.

13.3 The Hexagon

Using $R$-moves and $F$-moves any 2+1 D diagram (starting and ending in the vacuum) can be reduced to a complex amplitude. One might worry if the rules we have listed for evaluation of diagrams are self-consistent: i.e., does it matter in which order we apply the rules? Will we always obtain the same complex amplitude result? Indeed, given

Fig. 13.9 These moves, Reidermeister Type-II moves, are allowed in any anyon theory. See section 2.6.1
an $F$-matrix, only certain sets of (physically acceptable) $R$-matrices will have the property that the diagrammatic rules give a unique final result. In fact, it is even possible that for a given set of $F$-matrices that satisfy the pentagon, there may not even exist a set of consistent $R$-matrices!

When we discussed planar diagrams in chapter 12, the pentagon equation guaranteed self-consistency of $F$-matrices. Now, given some $F$-matrices that satisfy the pentagon equations, the consistency equations for $R$-matrices are known as the hexagon equations and are shown diagrammatically in Fig. 13.11.

The hexagon equations in graphical form.

In equations the hexagon conditions can be expressed as

\begin{align}
R_{ca}^e [F_{ab}^{cde}]_{eg} R_{gb}^{ec} & = \sum_f [F_{d}^{fabc}]_{ef} R_{df}^{ef} [F_{d}^{gabc}]_{fg} \\
[R_{d}^{e}]^{-1} [F_{d}^{abc}]_{eg} [R_{d}^{g}]^{-1} & = \sum_f [F_{d}^{fabc}]_{ef} [R_{d}^{f}]^{-1} [F_{d}^{gabc}]_{fg}
\end{align}

The top equation is the left diagram whereas the lower equation is the right diagram in Fig. 13.11. The left hand side of the equation corresponds to the upper path, whereas the right hand side of the equation corresponds to the lower path.

The structure we have now defined — a consistent set of (unitary) $F$ and $R$-matrices satisfying the pentagon and hexagon equations, is known as a unitary braided tensor category. All 2+1 D anyon theories must be of this form.

\^This is also sometimes known as a unitary ribbon tensor category due to the fact that Eq. 15.3 holds, which is always true for unitary theories with braidings. The unitary braided tensor category is also sometimes known as a premodular category.
Given a set of fusion rules, the pentagon and hexagon equation are very very strong constraints on the possible $F$- and $R$-matrices that can result. We mentioned in section 6.3 that given a set of fusion rules there are only a finite number of solutions of the pentagon. Similarly, once we have a solution of the pentagon, i.e., once the $F$-matrices are fixed, there are only a finite number of possible solutions of the hexagon equations. Both of these statements are known as rigidity. In particular this means that if you have an anyon theory (a solution of the pentagon and hexagon) it is not possible to perturb this solution a small amount and get another solution\(^5\).

With simple fusion rules, such as Fibonacci (as we saw in exercise 9.4) the fusion rules completely determine the $F$-matrices of the theory. In this particular case, there are exactly two consistent solutions to the Hexagon equations, corresponding to the left and right handed types of Fibonacci anyons (See Eq. 10.2 and exercise 13.1).

### 13.4 \(R\)-matrix Odds and Ends

#### 13.4.1 Appendix: Gauge Transforms and \(R\)

As in section 9.4 one can gauge make gauge transformations on the vertices of a theory. Given the transform shown in Fig. 9.8,

\[
\begin{array}{c}
\begin{array}{c}
\text{a} \\
\text{b}
\end{array}
\end{array}
\begin{array}{c}
\begin{array}{c}
\text{c}
\end{array}
\end{array}
\xrightarrow{\text{gauge transform}}
\begin{array}{c}
\begin{array}{c}
\text{a} \\
\text{b}
\end{array}
\end{array}
\begin{array}{c}
\begin{array}{c}
\text{c}
\end{array}
\end{array}
\]

Fig. 13.12 We have the freedom to make a gauge transform of a vertex by multiplying by a phase $u_{ab}^c$. The tilde on the right notates that the vertex is in the tilde gauge.

the $R$-matrix transforms as

\[
\tilde{R}_{c}^{ab} = u_{ab}^c R_{c}^{ab}
\]

Note that $R_{c}^{aa}$ is gauge invariant, as is the full braid $R_{c}^{ab} R_{c}^{ba}$ in Fig. 13.4

#### 13.4.2 Product Theories

Given two anyon theories $T$ and $t$, we can construct the product theory $T \times t$ as in section 8.3. If the theory $T$ has consistent $R$-matrices $R_{c}^{AB}$ and the theory $t$ has consistent $R$-matrices $R_{c}^{ab}$ (“consistent” here means there are $F$-matrices that satisfy the pentagon relation and the $F$’s and $R$’s satisfy the hexagon relations), then the product theory has a consistent $R$-matrices

\[
R_{(C,c)}^{(A,a)(B,b)} = R_{c}^{AB} R_{c}^{ab}
\]
Again, the point here is that in a product theory, the two constituent theories don’t “see” each other at all.

13.4.3 Appendix: Higher fusion multiplicities

When we have a theory with higher fusion multiplicities (i.e., $N^c_{ab} > 1$ for at least one fusion channel), then the vertices must be given indices as well as lines having indices as in section 12.5. In this case the $R$-matrix carries vertex indices as well, and is a unitary matrix with respect to these indices.

$$ R_{\mu\nu}^{ab} = \sum_{\nu} \left[R_{c}^{ab}\right]_{\mu\nu} $$

Fig. 13.13 Definition of $R$-matrix when there are higher fusion multiplicities. Here the vertices carry labels, and $R$ is a unitary matrix with respect to these labels.

Under a gauge transformation, as in Fig. 9.10 the $R$-matrix transforms as

$$ \tilde{R}_{c}^{ab} = \sum_{\mu,\nu} \left([u_{ab}]^{-1}\right)_{\mu,\mu'}\left[R_{c}^{ab}\right]_{\mu\nu}\left[u_{ab}\right]_{\nu,\nu'} \quad (13.4) $$

Exercises

Exercise 13.1 Fibonacci Hexagon

Once $F$-matrices are defined for a TQFT, consistency of the $R$-matrix is enforced by the so-called hexagon equations as shown in the figure diagramatically by Fig. 13.11 or the Fibonacci anyon theory, once the $F$-matrix is fixed as in Eq. 9.3 the $R$-matrices are defined up to complex conjugation (i.e., there is a right and left handed Fibonacci anyon theory — both are consistent). Derive these $R$-matrices. Confirm Eqs. 10.2 as one of the two solutions and show no other solutions exist.

Exercise 13.2 Evaluation of a Diagram

Consider the following diagram:

Evaluate this diagram in terms of $R$’s and $F$’s. Hint: First reduce the diagram to that shown in exercise 12.3.

Exercise 13.3 Gauge transform of $R$ and Hexagon

(a) Confirm the gauge transform Eq. [13.3]
(b) Show that a set of $F$-matrices and $R$-matrices satisfying the hexagon equations, Eq. 13.1 and 13.2 remains a solution after a gauge transformation. Remember that both $R$- and $F$- transform.

Exercise 13.4 Reidermeister Moves
(a) Use the $R$-matrix, and the completeness relationship, to derive the equivalence shown on the left of Fig. 13.9.
(b) How does the hexagon equation imply the equivalence shown in Fig. 13.14.
Hint: This is very subtle, but is almost trivial.

![Diagram](image)

**Fig. 13.14** This move is implied by the hexagon equation. (Similar with the straight line $f$ going under the other two, and similar if the left-to-right slope of $f$ is negative instead of positive.).

(c) Use Fig. 13.14 to show the equality on the right of Fig. 13.9
(d) Use the result of Fig. 13.14 along with completeness and the $R$-matrix to demonstrate Fig. 13.10

This exercise shows that equalities like those shown in Fig. 13.9 and 13.10 are not independent assumptions but can be derived from the planar algebra and the definition of an $R$-matrix satisfying the hexagon.
When we discussed knot invariants, like the Kauffman bracket invariant, we were allowed to deform a knot in arbitrary ways so long as we didn’t cut any strands. This is what we called isotopy invariance. We would very much like the diagramatic rules of our topological theories to obey such isotopy invariance. However, as we discussed in section 12.3 we really only expect invariance under a more limited set of moves which we called causal isotopy. In this chapter we take steps to achieve the higher level of isotopy invariance that we want in order to construct knot and link invariants.

We have already established the invariance of our rules under certain Reidemeister moves (see Figs. 13.9 and 13.10 and exercise 13.4). However, there is one much more crucial move that we need to have our theory. Whether we are considering a planar theory or a 2+1 dimensional theory, isotopy invariance requires the so-called zig-zag identity shown in Fig. 14.1 which is not a property of theories having only causal isotopy invariance as shown in Fig. 12.35.

Unfortunately, a set of $F$-matrices (even if they satisfy the pentagon self-consistency condition Eq. 9.7) does not generically satisfy this zig-zag identity Fig. 14.1. To see this, consider the manipulations shown in Fig. 14.2. With the physics normalization of diagrams we have been using, the zig-zag identity does not hold.

---

1This chapter continues the development of the diagrammatic algebra in some detail. Those who would like a brief and easier (albeit not as general) introduction to diagrammatic algebra should go straight to chapter 16.

2Meaning regular isotopy — i.e., we should treat strings as ribbons. See section 2.2.1.
14.1 Isotopy Normalization of Diagrams

To fix the zig-zag problem, we take a cue from the Kauffman bracket invariant and change our definition of diagrams just by a small bit. In particular, let a simple loop of particle \( a \), as shown in Fig. 14.3 be given a value which we call \( d_a \), which is sometimes called the “loop weight.” This is different from our prior definition where we set the loop weight to one as in Fig. 12.18. The change here only means that we will be working with unnormalized bras and kets. We will call this normalization “isotopy normalization”.

\[
\langle \text{state}|\text{state} \rangle = \frac{d_a}{\langle \psi|\psi \rangle} = d_a \text{ Isotopy Normalization}
\]

Fig. 14.3 Using a new normalization (which we call “isotopy normalization”) of bras and kets. Compare to Fig. 12.18.

We should not worry about working with unnormalized bras and kets — we are allowed to do this in quantum mechanics. The price for using unnormalized states is that expectations of operators are now given by

\[
\langle \hat{O} \rangle = \frac{\langle \psi|\hat{O}|\psi \rangle}{\langle \psi|\psi \rangle}
\]

instead of the usual expression for normalized states which just has the numerator.

We note that we would very much like to have

\[
d_a = \langle \text{state}|\text{state} \rangle > 0 \quad (14.1)
\]

so that we have a positive definite inner product, which is required by quantum mechanics. Often it is said that “negative normed states break unitarity” since they destroy the concept of the state living in a proper Hilbert space\(^3\). (Below we will also discuss \( d_a \) negative, despite the problems in doing so!).

Note also that for the identity particle \( d_I = 1 \) since we should be able to add and remove vacuum lines freely\(^4\).

With this idea of changing the normalization of kets, we declare the following:

**Henceforth, we will use isotopy normalization!**

With this new normalization, we can recalculate the value of a zig-zag analogous to that of Fig. 14.1. If we can arrange that the prefactor \( d_a [F^{a\bar{a}a}]_{II} \) is unity then straightening a zig-zag such as that in Fig. 14.1 will be an allowed transformation. In the simplest theories, we can

---

\(^3\) The definition of a Hilbert space requires a positive definite inner product.

\(^4\) Evaluation of an empty diagram also gives unity, since one can add or remove vacuum lines freely, we can think of the empty diagram as being equivalent to any number of loops of the vacuum \( I \).
Fig. 14.4 With the new isotopy invariant normalization of diagrams, straightening a zig-zag incurs a factor of \( d_a [F^a_{\bar{a}a}]_{II} \). We will choose the value of \( d_a \) so as to make this factor unity.

simply choose \( d_a \) such that this product is unity. However, this is not always going to be possible to do (this will be discussed in more detail in section 14.2). Instead, what we will always be able to do is to arrange that \( [F^a_{\bar{a}a}]_{II} \) is real and we choose \( d_a \) such that

\[
d_a = \frac{\epsilon_a}{[F^a_{\bar{a}a}]_{II}}
\]

(14.2)

where\(^6\)

\[
\epsilon_a = \pm 1
\]

(14.3)

Given this choice, the product \( d_a [F^a_{\bar{a}a}]_{II} \) in Fig. 14.4 is \( \epsilon_a = \pm 1 \) and the the zig-zag identity (Eq. 14.1) generally becomes modified to that shown in Fig. 14.5.

Fig. 14.5 The modified zig-zag identity. Here \( \epsilon_a \) is always arranged to be +1 or -1. The equality on the left is from Fig. 14.4. The equality on the right follows from Hermitian conjugation of the equality on the left (turning the diagrams upside down and reversing the arrows).

Thus by changing the normalization of a loop in Fig. 14.3 and choosing the value \( d_a \) of this loop appropriately, we arrange such that zig-zag lines as in Fig. 14.5 can be freely straightened out, up to a possible sign. In the simplest theories, \( \epsilon_a = +1 \) for all particle types and zig-zag lines can be straightened out freely without accumulating any sign. However, many theories will not allow this quite so straightforwardly. For the meantime, let us ignore the possible sign accumulated from a zig-zag and return to this sign in section 14.2.

Thus we have defined a new normalization of the loop Fig. 14.3 given by the choice of Eq. 14.2. As we will see in section 17.1 the normalization constant \( d_a \) will turn out (up to a possible sign\(^7\)) to be the same quantum constant\(^5\) This is arranged by gauge transform. See section 14.2.

\(^6\) I like to call \( \epsilon_a \) the “zig-zag phase” in absence of a better name. There is a temptation to identify it with the Frobenius-Schur indicator \( \kappa_a \) (which we will meet in Eq. 14.4) but these may not be the same if we choose \( d_a \) negative.

\(^7\) Note that our conventions allow \( d_a \) to be positive or negative. We will discuss potential issues with \( d_a < 0 \) in section 14.2 below. Some references, including Bonderson [2007] and Kitaev [2006], define \( d_a \) to be the absolute value of this quantity so it is the same as \( d_a \). We have already seen examples where \( d_a < 0 \) (see exercise 2.2).
dimension $d_a$ that we found in Eq. 5.10 from the Hilbert space dimension of fusing anyons together!

$$\begin{align*}
\langle a \mid \text{Isotopy Normalization} \mid b \rangle &= \left(\frac{d_a d_b}{d_c}\right)^{1/4} \langle a \mid \text{Physics Normalization} \mid b \rangle \\
\end{align*}$$

**Fig. 14.6** New “isotopy” normalization for vertices\(^8\). Note that this is consistent with Fig. 12.18 by setting $c = I$ with $a = b$ (and note that $d_I = 1$).

Having changed the normalization of our kets, for consistency we need to change the normalization of fusions and splittings vertices as well. Thus we define new normalization of vertices as shown in Fig. 14.6.

With this new normalization, the orthonormality of trees is now different from what we previously assumed. For example, Fig. 12.32 should now have a factor of $\sqrt{\frac{d_a d_b d_c d_d}{d_e}}$ on the right hand side.

$$\begin{align*}
\delta_{cd} \sqrt{\frac{d_a d_b}{d_c}} \\
\end{align*}$$

**Fig. 14.7** Bubble diagram with isotopy invariant normalization of diagrams. See Eq. 14.8 for how to interpret the square root in cases where $d < 0$.

Similarly our bubble diagram Fig. 12.17 and our completeness diagram Fig. 12.20 need to be modified as shown in Fig. 14.7 and Fig. 14.8\(^9\).

$$\begin{align*}
a \rightarrow b &= \sum_c \sqrt{\frac{d_c}{d_a d_b}} \\
\end{align*}$$

**Fig. 14.8** Insertion of a complete set of states with isotopy invariant normalization of diagrams. See Eq. 14.8 for how to interpret the square root in cases where $d < 0$.

A crucial point is that the $F$-matrix does not need any alteration when we switch from physics normalization to isotopy normalization\(^10\). One can check that in changing normalizations both sides of Fig. 9.1 (equivalently Fig. 12.29) are multiplied by the same factor of $\left(\frac{d_a d_b d_c d_d / d_e}{d_a d_b d_c d_d / d_e}\right)^{1/4}$.

With this isotopy invariant normalization the rules for evaluating planar diagrams are exactly the same as those described in section 12.4 except that loops are now normalized with $d_a$ as in Fig. 14.3 and our orthonormality relationships (Fig. 12.20 and Fig. 12.17) are altered to those shown in Fig. 14.8 and Fig. 14.7.

The same $R$-matrix rules can be applied to diagrams with over- and under-crossings as in chapter 13. The use of the $R$-matrix is unchanged. Be warned, however, that in Fig. 13.7 and 13.8 we have used the completeness relationship Fig. 12.20 which now needs to be modified to

\(^8\)One might be worried how one handles the fractional power when one of the $d$’s is negative. In fact we will only need to consider cases where the factor inside the brackets ends up positive. (See Eq. 11.17 below).

\(^9\)Once again if $N_{ab}^c > 1$ there are additional indices at the vertices and these must match as well. See section 12.3.

\(^10\)Indeed, the reason why we changed the value of all vertices, as in Fig. 14.6 and not just rescale the vertex corresponding to a simple loop as in Fig. 14.3 is in order to keep $F$ from changing.
also have zig-zags as in Fig. 14.5 without incurring any minus signs. Theories of this type are fairly simple to work with. Theories with all positive Frobenius-Schur indicators are sometimes called unimodal or unimodular.

Fig. 14.8 so that when we evaluate crossings we now obtain, for example, Fig. 14.9

\[ \sum_c \sqrt{\frac{d_c}{d_ad_b}} R_{ab}^{cd} = \sum_c \sqrt{\frac{d_c}{d_ad_b}} R_{ab}^{cd} \]

Fig. 14.9 Resolving a crossing with isotopy normalization. Compare to Fig. 13.7. See Eq. 14.3 for how to interpret the square root in cases where \( d < 0 \).

14.2 Gauge Choice and Frobenius-Schur Indicator

Let us now return to the zig-zag in Fig. 14.1 and our choice of the quantity \( d_a \) in Eq. 14.2. First, we claimed that we can always arrange to have \( [F_a^{aad}]_{II} \) \( 1 \) be real. With a gauge choice, we can fix the phase of \( [F_a^{aad}]_{II} \) any way we like, at least for cases where \( a \neq \bar{a} \). Let us see how this can be done. On the far left of Fig. 14.4 we have a vertex \(|aa\rangle\) as well as a vertex which we write as \(|\bar{a}\bar{a}\rangle\) (compare to Fig. 14.10). Note that, at least when \( a \neq \bar{a} \) these two vertices are \emph{not} Hermitian conjugates of each other (recall that when Hermitian conjugating a diagram arrows get reversed as well as reflecting the diagram). By making separate gauge transforms on these two states, these kets can be redefined by an arbitrary phase as discussed in section 9.4, and this phase then ends up in \( [F_a^{aad}]_{II} \) (See the transformation in Eq. 14.2). Thus by a gauge choice we can choose any phase for \( [F_a^{aad}]_{II} \) as long as \( a \neq \bar{a} \). It is often (albeit not always) convenient to choose \( [F_a^{aad}]_{II} \) to be real and positive so that we can have \( d_a \) positive and \( \epsilon_a = +1 \) positive as well.

However, if \( a = \bar{a} \), it is not possible to change \( [F_a^{aad}]_{II} \) by gauge transform. In this case the kets \( |a\bar{a}\rangle \) and \( |ar{a}a\rangle \) are equal and we do not have the freedom to gauge transform them separately. It is easy to show that when \( a = \bar{a} \), the factor of \( [F_a^{aad}]_{II} \) must be real (See appendix 14.6 for a three line proof). The sign of \( [F_a^{aad}]_{II} \) is then a gauge invariant quantity, known as the Frobenius-Schur indicator\(^{11}\)

\[ \kappa_a = \text{sign}[F_a^{aad}]_{II} \epsilon_a \text{sign}[d_a] \quad . \quad \text{(14.4)} \]

If the Frobenius-Schur indicator is positive for all the self-dual particles in a theory, then we can set \( \epsilon_a = +1 \) for all particles and we can also have \( d_a \) positive for all particles. This means that we can both have a positively normed inner product, and we can freely straighten out zig-zags as in Fig. 14.5 without incurring any minus signs. Theories of this type are fairly simple to work with.\(^{12}\)

However, when the Frobenius-Schur indicator of a self-dual particle is negative, things are more complicated as it looks like we must give

\[ a \bar{a} = |a\bar{a}\rangle \]

\[ \bar{a} a = |\bar{a}a\rangle \]

\[ a \bar{a} = |a\bar{a}\rangle \]

\[ \bar{a} a = |\bar{a}a\rangle \]

Fig. 14.10 Cups (top two) and Caps (bottom two). The vertex \( |aa\rangle \) (top figure) and the vertex \( |\bar{a}\bar{a}\rangle \) (second figure) can be assigned different phases as a gauge choice (See section 9.4). The third figure here is the Hermitian conjugate of the top figure and must have the conjugate phase choice. The bottom figure is the Hermitian conjugate of the second figure. In Fig. 14.1 the leftmost figure includes \( |aa\rangle \) and \( |\bar{a}\bar{a}\rangle \), whereas the phases cancel in the loop formed in the middle picture of Fig. 14.4 from \( |aa\rangle \) and \( |\bar{a}\bar{a}\rangle \). Thus choosing gauges we can choose any phase for \( [F_a^{aad}]_{II} \) unless \( a = \bar{a} \) (See section 14.6 for discussion of the effects of gauge transform on \( F \)).

\(^{11}\)For particles which are not self-dual there are several different definitions of what people call the Frobenius-Schur indicator. Some references just define it to be zero for such particles. Other references define it to be \( \epsilon_a \text{sign}[d_a] \) and others define it to be unity! To avoid confusion we will not use the phrase Frobenius-Schur in the context of non-self-dual particles.

\(^{12}\)Theories with all positive Frobenius-Schur indicators are sometimes called unimodal or unimodular.
up either isotopy invariance or positively normed inner product. Unfortunately, many anyon theories, even very simple ones like $SU(2)\mathbb{I}$ (the semion theory) have negative Frobenius-Schur indicators. The Frobenius-Schur sign associated with a zig-zag, though perhaps surprising, is a genuine physical quantity which occurs in more familiar places as well. In section [14.7] we show that such a Frobenius-Schur sign actually occurs for simple systems such as spin-half particles! Independent of why this sign occurs, theories that have self-dual particles with negative Frobenius-Schur indicators seem to force us to choose between several imperfect options:

**Option A: Negative $\epsilon_a$**

Here (for particles with negative Frobenius-Schur indicator) we choose $\epsilon_a = -1$, but $d_a > 0$. With $d_a > 0$ we have a positive definite inner product, and therefore a theory that can properly represent a (necessarily unitary) quantum mechanical system. On the other hand, with $\epsilon_a = -1$, we incur a minus sign for straightening out any zig-zag as in Fig. [14.5] so our diagrammatic rules are not completely isotopically invariant.

Unless otherwise stated we will work with Option A such that all $d_a > 0$ and self-dual particles with negative Frobenius-Schur indicator have $\epsilon_a = -1$ (and all other particles have $\epsilon_a = +1$).

**Option B: Negative $d_a$**

Here (for particles with negative Frobenius-Schur indicator) we choose $\epsilon_a = +1$, but $d_a < 0$. Choosing $\epsilon_a > 0$ means we have isotopy invariant diagram rules. However, negative $d_a$ means a non-positive-definite inner product (See Eq. [14.1]) which is inappropriate for quantum mechanical systems. Nonetheless, one will often see theories with negative $d_a$ in the physics literature attempting to describe quantum mechanical systems. While this appears problematic, often one can reinterpret this as a valid theory with a certain bookkeeping trick whereby we redefine our inner product and signs are pushed around from one place to another to make it look like we have $d_a < 0$ and $\epsilon_a = +1$ whereas what we actually have is the reverse $d_a > 0$ and $\epsilon_a = -1$, and thus we are describing a valid quantum mechanical theory. The interpretation of negative $d_a$ as a bookkeeping trick is elaborated in section [14.5]. It will turn out that that will be precisely equivalent to Option A. Nonetheless it may be convenient at times to work with negative $d_a$ so as to obtain isotopy invariant rules. A good example of this is discussed in chapter [30].

**Option C: Do Something Physically Different!**

There is yet a third possibility — to do something which is physically slightly different. It turns out that there is a way to preserve both the usual definition of the positive definite inner product (i.e., describe a
maintain isotopy invariance, which is what we will discuss next.

14.3 Isotopy Invariant Unitary Rules

Despite the fact that many anyon theories have particles with negative Frobenius-Schur indicators, which appear to break isotopy invariance, it is always possible to construct an isotopy invariant set of rules for evaluating knot and link diagrams while at the same time maintaining unitarity and a proper inner product. Indeed, such rules are natural in the language of category theory (which we will not discuss). To understand what this alternative approach is, it is worth thinking back to what we learned about Chern-Simons theory (which often does have particles with negative Frobenius-Schur indicators).

14.3.1 Isn’t Chern-Simons Theory Isotopy Invariant and Unitary?

Yes! All the way back in chapter 5 we explained that Chern-Simons theory is based on a topologically invariant (diffeomorphism invariant) action, and it is a unitary quantum theory. So why do we now seem to have this contradiction that we either have to give up unitarity (i.e., accept a non-positive-definite inner product) or give up isotopy invariance (i.e., accept a sign when we deform certain lines to insert a wiggle as in Fig. 14.5). What went wrong here? Why do negative Frobenius-Schur indicators show up in Chern-Simons theory?

The discussion of this section is taken from a (yet unpublished) paper by Joost Slingerland and myself. See also references at the end of this chapter.

---

Fig. 14.11 Drawing a particle line as a ribbon rather than a single line. Right-handed framing means the dotted line is to the right of the solid line when walking in the direction of the arrow. Left-handed framing means the dotted line is to the left of the solid line when walking in the direction of the arrow. Framing a right-handed particle $a$ can (only) annihilate a right-handed particle $\bar{a}$ as shown in Fig. 14.12. Note that we cannot turn right-handed framing into left-handed framing by performing a half-twist (flipping over the ribbon) because the ribbon is meant to have a well-defined front and back (say, two different colors on the two different sides): a right-handed framed...
ribbon turned over is inequivalent to a left-handed framed ribbon.

We can then choose a convention that all particles of a particular type are framed, say, right-handed. However, we run into a problem when we try to establish a convention for particles that are self-dual: since we do not draw arrows on particle lines, we cannot distinguish right- from left-handed framing. To consistently describe the framing, we need to establish a bit more notation, known as flags.

The scheme here is based on the idea that we should introduce a new degree of freedom whose physical meaning is related to the framing. A version of this scheme is described by Kitaev [2006] which we will follow (This is essentially equivalent to the discussions of Bakalov and Kirillov [2001]; Turaev and Virelizier [2017]).

While one can apply this scheme to all the particles in the theory, for non-self-dual particles, and self-dual particles with positive Frobenius-Schur indicators, it is easy enough to just choose a gauge so that you can remove zig-zags as in Fig. 14.5 with \( \epsilon = +1 \). As such, we focus here only on the self-dual particles with negative Frobenius-Schur indicators where the minus signs from straightening zig-zags arise.

We assign each cup (pair creation) and cap (pair annihilation) a big triangular arrow (called a “flag”) which can either point left or right as shown in Fig. 14.13. These are not the same arrows we have been using to distinguish particles from their antiparticles. Indeed, the particles we are focusing on here are self-dual! These flags are supposed to indicate the framing of the ribbon.

Hermitian conjugation does not flip the direction of the flag as shown in Fig. 14.14. Thus the corresponding inner product is defined to be positive definite as shown in Fig. ??.

Straightening a wiggle in a ribbon corresponds to cancelling a cup/cap pair pointing in opposite directions as shown in Fig. 14.16.

We can now add a rule which is outside of what can be done in Chern-Simons theory, but will allow us to use the same notation to describe the options described in section 14.3. Let us declare that the direction of a flag can be reversed at the price of a factor of the Frobenius-Schur
Isotopy Invariant Unitary Rules

\[ \begin{array}{c}
\begin{array}{c}
\includegraphics[width=0.1\textwidth]{fig14_15a}
\end{array} \\
\begin{array}{c}
\includegraphics[width=0.1\textwidth]{fig14_15b}
\end{array}
\end{array} = d_a > 0 \]

**Fig. 14.15** The inner product is positive definite. (Flags must align to join ribbons).

\[ \begin{array}{c}
\begin{array}{c}
\includegraphics[width=0.1\textwidth]{fig14_16a}
\end{array} \\
\begin{array}{c}
\includegraphics[width=0.1\textwidth]{fig14_16b}
\end{array}
\end{array} = d_a > 0 \]

**Fig. 14.16** Straightening a wiggle corresponds to canceling a cup/cap pair pointing in opposite directions.

The indicator (which is \( \kappa_a = -1 \) since these are the only particles we are concerned with here)

\[ \kappa_a = 1 \]

Diagrams thus have different values depending on how the flags are to be added to the cups and caps of the diagram (for all self-dual particles with negative Frobenius-Schur indicator). If we make the rule that all flags always point right, we recover the above “Option A” (section 14.2), that is, removing a zig-zag incurs a minus sign (this can be seen from Eqs. 14.5 and ??).

However, another possible choice (motivated by Chern-Simons theory) is to say that flags should be put on cups and caps so that they alternate directions as you walk along any line. Because zig-zags with alternating flags can be freely straightened, we then have an isotopy invariant diagrammatic set of rules. This scheme of decorating with alternating flags then provides a way that any braided anyon theory can be converted into a knot (or link) invariant with full isotopy invariance\(^\text{16}\). Note, however, that it provides a different output from our above “gauge-fixed” choice (Option A).

One should be cautious that we defined our \( F \)-matrices without ever

\(^{16}\text{Meaning regular isotopy invariance: strands should be treated as ribbons.} \)
specifying the direction of any flags (in some F-matrix diagrams with cups and caps, such as in Figs. 14.2 and 14.4 we would be able to decorate these cups and caps with flags). We should realize that in all prior sections, we were implicitly working with the “Option A” version of diagrams — in other words, our definitions of the F-matrix assumes that all flags are pointing to the right. If flags are not all aligned right, we should first flip the flags using Eq. 14.5 before using the F-matrix.

Let us now consider the isotopy invariant rules for evaluation of a knot or link diagram, as defined in this section (where flags alternate direction). Since we have isotopy invariance we can first use this the isotopy invariance to simplify the knot or link as much as possible. Then before applying F-moves, we should flip flags using Eq. 14.5 to get to a gauge fixed diagram (with all flags pointing right) and continuing our evaluation. Once we have a gauge fixed diagram we must keep careful track of zig-zags which may now incur minus signs if they are straightened.

So which convention should one use? The answer is “it depends”. For many mathematics applications (certainly for knot theory) one wants to work with an isotopy invariant set of rules as described in this section. However, for certain diagrammatic calculations, Option A will be more appropriate. We again emphasize that the phases associated with straightening zig-zags can actually be physical as discussed more in section 14.7.

14.3.2 What Have We Achieved?

One of our original hopes for defining a TQFT, way back in chapter 7, was some prescription that would turn a labeled knot or link diagram into a complex amplitude (see Fig. 7.1) where the result would be unchanged by any smooth deformation of space-time (treating the strands of the knots as ribbons, i.e., we are allowed regular isotopy of the diagram). We have now achieved this goal using the method of section 14.3. We should be a bit cautious though, as once we start evaluating diagrams using F-moves, we may decide it is more convenient to switch to a gauge fixed evaluation (Further in some cases we may have fusion diagrams which lose full isotopy invariance due to certain properties of the fusion vertices as we will see in section 14.4).

Note that in chapter 7 when we were defining a TQFT we wanted to more generally have a prescription for turning a knot or link \( \text{embedded in an arbitrary closed manifold} \) into a complex number output. This generalization will indeed be possible, and we will return to this issue in chapter 22. However, for now we note that our scheme gives unity for an empty diagram (which we can think of as any number of loops of the identity particle with \( d_I = 1 \)) so our diagrammatic evaluation corresponds to

\[
\text{diagram} = \frac{Z(S^3 \text{ with labeled link embedded})}{Z(S^3)} \quad (14.6)
\]
\[ Z(S^2 \times S^1 \text{ with labeled link embedded}) \]

with the caveat that the link does not go around the cycle of the \( S^1 \) in the latter case. Note that this normalization matches that of Eq. 5.18.

### 14.4 Impediments to Isotopy Invariance in Fusion Diagrams

Once we consider diagrams with fusion vertices, full isotopy invariance can be even harder to obtain. Even neglecting issues with Frobenius-Schur indicator, we are not guaranteed that we can deform lines in any way we like in the plane. For example, the right hand side of Fig. 12.35 cannot generically be turned into an equality. In Fig. 14.17 and 14.18 we give similar examples (Recapitulating the calculation in Fig. 12.37 but now using isotopy normalization) of turning-up transformations that generically incur nontrivial factors.

**Fig. 14.17** To evaluate the diagram on the left, the vacuum line is inserted and an \( F \)-move is made. The bubble is then removed with Fig. 14.7. Note that if we were to use the physics normalization, the prefactor of \( \sqrt{d_a d_c / d_b} \) would be absent (See Fig. 12.37). Generally we should not expect that the prefactors of \( d \)'s and \( F \) obtained on the right should cancel each other. In chapter 16 we focus on precisely the theories where this does turn out to be unity as is required for full isotopy invariance. More generally, as we will discuss in section 14.8.1, the transform from left to right in this figure is unitary, meaning the resulting factor on the right \( \sqrt{d_a d_c / d_b} [F_a^{c\bar{c}a}]_b I \) is just a magnitude one complex phase.

**Fig. 14.18** The mirror image of Fig. 14.17. Here we use the fact that \( F \) is Hermitian, so \( F^{-1} = [F^*]^T \).

Thus it seems that our most general theory with fusions and with causal isotopy invariance cannot achieve full planar isotopy invariance. Perhaps this is not surprising. Even if we can deform space-time world lines into each other, we might still expect that there would be some minor difference between a process on the far left and far right of Fig. 14.17. On the far left \( c \) and \( \bar{c} \) are produced from the vacuum then \( \bar{c} \) and \( a \) come...
Seeking Isotopy together to form $b$ whereas on the far right, $a$ simply turns into $c$ and $b$. Fortunately, many topological theories are not this complicated: as we will see in chapter 16 there are many theories where one does have full planar isotopy invariance, and the prefactor incurred in the process shown in Figs. 14.17 and 14.18 turns out to be unity. MORE HERE.

14.5 Appendix: Bookkeeping Scheme with Negative $d$

This bookkeeping scheme shows how one can push the minus sign from $\epsilon_a$ onto $d_a$, redefine the inner product and reproduce the results of “Option A” in section 14.2 by using “Option B”\textsuperscript{18}.

We would like to work with a situation where isotopy invariance is assured, i.e., where $\epsilon_a = +1$ for all particle types. For any self-dual particle with $\kappa_a = -1$ we choose $d_a < 0$ to allow $\epsilon_a = +1$. For non-self-dual particles we then choose a gauge such that the following identity is satisfied\textsuperscript{19}

$$\text{sign}[d_a] \text{ sign}[d_b] = \text{ sign}[d_c] \quad \text{when} \quad N_{ab}^c > 0 \quad (14.7)$$

Note that choosing this may involve choosing $d_a < 0$ for certain non-self-dual particles, but we can do this with a gauge choice just as well as we can choose $d_a > 0$. The sign condition Eq. (14.7) assures that that factor $d_a d_b / d_c$ in Fig. 14.6 is positive, so that for example, the square roots in Figs. 14.7 and 14.8 have positive arguments. This nonetheless, leaves an ambiguity as to whether we should take the positive or negative square root. The convention we will use below is that

$$\sqrt{d_a d_b / d_c} = \begin{cases} \text{negative} & d_a < 0 \text{ and } d_b < 0 \\ \text{positive} & \text{otherwise} \end{cases} \quad (14.8)$$

Having chosen $\epsilon_a = +1$ for all particles yet we may generally allow some $d_a < 0$. It may sound problematic to have some $d_a < 0$ since the negative normed state in Fig. 14.3 seem like they would violate the principles of unitary in quantum mechanics. However, with a small reinterpretation of the meaning of our inner product, we can reinterpret our diagrammatics as representing a well behaved unitary theory.

Our reinterpretation of this diagrammatic algebra is quite simple. We evaluate diagrams using the rules given in section 14.1. That is, we use the rules from section 12.4 except that loops are now normalized with $d_a$ as in Fig. 14.3 and our orthonormality relationships (Fig. 12.20 and Fig. 12.17) are altered to those shown in Fig. 14.7 and Fig. 14.8 (noting the choice of sign in square roots given by Eq. 14.8). If there are over- and under-crossings, these can be evaluated using the $R$-matrix as in Fig. 14.9. Crucially, since we have set all $\epsilon_a = 1$, zig-zags like Fig. 14.5 can be freely straightened out (although recall that diagrams like 14.17 may not be freely straightened in general).

In the case where there are some $d_a < 0$, we call the result of this
evaluation the non-unitary evaluation of the diagram as it corresponds to the non-unitary inner product. However, we now insert one additional rule into our list:

(0) Before evaluating a diagram, count the number of negative-$d$ caps, and call it $n$. After fully evaluating the diagram multiply the final result by $(-1)^n$.

Here a negative-$d$ cap occurs when we go forward in time and two particles with $d < 0$ come together to annihilate or form a particle having $d > 0$. (See examples in Figs. 14.19 and 19.4). Another way of counting the negative-$d$ caps is to imagine erasing all lines in the diagram which have $d > 0$. This leaves only a set of closed loops (due to Eq. 14.7). We then just need to count caps in this set of closed loop of the form shown in the left of Fig. 19.4.

With these new rules, we are now describing a unitary positive-normed quantum theory — we call this evaluation of a diagram, including rule 0, the unitary evaluation of the diagram. To understand the intuition behind these rules, consider a self-dual particle with negative Frobenius-Schur indicator. For such particles a zig-zag like in Fig. 14.5 is supposed to incur a minus sign — however, in our scheme we have set $\epsilon = +1$ and instead made $d$ negative. Since $\epsilon = +1$ in the diagrammatic algebra, there is no sign associated with straightening a zig-zag. However, the zig-zag in Fig. 14.5 has a negative-$d$ cap, so in the final evaluation of the diagram (applying rule 0) we correctly obtain the required minus sign.

As a simple example, consider the evaluation of a single loop as in Fig. 14.3 where $d_a < 0$. Before evaluating the loop we count that there is a single negative-$d$ cap on the top of the loop (as in the left of Fig. 19.4). We evaluate the diagram with the rules of section 16.1.2, to obtain $d_a < 0$ as the nonunitary evaluation. However, applying rule (0) this quantity is then multiplied by $-1$, giving the final result for the $-d_a = |d_a| > 0$. This is the result of the unitary evaluation of the diagram, and it is positive as we would hope for a positive definite inner product for a diagram that can be written as $\langle \text{state} | \text{state} \rangle$ (See Fig. 14.3).

As a second example, consider the diagram Fig. 14.21 and let us assume that $d_a, d_b < 0$ and $d_c > 0$. The (nonunitary) evaluation of the diagram (without rule 0) gives $-d_c \sqrt{d_ad_b/d_c}$, the square root coming from Fig. 14.7 and the sign from the rule Eq. 14.8 of how to handle square roots with negative $d$'s. However, applying rule 0, there is a single negative-$d$ cap (from the vertex with $a$ and $b$ coming in from the bottom, and $c$ going out the top), and hence the unitary evaluation of this diagram is $+d_c \sqrt{d_ad_b/d_c}$. Note that this is positive as it should be for a diagram that can be written as $\langle \text{state} | \text{state} \rangle$ analogous to Fig. 14.3.

As a third example, consider the same diagram Fig. 14.21 but consider the case where $d_a, d_c < 0$ and $d_b > 0$. Here the nonunitary evaluation gives $d_c \sqrt{d_ad_b/d_c}$, but applying rule 0, with a single negative-$d$ cap (the top of the $c$ loop) we obtain a final result of the unitary evaluation given by $-d_c \sqrt{d_ad_b/d_c}$. Note that this is also positive as it should be.

The situation described in this section — having a theory which allows
straightening of zig-zag, but has negative $d_a$ is quite common. It is very useful to be able to interpret such theories as being unitary theories with this additional rule 0.

14.6 Appendix: \([F_a^{aaa}]_{II}\) is real

Let $a$ be a self-dual particle (i.e., $a = \bar{a}$). Working with the physics normalization we already showed (Fig. 14.2) that

$$a \bigtriangledown = [F_a^{aaa}]_{II} a$$

Similarly, using an inverse $F$-move, and the fact that $F$ is unitary (See section 9.5.2) we derive

$$a \bigtriangleup = [F_a^{aaa}]^*_{II} a$$

Equivalently the last diagram can be derived as being the Hermitian conjugate the previous diagram.

Finally, assuming only causal isotopy invariance, the equality

$$[F_a^{aaa}]_{II} = [F_a^{aaa}]^*_{II}$$

then shows that \([F_a^{aaa}]_{II}\) must be real.

14.7 Appendix: Spin 1/2 Analogy and Why We Have a Frobenius-Schur Sign

It may seem a bit odd that a zig-zag in a space-time line (as in Fig. 14.5) can incur a minus sign. While this might appear a bit strange it turns out that there is a familiar analog in angular momentum addition — where the particle types (the labels $a, b, c$ etc) correspond to the eigenvalue of total angular momentum squared $J^2$.

Consider three spin-1/2 particles which all taken together are in an eigenstate of $J = 1/2$. We can describe the possible states of the system with fusion trees as in Fig. 14.22 (see also Fig. 9.1) — in this case where $a, b, c$ and $e$ are all labeled with $J = 1/2$. In Fig. 14.22 we can (on the left of the figure) consider either the fusion of the left-most two particles to some angular momentum $d = 0$ (meaning a singlet) or $d = 1$ (meaning a triplet), or we can (on the right of the figure) consider fusion of the right-most two particles to either $f = 0$ or $f = 1$. The $F$-matrix
that relates these two descriptions of the same space is given by \[ [F_{\frac{1}{2},\frac{3}{2}}]_{df} \]
which is often known as a \(6j\) symbol in the theory of angular momentum addition. The analogy of negative Frobenius-Schur indicator here is the fact that \([F_{\frac{1}{2},\frac{3}{2}}]_{00}\) is negative.

\[
\sum_{f} \left[ F_{abc} \right]_{df} \]

Fig. 14.22 The \(F\)-move.

Let us try to see how this happens more explicitly. Given that the total spin is \(1/2\) we can focus on the case where the total \(z\)-component of angular momentum is \(J_z = 1/2\) as well. The state where the leftmost two particles fuse to the identity (or singlet \(J = d = 0\)) can then be written explicitly as

\[
|\psi\rangle = \frac{1}{\sqrt{2}} \left( |\uparrow_1 \downarrow_2 \rangle - |\downarrow_1 \uparrow_2 \rangle \right) \otimes |\uparrow_3 \rangle \tag{14.9}
\]

where the subscripts are the particle labels given in left to right order. This wavefunction is precisely analogous to the lower half (the “ket”) of the far left hand picture in Fig. 14.2.

On the other hand, we could use a basis where we instead fuse the rightmost two particles together first, as in the righthand side of Fig. 14.22. We can write the state where the right two fuse to \(J = f = 0\) analogously as

\[
|\psi'\rangle = |\uparrow_1 \rangle \otimes \left( |\uparrow_2 \downarrow_3 \rangle - |\downarrow_2 \uparrow_3 \rangle \right) \frac{1}{\sqrt{2}} \tag{14.10}
\]

which is precisely analogous to (but the Hermitian conjugate of) the top half (the “bra”) of the left hand side of Fig. 14.2.

It is easy to check that the inner product of these two states \(|\psi\rangle\) and \(|\psi'\rangle\), corresponding to the value of the left diagram of Fig. 14.2 is \(^{20}\)

\[
\langle\psi'|\psi\rangle = -1/2
\]

By redefining the normalization of these states, we can arrange for this overlap to have unit magnitude. However, the sign cannot be removed. The situation is the same for any two half-odd-integer spins fused to a singlet.

### 14.8 Appendix: Some Additional Properties of Unitary Fusion Categories

Unitary fusion categories (the theories we have been discussing!) have two useful properties which we now present. \(\text{bWe do not prove these properties here. More detailed discussion is given by [Kitaev] [2006]. More}^5\)
detailed discussions are given for example in Jones and Penneys [2017] or Etingof et al. [2005]. These latter references are quite mathematical.

14.8.1 Pivotal Property

A property that may seem obvious is known as the pivotal property. This states that there should be isomorphisms between a vertex with a downturned line and one with an upturned line, such as that shown in Fig. 14.23. While this seems like a rather small statement (which is a property of any unitary fusion category) it turns out to be quite powerful. One can deduce from this that the transformations in Figs. 14.17 and 14.18 are unitary — meaning that the constants on the right hand side have unit magnitude

\[ \sqrt{\frac{d_a d_c}{d_b}} [F^c a]_{b1} = \sqrt{\frac{d_a d_c}{d_b}} [F^a c]_{b1} = 1 \] (14.11)

See Kitaev [2006]; Bonderson [2007]. In the more general case where the fusion multiplicity \( N_{ab}^c \) is greater than one, the vertices have additional indices \( \mu \) and \( \nu \) and the transform is a unitary matrix in these indices. An example of this is given in Fig. 14.24.

From this type of identity one can successively turn up and down legs at vertices to obtain the identity in Fig. 14.25 where the prefactor \( C \) in the figure is also a unit magnitude complex scalar (or a unitary matrix in the case where the vertex has an additional index).

Quite a few more identities can also be derived from the pivotal property. Detailed discussions of this property (and its meaning) are given by Kitaev [2006]; Bartlett [2016]. One particularly useful identity is given

\[ a \quad b \quad \leftrightarrow \quad a \quad c \quad b \]

Fig. 14.23 A theory is pivotal if there exist isomorphisms between the states of the Hilbert spaces described by pairs of vertices that differ by downturned and upturned lines.
Appendix: Some Additional Properties of Unitary Fusion Categories

14.8.2 Spherical Property

Theories which are unitary (describing real quantum mechanical particles) have an additional property called being “spherical”. Given a diagram $X$ with a line coming out the top and a line coming in the
bottom. The so-called left trace is defined by connecting up the top line with the bottom line in a loop going to the left, as in the left of Fig. 14.28. The right trace is defined similarly, except that the loop goes to the right of the diagram $X$ as in the right of Fig. 14.28. If the left trace is always equal to the right trace we say that the theory is spherical. The name here comes from the idea that we could pull the string around the back of a sphere in order to turn a left trace into a right trace as shown in Fig. 14.29. However, the spherical property is actually stronger than Fig. 14.29 suggests since it allows us to turn a right trace into a left trace even when there are other objects on the sphere which might prevent us from dragging a string all the way around the back of the sphere.

\[ a \ x \ x \ a \]

**Fig. 14.28** The Spherical Property sets the left trace equal to the right trace as shown in the picture.

\[ a \ x \ x \ a \]

**Fig. 14.29** The naming “spherical” comes from the idea that we can pull the string around the back of a sphere (as indicated by the black arrows) to turn a left trace into a right trace.

An obvious result of the spherical property is that $d_a = d_a$.

### 14.9 Appendix: Higher Fusion Multiplicities

When we have a theory with higher fusion multiplicities (i.e., $N_{ab} > 1$ for at least one fusion channel), then the vertices must be given indices. The first two equations of this appendix is identical to that of section 12.5 except that here we have changed the normalization from physics normalizaton to isotopy invariant normalization.
Fig. 14.30 The bubble diagram when there are fusion multiplicities. This diagram is drawn in the isotopy invariant normalization. Compare to Fig. 12.39.

\[ \sum_{c, \mu, \nu} \sqrt{\frac{d_a d_b}{d_c}} \delta_{cd} \delta_{\mu \nu} \]

Fig. 14.31 Insertion of a complete set of states. When there are fusion multiplicities, these must be summed over as well \( \mu \in \mathbb{N}^e_c \). This diagram is drawn in the isotopy invariant normalization. Compare to Fig. 12.40.

We can combine Fig. 14.31 with Fig. 13.13 to find the generalization of Fig. 14.9.

\[ \sqrt{\frac{d_c}{d_a d_b}} \left[ R_{ab}^{cd} \right]_{\mu \nu} \]

Fig. 14.32 Resolving a crossing with isotopy normalization and higher fusion multiplicity. See Eq. 14.8 for how to interpret the square root in cases where \( d < 0 \).

Further Reading

This is some reading.

Exercises

Exercise 14.1 Higher Fusion Multiplicities
Derive Eq. 14.32.
Twists

Recall from chapter 2 that we considered the procedure of pulling tight a curl in a ribbon as shown in Fig. 15.1 (compare Fig. 2.7). Pulling tight results in a twisted ribbon, which (viewing time as going vertically) corresponds to a particle rotating around its own axis, while at a fixed point in space, as time progresses. The twist in the ribbon can be removed at the cost of a complex phase which we call $\theta_a$, known as the particle’s twist factor. In other words, the particle rotating around its own axis accumulates a phase $\theta_a$ compared to a particle that does not twist. This phase is something we should expect for any particle with a spin, since rotating a spin in quantum mechanics accumulates a phase.

In our diagrammatic notation, we do not draw ribbons. Rather to represent a particle twisting around itself we use blackboard framing as discussed in section 2.2.2 and we always imagine the ribbon lying flat in the plane. We thus formally define the twist factor as given in Fig. 15.2.

1. If one evaluates a diagram using “Option B” from section 14.2 (Detailed in 14.5), during the non-unitary part of the evaluation one must use $\text{sign}(d_a)\theta_a$ instead of $\theta_a$ to remove a curl, since removing a curl also removes a cap.

Invoking the Hermitian conjugation principle (if we reflect a diagram around a horizontal axis, and reverse the arrows so they remain pointing in the same direction, we complex conjugate its amplitude, see Fig. 12.4) we similarly have the mirror image diagrams shown in Fig. 15.3.

It is easy to confirm that the twist factor $\theta_a$ can only be a unit magnitude phase as expected. Proof of this is given in Fig. 15.4.

1 If one might think that $\theta_a$ has been acting strange for a while — but it is just a phase. Ha ha!
One further interesting fact about the twist factor is that

\[ \theta_a = \theta_{\bar{a}} \]

which can be seen from the equality of the diagrams shown in Fig. 15.5.

The twist factor is related to the so-called topological spin, or conformal scaling dimension, usually called \( h_a \), via the relation

\[ \theta_a = e^{2\pi ih_a} \]

This phase accumulated from a \( 2\pi \) rotation is what we typically get in quantum mechanics from the operator \( e^{2\pi i\hat{S}} \) with \( \hat{S} \) the spin operator and we set \( \hbar = 1 \). The vacuum, or identity particle, should have zero scaling dimension, \( h_I = 0 \).

Note that many quantities of interest will depend only on the twist factor \( \theta_a \), i.e., the fractional part of the topological spin, \( h_a \) mod 1. Indeed, we will see that many of the topological properties of a system are independent of the integer part of the topological spin, and care only about the fractional part. That said, in chapter *** below we will also find cases where the integer part of \( h_a \) is important too.

Recall also the famous spin-statistics theorem (as discussed near Fig. 2.7), which tells us that the twist factor should give us the phase for exchanging two identical particle, and is thus intimately related to the anyonic statistics of particles. Of course two cases are very well known to us: if the spin \( h_a \) is an integer, then \( e^{2\pi ih_a} \) is the identity, and the particle is a boson. If \( h_a \) is a half-odd-integer, then the phase is \(-1\) and the particle is a fermion.

### 15.1 Relations between \( \theta \) and \( R \)

Braiding and twisting are very closely related to each other. In fact, twist factors \( \theta \) are related to the \( R \)-matrices we introduced in chapters 10 and 13 in several different ways.

First, let us try to evaluate the curled ribbon in Fig. 15.2 using the \( R \)-matrix as in Fig. 15.6.

\[ \theta_a = \sum_c \frac{d_c}{|d_a|} R_{ae}^{aa} N_{ea}^{aa} \]

\[ \theta_{\bar{a}} = \sum_c \frac{d_c}{|d_{\bar{a}}|} R_{ae}^{aa} N_{ea}^{aa} \]

This manipulation establishes the relation\(^3\)\(^4\)

\[ \theta_a = \sum_c \frac{d_c}{|d_a|} R_{ae}^{aa} N_{ea}^{aa} \quad (15.1) \]

---

\(^3\)Note that \( \theta_a \) is gauge independent, whereas \( R_{ab}^{cd} \) is generally gauge dependent. (Although \( R_{ab}^{cd} \) is gauge independent as well).

\(^4\)If \( d_a < 0 \) as in the case of using “Option B” from section 14.2, the curl correctly gives \( \text{sign}(d_a) \theta_a \) as mentioned in note 11 above.
where we are assuming \( N_{aa}^{\alpha\alpha} = 0 \) or 1. (See section 15.2 for the case where \( N_{aa}^{\alpha\alpha} > 1 \).

A second, and different, relationship can be derived via the manipulations shown in Fig. 15.7. One might be tempted to identify the left of

\[
\begin{array}{c}
\quad \epsilon_a = \epsilon_a \\
\quad \epsilon_a = \epsilon_a \\
\quad \theta_a \theta_b \\
\end{array}
\]

Fig. 15.7 This curl is evaluated with the \( R \) matrix. Note that this diagram is not precisely equal to \( \theta^*_a \) (See Fig. 15.8).

Fig. 15.7 with the twist factor \( \theta \) but this is not quite right when we look at it carefully, as shown in Fig. 15.8. From Fig. 15.7 and 15.8 we derive\(^5\)

\[
\begin{array}{c}
\quad \epsilon_a = \epsilon_a \\
\quad \epsilon_a = \epsilon_a \\
\quad \theta_a \theta_b \\
\end{array}
\]

Note that in this equation, and Figs. 15.6 and 15.7 we are assuming the gauge choice discussed in section 14.5, and we are assuming a unitary theory. If we are doing a non-unitary evaluation, then rule 0 is dropped in Fig. 15.8 losing the factor of sign(\( d_a \)).

\[
R_I^{\alpha a} = \text{sign}([F_a^{\alpha a}])_{II} \theta_a^* .
\]  \hspace{1cm} (15.2)

For self-dual particles \( \text{sign}([F_a^{\alpha a}])_{II} \) is the Frobenius-Schur indicator. For non-self-dual particles we usually choose \( \text{sign}([F_a^{\alpha a}])_{II} = +1 \)\(^6\)

The final relationship between \( R \) and \( \theta \) is called the ribbon identity\(^7\)

\[
R^{ba}_c R^{cb}_c = \frac{\theta_c}{\theta_a \theta_b}
\]  \hspace{1cm} (15.3)

which can be derived by the geometric manipulations in Fig. 15.9.

\[
\begin{array}{c}
\quad \epsilon_a = \epsilon_a \\
\quad \epsilon_a = \epsilon_a \\
\quad \theta_a \theta_b \\
\end{array}
\]

Fig. 15.9 Deriving the ribbon identity. The middle is the non-obvious geometric step. See also exercise 15.3.
The middle step in Fig. 15.9 is perhaps non-obvious, but is clarified if viewed as a ribbon diagram as in Fig. 15.10. See also exercise 15.3.

15.2 Appendix: Higher Fusion Multiplicities

In the case where we have fusion multiplicities $N_{ab}^c > 1$, then we have the following more general equations. Eq. 15.1 is generalized to

$$\theta_a = \sum_{c, \mu} \frac{d_c}{|d_a|} [R_{c}^{a\mu}][R_{c}^{\mu c}] (15.4)$$

and the ribbon identity Eq. 15.3 is generalized to

$$\sum_{\nu} [R_{c}^{b\nu}][R_{c}^{\nu c}] = \theta_a \theta_b \delta_{\mu \lambda} (15.5)$$

Exercises

Exercise 15.1 Fibonacci Twists

For Fibonacci anyons, the twist factor is $\theta_r = e^{\pm 4\pi i/5}$ (With ± being for right or left-handed theories respectively). Check that these twist factors agree with the $R$-matrices (Eq. 10.2) and Eq. 15.1 and Eq. 15.2

Exercise 15.2 Using Geometric Moves I

(a) Using the allowed moves in Fig. 13.9 show the equivalence of the left and right of Fig. 15.4 (b) Similarly, show the equivalence of the left and right of Fig. 15.5 (c) Similarly show the equivalence of the middle two figures in Fig. 15.8

Exercise 15.3 Using Geometric Moves II
Demonstrate the middle step of Fig. 15.9 by using allowed geometric moves such as Fig. 13.9 and Fig. 13.10 and Fig. 13.11. You may also need the pivotal identity Fig. 14.26.

Exercise 15.4 Gauge Independence of Ribbon Identity
Show that the ribbon identity Eq. 15.3 is gauge independent.

Exercise 15.5 Higher Fusion Multiplicities
Derive Eq. 15.4 and Eq. 15.5.
Theories with Tetrahedral Symmetry (or Full Isotopy)

In chapters 12 through 15 we carefully developed the principles of anyon diagrammatics in quite a bit of generality. In the current chapter we aim for a slightly simplified and abbreviated, but still extremely useful, version of the diagrammatic rules developed (roughly axiomatically) in the prior chapters.

Our original intent for a TQFT was to develop rules that would map a labeled knot or link diagram into a complex amplitude output (as in Fig. 7.1) in a way that would be invariant under any smooth deformations (isotopy) of space-time. Most generally in topological theories, we found that there could some restrictions on what sort of deformations of space-time would leave the output unchanged (See for example section 14.3). In the current section we will focus on a simpler class of theories where these impediments are lifted. In particular the topological theories of this chapter have the property that they give the same output amplitude for any smooth deformation of space-time (treating world lines as ribbons). In other words, in this chapter we assume our theories have “full isotopy” invariance (or “regular isotopy” invariance as discussed in section 2.6.1). An example of such full isotopy is shown in Fig. 16.1. Such invariance is essentially equivalent to something we call “tetrahedral symmetry” which we will describe further below.

![Diagram](image)

**Fig. 16.1** For a theory with full isotopy invariance (regular isotopy invariance) these two diagrams must evaluate to the same result since one can be continuously deformed into the other treating the lines as ribbons.
16.1 Planar Diagrams

We start by considering only planar diagrams, so we do not allow over- and under-crossings (which we re-introduce in section 16.2). Because of our specialization to these fully isotopy invariant theories, our rules for diagrammatic manipulation will be slightly easier than those in chapter 12.

As in chapter 12 there is still a bra and ket interpretation of diagrams. Roughly we can think of cutting a diagram in half and viewing one side as a bra and the other as a ket. We can also roughly think of these diagrams as being world lines of particles moving in 1+1 dimensions.

16.1.1 Planar Diagrammatic Rules

In this chapter all (regular) isotopy of lines is allowed. In particular we are freely allowed to make the deformation shown in Fig. 16.2. In the language of Fig. 14.5 we are assuming all $\epsilon_a = +1$.

As in previous chapters we would like to use $F$-matrices to help us convert one diagram into another. Although we previously found that bending lines up and down (as in Fig. 14.17) can incur nontrivial factors, in this chapter we instead assume no such nontrivial factors so we may turn up and down legs freely. Our $F$-matrix can thus be written as in Fig. 16.3. Note that the conventions we use in this chapter are different from that of the previous chapter but instead match those introduced by Levin and Wen [2005].

In this chapter, the orientation of this diagram (how we direct the legs compared to some direction we call time) does not matter. Further, we can freely rotate the diagrams in Fig. 16.3 and we can bend legs up and down freely as well. For example, the same $F$-matrix as in Fig. 16.3 applies to Fig. 16.4.

We can compare the definition of $F$-matrix in Fig. 16.3 to our prior definition of the $F$-matrix shown in Fig. 9.4. Since we now assume that we can bend legs up and down freely, we can bend legs in Fig. 9.4 and reverse arrows to make it look like Fig. 16.3 and we thereby derive the...
relation between the two definitions

\[ F_{ecf}^{bad} = [F_{e}^{abc}]_{df} \]  

(16.1)

Again the idea of the \( F \)-matrix is to write a single diagram (on the left of Eq. 16.3) as a sum of diagrams on the right. By successively applying such \( F \)-moves to parts of complicated diagrams we can restructure any given diagram in a multitude of ways.

There are several further useful rules for diagram evaluation. First, we need to give a value to the labelled loop as in Fig. 16.5. As in the case of the Kauffman bracket invariant, the value of the loop\(^2\) will be called \( d \), here indexed with a subscript \( a \) for each possible particle type \( a \).

\[ a = d_a = \overline{d}_a \]

Fig. 16.5 The value of a loop labeled \( a \) is given by the quantum dimension \( d_a \). Here we have invoked the spherical assumption to give us \( d_a = \overline{d}_a \).

It is always true that \( d_f = 1 \), meaning that loops of vacuum can be freely added or removed from a diagram. As emphasized in section 14.1 giving the loop this normalization implies we are working with non-normalized kets (see Fig. 14.3 and also note 20 of chapter 2). For now we will allow \( d_a \) to be either positive or negative and we will discuss the meaning of this further in section *** below.

Secondly we define the contraction of a bubble as shown in Fig. 16.6.

\[ a = \overline{d}_a \delta_{cd} \sqrt{d_a d_b d_c} \]

Fig. 16.6 Contraction of a bubble for fully isotopy invariant theories. In cases where some \( d \)'s are negative we interpret the sign outside the square root as negative if and only if both \( d_a < 0 \) and \( d_b < 0 \). We must have \( N_{ab}^c = 1 \) or the value of the diagram is zero. The case with fusion multiplicities \( N_{ab}^c > 1 \) is given in Fig. 16.28.
Theories with Tetrahedral Symmetry (or Full Isotopy)

This identity is the same as Fig. 14.7 only written sideways (in this chapter the orientation of the diagram on the page does not matter!). Physically we should think of this as being a version of the locality rule of section 8.2 — looked at from far away, one does not see the bubble. In particular this locality rule implies the “no-tadpole” rule, that any diagram of the sort shown in Fig. 16.7 must vanish unless the incoming line is the vacuum.

\[ \sum_a \sqrt{d_a d_b} \]

We also have again the completeness relation as shown in Fig. 16.8.

\[ a \quad b = \sum_c \sqrt{d_c} \]

This relation is precisely the same as Fig. 14.8 only now we can orient the diagram in any direction.

16.1.2 Summary of Planar Diagram Rules For Fully Isotopy Invariant Theories

Given the rules established in section 16.1.1, we can evaluate any planar diagram and turn it into a complex scalar number made up of factors of \( F \)'s and \( d \)'s—very similar to what we did with the Kauffman bracket invariant, only without over- and under-crossings here. Here are a summary of the rules for diagram evaluation in the case of fully isotopy invariant planar theories. These rules are analogous to those presented in section 12.4.1 only here the rules are simpler.

1. One is free to continuously deform a diagram in any way as long as we do not cut any strand (for this section we assume no over- or under-crossings).
2. One is free to add or remove lines from a diagram if they are labeled with the identity or vacuum (I). See the example in Fig. 16.9.
3. Reversing the arrow on a line turns a particle into its antiparticle (See Fig. 5.3).
(4) A line must maintain its quantum number unless it fuses with another line, or splits.

(5) Vertices are allowed for multiplicities $N_{ab} > 0$ (See section 8.3). This includes particle-antiparticle creation and annihilation processes where $N_{ab} = 1$ (an example is shown in Fig. 16.9).

(6) One can use $F$-moves to change the structure of diagrams.

(7) One can use relations Fig. 16.8 and 16.6 to change the structure of diagrams.

(8) Every diagram can be reduced to a set of loops which can each be evaluated to give $d_a$ for each loop of type $a$.

16.1.3 Negative $d_a$ and Unitarity

We have allowed theories with $d_a$ to be negative. We will assume, however, that

$$\text{sign}[d_a]\text{sign}[d_b] = \text{sign}[d_c] \quad \text{whenever} \quad N_{ab}^c > 0 \quad (16.2)$$

as we described previously in section 14.5 (and as mentioned there, this is not a particularly stringent condition).

As discussed extensively in section 14.2 if we think of a loop as being an inner product $\langle \text{state}|\text{state} \rangle$ as in Fig. 14.3, a negative $d_a$ implies a non-positive-definite inner product, which is forbidden in quantum mechanics. This apparent problem is discussed in detail in sections 14.1–14.2. Here we will very briefly adapt the scheme discussed in section 14.5 to the current situation.

Here we accept that our diagrammatic algebra has negative $d_a$’s. Evaluation of such diagrams with negative $d_a$’s we call the non-unitary evaluation. However, with a small reinterpretation of the diagrams we can still think of these diagrams as describing a unitary theory.

To implement this reinterpretation of the diagrams, as discussed in section 14.5 we add two simple rules to our list for evaluation of diagrams.

(0') We must break the space-time symmetry and define a time direction (often up on the page).

(0) Before evaluating a diagram, count the number of negative-$d$ caps, and call it $n$. After fully evaluating the diagram multiply the final result by $(-1)^n$.

Recall from section 14.5 a negative-$d$ cap occurs when we go forward in time and two particles with $d < 0$ come together to annihilate or form a particle having $d > 0$. (See examples in Figs. 16.10 and 16.11).

These modifications guarantee we are describing a unitary theory. For example, if we take as simple loop like Fig. 16.5 with $d_a < 0$ the naive evaluation (before application of rule 0) gives a negative result. However, the diagram has one negative-$d$ cap and so the result is multiplied by $(-1)^1$ thus giving a positive result as we should expect for a diagram.

**Fig. 16.10** With time going vertical, the left diagram is a negative-$d$ cap if and only if $d_a < 0$ and $d_b < 0$. (The directions of the arrows do not matter, and if the particles are self-dual we do not draw arrows). The right diagram is never a negative-$d$ cap.

**Fig. 16.11** With time going vertical, the left diagram is a negative-$d$ cap if and only if $d_a < 0$. The right diagram is never a negative-$d$ cap. We can think of these diagrams as being the same as the diagrams in Fig. 16.10 with $c$ being the identity. The directions of the arrows do not matter.
that can be interpreted as \langle \text{state} | \text{state} \rangle \text{ as in Fig. 14.3. More examples of how these evaluations work are given in section 14.5.}

### 16.1.4 Constraints and Examples

There are many constraints on our diagrammatic algebras for planar isotopy invariant theories. Here we give such constraints and explain where they all come from.

**Constraint: The Pentagon**

The consistency condition on $F$-matrices given in Eq. 9.7 can be converted to the notation of this chapter (See Eq. 16.1) to give

\[
F_{cdef} F_{abf} = \sum_h F_{gh} F_{edkh} F_{cbh} (16.3)
\]

**Constraint: Relating $F$ to $d$**

For any theory with full planar isotopy the value of $d_a$ should be fixed by the $F$-matrices:

\[
d_a = \frac{1}{F_{aaf}} (16.4)
\]

This is demonstrated by the manipulations of Fig. 14.4 converted into the notation of the current chapter. Recall that we are assuming $\epsilon_a = +1$.

**Constraint: Inversion**

One can perform an $F$-move on the right hand side of Fig. 16.3 to bring it back into the form on the left. We obtain the diagrammatic relation shown in Fig. 16.12.

\[
F_{ba} = \sum_{f, g} F_{bad} F_{ecf} F_{cbf} (16.5)
\]

**Constraint: Rotation**

Rotating the diagram in Fig. 16.3 by 180 degrees and comparing it to the original diagram, one derives

\[
F_{adc} = F_{ba} \quad (16.6)
\]
Constraint: Turning Up and Down

For a theory to be fully isotopy invariant, we must be able to freely make the moves shown in Fig. 14.17. As shown there, this requires

\[ 1 = \sqrt{d_a d_c} / d_b [F_{\alpha \beta}]_{1b} \]

or in the notation of this chapter

\[ F_{\alpha \beta}^{cd} = \sqrt{d_b / d_a d_c} \]

whenever \( b \times c = a + \ldots \), with the sign of the square root taken negative if and only if \( d_a \) and \( d_c \) are both negative.

Constraint: Unitarity

As mentioned above in Fig. 16.3, the \( F \)-matrix, being a change of basis, must be unitary. This means that

\[ \sum_f F_{\alpha \beta f} F_{\alpha \beta f}^* = \delta_{dd'} \]

\[ \sum_d F_{\alpha \beta d} F_{\alpha \beta d}^* = \delta_{ff'} \]

or equivalently \( [F_{\alpha \beta}]^\dagger = [F_{\alpha \beta}]^{-1} \) Comparing the former to Eq. 16.5 we obtain

\[ [F_{\alpha \beta f}]^\dagger = F_{\alpha \beta f}^{*abcd} \]

Constraint: Hermitian Conjugation

Using reflection across the horizontal axis as in Fig. 12.4 we can reflect the \( F \)-matrix equation Fig. 16.3 and compare the reflected to the unreflected diagram to obtain

\[ F_{\alpha \beta f}^{cd} = [F_{\alpha \beta f}]^* \]

\[ = [F_{\alpha \beta d}]^* \]

\[ = [F_{\alpha \beta d}]^* \]

\[ = F_{\alpha \beta f}^{*abcd} \]

where in the second line the first line has been used in combination with Eq. 16.6 whereas in the third line the first line has been used in combination with Eq. 16.10.

Constraint: Reflection

An independent condition that is very often imposed is that the \( F \)-matrix should be invariant under left-right reflection. Compare the diagram shown in Fig. 16.13 to that of Fig. 16.3.
Theories with Tetrahedral Symmetry (or Full Isotopy)

If a theory has left-right reflection symmetry, then we must have a further constraint

$$F_{eef}^{bad} = F_{eef}^{cde}$$

(16.14)

While this additional condition is not required for a planar diagram algebra, and one can even have full isotopy invariance in two dimensions without it, it is often assumed. For isotopically invariant three dimensional theories, such a symmetry is necessary since one can view the diagrams either from the front or the back.

Using Eq. 16.14 along with Eq. 16.13 gives us the natural seeming constraint

$$F_{eef}^{bad} = [F_{eef}^{bad}]^*$$

(16.15)

Example: Evaluating a bubble

As an example of showing how further constraints are derived, let us use $F$-moves to evaluate the bubble shown in Fig. 16.14

$$= \delta_{ad} F_{aef}^{cab}$$

$$= \delta_{ad} F_{aef}^{cab} d_c$$

Fig. 16.14 Evaluation of a bubble diagram. In the first step, as usual we can flip the direction of an arrow and turn a particle into its antiparticle. In the second step we apply an $F$-move (compare to Fig. 16.3). Then by the no-tadpole (locality) rule (Fig. 16.7) , we can set $f$ to the vacuum particle $I$ and hence $a = d$.

However, we also know the value of the diagram in Fig. 16.14 from Fig. 16.6 which gives us $\sqrt{d_c d_b / d_a}$. Thus we derive $F_{aef}^{cab} d_c = \sqrt{d_c / d_a}$, or equivalently (while replacing $b$ with $b$ for simplicity and using $d_b = d_b$) we have

$$F_{aef}^{cab} = \sqrt{d_b / d_a d_c}$$

(16.16)
whenever \( c \times b = a + \ldots \) where the sign of the square root is taken negative if and only if \( d_a \) and \( d_b \) are both negative. Note that Eq. 16.16 could also be obtained from Eq. 16.7 with Eq. 16.13.

**Example: The Theta diagram**

A commonly considered diagram is the Theta diagram \( \Theta(a, b, c) \) shown in Fig. 16.15. This diagram is easily evaluated by using Fig. 16.6 along with the value of a single bubble Fig. 16.5.

\[
\Theta(a, b, c) = \frac{a \cdot b \cdot c}{c} = a \cdot b \cdot c = d_c \sqrt{\frac{d_a d_b}{d_c}} = \sqrt{d_a d_b d_c}
\]

Fig. 16.15 The Theta diagram. This is evaluated by using Fig. 16.6 along with the value of a single bubble Fig. 16.5. The sign on the square root is taken negative unless all three \( d_a, d_b \) and \( d_c \) are positive.

**Example: The tetrahedral diagram**

Let us consider one more evaluation known as the tetrahedral diagram as shown in Fig. 16.16. At this point we are considering this as a planar diagram even though it looks three dimensional! However, we usually consider diagrams to be well defined if they live on the surface of a sphere, so if we want to think about this as being three dimensional, we should think of this as living on a spherical surface.

\[
\Theta(a, b, c, d) = F_{ec}^{bad} \sum_{g} d_f \sqrt{\frac{d_a}{d_f}} \sqrt{\frac{d_b}{d_f}} \sqrt{\frac{d_c}{d_f}} \equiv G_{ef}^{bad}
\]

Fig. 16.16 Evaluation of the tetrahedral diagram. The first step is just smooth deformation. The second step is application of an \( F \) move. Using Fig. 16.6 the index \( g \) must be equal to the index \( f \) and we obtain some factors of \( \sqrt{d} \). Finally we are left with a single loop of \( f \) which gives a factor of \( d_f \) to give the final result which we give the name \( G \). As in Fig. 16.4 the square roots are taken negative if and only if both \( d \)'s in the numerator of the square root are negative.

For theories with full planar isotopy, the tetrahedral diagram has some obvious symmetries. For example, we should have rotational symmetry in the plane as shown in Fig. 16.17 which implies the identity (note the definition of \( G \) in Fig. 16.16)

\[
G_{ec}^{bad} = G_{fe}^{deb} \tag{16.17}
\]
Theories with Tetrahedral Symmetry (or Full Isotopy)

Fig. 16.17 An obvious rotational symmetry of the tetrahedral diagram.

Another symmetry comes from Eq. 16.6

\[ G_{ba} = G_{ec} \]

which we draw as shown in Fig. 16.18.

Fig. 16.18 The first step is the identity in Eq. 16.18 and the second step is a rotation as in Fig. 16.17. Although this is actually a planar diagram it appears as a rotation in 3D.

Although the diagram shown in Fig. 16.18 is a planar diagram, from the far left to the far right, it appears as if it is a rotation in 3D. Using Fig. 16.17 and 16.18 we can rotate this tetrahedron in any way we like.

If one assumes the reflection symmetry Eq. 16.14 then one can also take the mirror image of the tetrahedron as well to obtain an equivalence between 24 tetrahedral diagrams related by symmetries.\(^5\)

16.2 Braiding Diagrams Revisited

So far in this chapter we have considered planar theories only. Extension to fully (regular) isotopy invariant three dimensional\(^6\) theories follows almost exactly the expositions of chapters 13. Here we will recapitulate the key points.

First, any regular isotopy (See section 2.6.1) of diagrams is allowed and does not change the value of the diagram. This means that as long as we treat the lines in a diagram as ribbons, we can deform the diagram into any shape we like. Often this sort of regular isotopy can turn a diagram with braiding into a planar diagram which can then be evaluated using the rules of section 16.1.2. An example of this is shown in Fig. 16.1.

Most generally, however, we will not be able to eliminate all over- and under-crossing of lines just by using isotopy (i.e., by deforming a diagram). To handle crossings, we invoke the \(R\)-matrix discussed in chapter 13. The basic moves we need are summarized in Fig. 16.19 (which just repeat results previously discussed in Fig. 14.3). In the current chapter, where we consider fully isotopy invariant theories, the orientation of the crossing does not matter. So, for example, the same \(R\)-
matrix formula applies to the crossing in Fig. 16.19 (top) as in Fig. 16.20.

\[
\sum_c \sqrt{d_c/d_ad_b} R_{ab}^c
\]

\[
\sum_c \sqrt{d_c/d_ad_b} [R_{ab}^c]^{-1}
\]

Fig. 16.19 Resolving a crossing with isotopy normalization. The square roots are taken negative if any only if \(d_a\) and \(d_b\) are both negative.

Fig. 16.20 With fully isotopy invariant theories, we can rotate diagrams freely. Thus the uncrossing formula here is identical to that in Fig. 16.19 (top).

The \(R\)-matrix moves in Fig. 16.19 allow us to take any diagram with over- and under-crossings and turn it into a planar diagram\(^7\) which can then be evaluated using the rules of section 16.1.2.

We thus add two rules for evaluation of diagrams in three dimensions to our previously stated rules for planar diagrams of section 16.1.2:

1) One is free to continuously deform diagrams in three dimensions in any way as long as we do not cut any strands (and strands are treated as ribbons). In other words we have regular isotopy of diagrams (See section 2.6.1). [This rule replaces rule (1) from the list in section 16.1.2]

9) Over- and under-crossings can be turned into planar diagrams using the \(R\)-matrix as in Fig. 16.19.

16.2.1 Constraints

There are several further constraints on the braiding diagrammatic algebra which we now mention.

Constraint: Rotation

As mentioned in Fig. 16.20 we can rotate crossings freely. By turning a crossing entirely upside-down and comparing to the original crossing,
we obtain an identity which holds for isotopy invariant theories

$$R_{ab}^c = R_{\bar{c} \bar{a} \bar{b}}^c.$$  \hfill (16.19)

**Constraint: Hermitian Conjugation**

As in Eq. ?? we can use Hermitian conjugation to derive

$$[R_c^{ab}]^{-1} = [R_c^{ba}]^*.$$  \hfill (16.20)

**Constraint: Hexagon Equations**

As discussed in section 13.3 there are consistency conditions between $F$ and $R$ matrices known as hexagon equations (Eqs. 13.1 and 13.2). In the notation of the current chapter these can be written as

$$R_{e}^{ca} F_{d \bar{g} f}^{\bar{c} \bar{a} e} R_{g}^{cb} = \sum_f R_{d}^{\bar{g} \bar{c} \bar{a} e} F_{d \bar{g} f}^{\bar{b} \bar{a} f}$$ \hfill (16.21)

$$[R_c^{ca}]^{-1} F_{d \bar{g} f}^{\bar{c} \bar{a} e} [R_c^{cb}]^{-1} = \sum_f R_{d}^{\bar{g} \bar{c} \bar{a} e} [R_{f}^{c \bar{a} f}]^{-1} F_{d \bar{g} f}^{\bar{b} \bar{a} f}$$ \hfill (16.22)

**Relation to Twists**

As detailed in chapter 15 each particle has a twist factor $\theta^a = \theta^{\bar{a}}$ (with $\theta_I = 1$) describing twisted strands as chapter 15 which we show again here for completeness in Fig. 16.21.

![Fig. 16.21 Definition of Twist Factors (See chapter 15 for more details)](image)

In the current chapter the direction the twist is drawn on the page is not important (only its chirality).

The $R$-matrix and the twist factors are related in several ways. The identities Eq. 15.1, 15.2, and 15.3 hold where for fully isotopy invariant theories we set $\epsilon_a = +1$ for all particles.

The situation described in this section — having a theory which is fully isotopy invariant but has negative $d_a$ — is quite common. Fundamentally, as discussed in sections 14.1, 14.2 the need to use negative $d$ comes from negative Frobenius-Schur indicators. There are many topological theories of this type — including very simple theories the semion theory $SU(2)_1$ and more generally theories like $SU(2)_k$. 


16.3 Gauge Transformations

As in sections 9.4 and 13.4.1 it is possible to make gauge transformations of vertices, which makes resulting changes on $F$-matrices (and $R$-matrices, see section *** ) as shown also in Fig. 16.22.

\[ \begin{array}{c}
 a \\
 b \\
 c
\end{array} \quad u_{cb}^{ab} \quad \begin{array}{c}
 a \\
 b \\
 c
\end{array} \]

**Fig. 16.22** We have the freedom to make a gauge transform of a vertex by multiplying by a phase $u_{cb}^{ab}$. The tilde on the right notates that the vertex is in the tilde gauge.

However, since in an isotopy invariant theory we have Fig. 16.23 we conclude that we must have

\[ u_{cb}^{ab} = u_{cb}^{bc} = u_{bc}^{ca} \]

Depending on our bookkeeping of Frobenius-Schur indicators, there may be additional signs if one or more of the particle types is self-dual (See section *** below).

\[ \begin{array}{c}
 a \\
 b \\
 c
\end{array} = \begin{array}{c}
 a \\
 b \\
 c
\end{array} \]

**Fig. 16.23** This is an allowed identity in isotopy invariant theories. (Compare the more general Fig. 14.25.)

Given such a gauge transform, the $F$- and $R$-matrices transform as

\[ \tilde{F}_{ccf}^{bad} = \frac{u_{cb}^{k\ell} u_{c\ell}^{af}}{u_{db}^{bc} u_{de}^{ce} F_{ccf}^{bad}} \]

(16.23)

\[ \tilde{R}_{cb}^{ab} = \frac{u_{cb}^{ba}}{u_{cb}^{ab}} R_{cb}^{ab} \]

(16.24)

16.4 Appendix: Higher Fusion Multiplicities

As in section 9.5.3 when fusion multiplicities are greater than one, the vertices have additional indices which we label with greek indices $\mu, \nu, \ldots$. For example, if $a$ and $b$ fuse to $c$ with $N_{cb}^{ab} > 1$ then the vertex will have an additional index $\mu \in 1 \ldots N_{cb}^{ab}$. Note that compared to section 9.5.3 we do not put black dots on the vertices here. In the conventions of the current chapter we would then have
Theories with Tetrahedral Symmetry (or Full Isotopy)

\[ \sum_{f \lambda \tau} \left[ F_{ecf}^{bad} \right]_{\mu \nu}^{\lambda \tau} \]

Fig. 16.24  $F$-matrix for isotopy invariant theories with fusion multiplicity.

\[ \delta_{cd} \delta_{\mu \nu} \sqrt{\frac{d_a d_b}{d_c}} \]

Fig. 16.25  The locality principle for the isotopy invariant diagrammatic algebra with fusion multiplicity.

\[ \sum_{c, \mu, \nu} \sqrt{\frac{d_c}{d_a d_b}} [R_{c}^{ab}]_{\mu \nu} \]

Fig. 16.26  Insertion of a complete set of states with isotopy normalization and fusion multiplicity.

\[ \sum_{c, \mu, \nu} \sqrt{\frac{d_c}{d_a d_b}} [R_{c}^{ab}]_{\mu \nu}^{-1} \]

Fig. 16.27  Resolving a crossing with isotopy normalization and fusion multiplicity. The square roots are taken negative if any only if $d_a$ and $d_b$ are both negative.

Chapter Summary

- This is an item
Further Reading

This is some reading.
Levin-Lin?
Summary of properties of category?

Exercises

Exercise 16.1 Triangle Bubble Collapse
A useful lemma is the collapsing of a triangular bubble

\[ F^g_{s,a} \sqrt{d_j d_s / d_k} \]

Derive this lemma.
In this chapter we will explore some further structure that is inherent in topological theories.

### 17.1 Quantum Dimension

Recall that we defined $d_a$, the quantum dimension, in terms of how fast the Hilbert space grows as we fuse together many $a$ particles (See Eq. 3.8)

$$\text{Dim of } M \text{ anyons of type } a \sim d_a^M$$

An alternative definition (Eq. 8.10) is that

$$d_a = \text{the largest eigenvalue of the matrix } N_a$$

(17.1)

We have claimed several times that these quantum dimensions are (up to a possible sign) equal to the value of a loop $d_a$ in our diagrammatic algebra. In this section we will finally prove this important result.

To make a connection to our diagrammatic algebra, let us consider fusing two loops labeled $a$ and $b$ as shown in Fig. 17.1

$$\sum_{c,\mu} \sqrt{d_c d_a d_b} \mu = \sum_{c,\mu} \sqrt{d_c d_a d_b} \mu$$

$$= \sum_{c,\mu} \sqrt{d_c} \mu = \sum_{c} N_{ab}^{c} \mu$$

Fig. 17.1 Fusing two loops into a single loop. In the first line we use the completeness relation Fig. [4.33], then we deform to the second line and finally in the last step we remove the bubble using Fig. [14.30].

The result seems rather natural, that $a$ and $b$ can fuse together to form $c$ in all possible ways. The derivation uses the completeness relation in

1Recall that $N_a$ is defined as the matrix with components $[N_a]_{c}^{\mu} = N_{ab}^{c}$ which are the fusion multiplicities.

2This result holds very generally. One might worry that for general theories, without full isotopy invariance, going from the first line to the second line might be problematic. However, it turns out that one does not need full isotopy invariance, just the pivotal property is enough to get to the second line (See section [4.8.1] and exercise [7.1].)
Further Structure

the first line (Fig. 14.8), then we deform to get to the second line, and finally in the last step we remove the bubble using Fig. 14.7.

Now the value of a loop in our diagrammatic algebra is \(d_a\). However, if \(d_a\) is negative we must implement rule 0 from section 14.5 and we always obtain a positive definite result \(|d_a|\) for a single loop (which is appropriate for a diagram which can be interpreted as \langle\text{state}|\text{state}\rangle, see Fig. 14.3). Thus the diagramatic manipulations of Fig. 17.1 give us

\[
|d_a| |d_b| = \sum_c N_{ab}^c |d_c|
\]  

(17.2)

Eq. (17.2) has an interesting interpretation if we define a vector \(\vec{d}_{abs}\) to have components \(|d_c|\) with the vector index being \(c\). We can then rewrite the Eq. (17.2) as an eigenvalue equation

\[
|d_a| \vec{d}_{abs} = N_a \vec{d}_{abs}
\]

Since \(N_a\) is a matrix of nonnegative numbers\(^4\), and \(\vec{d}_{abs}\) is a vector of positive numbers, we identify \(d_{abs}\) as the so-called Perron-Frobenius eigenvector of the matrix \(N_a\), and its eigenvalue \(|d_a|\) is guaranteed by the Perron-Frobenius theorem (see appendix 17.7) to be the largest\(^5\) eigenvalue of \(N_a\) which, by Eq. (17.1) gives us

\[
d_a = |d_a|
\]

as claimed. This derivation does not rely on the theory having a well-defined braiding.

17.2 The unlinking \(\tilde{S}\)-matrix

First let use the locality principle (or no-transmutation) principle (See Fig. 5.1) to show that a closed loop of type \(a\) around a world line of type \(x\) gives some constant which we call \(\tilde{S}_{ax}\) as shown in Fig. 17.2.

(In fact using the \(R\)-matrix we can explicitly derive this identity and evaluate \(\tilde{S}_{ax}\) in terms of twist factors \(\theta\), the fusion multiplicities \(N_{ax}^c\), and the quantum dimensions \(d\). See exercise 17.4. However, we will not need this explicit expression.) Note in particular that \(\tilde{S}_{Ix} = 1\) since the identity loop can be removed for free, and \(\tilde{S}_{aI} = d_a\) since a single loop\(^6\) of \(a\) gives \(d_a\).

\[
\begin{array}{c}
\text{Fig. 17.2} \quad \text{The locality principle tells us that the value of a loop of } a \text{ around a world line } x \text{ is some number which we call } \tilde{S}_{ax}. \quad (\text{Indeed, we can use the } R\text{-matrix to calculate } \tilde{S}_{ax}. \text{ See exercise } 17.4)
\end{array}
\]
The (modular) $S$-matrix

Recall from section 7.3.1 that we defined the $S$-matrix (Eq. 7.6) in several ways. On the one hand we defined

\[ S_{ab} = Z(S^3; \text{a loop linking b loop}) \]  

(17.4)

whereas on the other hand, we said that $S$ was (under certain conditions that the theory has no transparent particles) a unitary transformation between two different bases for describing the Hilbert space of a torus.

Now recall our normalization of diagrams is given by diagram = $Z(S^3; \text{diagram})/Z(S^3)$ (See Eq. 14.6), so we can write\(^7\)

\[ \sum_c \sqrt{d_c} \frac{d_a}{d_a d_b} c = \sum_c N_{a b} \tilde{S}_{c x} \]

(17.3)

This result holds for any anyon theory with a well defined braiding (i.e., that satisfies the hexagon relationship). Note that in the special case where \( x \) is the identity we just recover Eq. 17.2.

17.3 The (modular) $S$-matrix

As mentioned near Fig. 7.13, there is not agreement in the literature as to which way the arrows should point in the definition of $S_{ab}$. Our convention matches that of Kitaev [2006], which seems to be more common.
Further Structure

\[ S_{ab} = \frac{1}{D} \]

where we have defined

\[ D = \frac{1}{Z(S^3)} \]

Using isotopy of diagrams and Hermitian conjugation it is easy to establish (see exercise 17.3) that

\[ S_{ab} = S_{ba} = S_{\bar{a}\bar{b}} = S_{\bar{b}\bar{a}} = S_{\bar{a}}{}^* \bar{b} = S_{\bar{b}}{}^* \bar{a} = S_{\bar{a}}{}^* \bar{b} = S_{\bar{b}}{}^* \bar{a} \]

And further by setting one of the indices to the vacuum \( I \), we are left with a single loop that evaluates to the quantum dimension \( D \), hence giving us

\[ S_{Ia} = S_{aI} = d_a / D. \]

Let us now evaluate the \( S \)-matrix in terms of our unlinking matrix \( \tilde{S} \).

Comparing to the definition of \( S \) in Fig. 17.5 we obtain

\[ S_{ab} = \tilde{S}_{ab} d_b / D = \tilde{S}_{ab} S_{Ib} \]

or equivalently

\[ \tilde{S}_{ab} = \frac{S_{ab}}{S_{Ib}} \]

Plugging this into Eq. 17.3 gives us

\[ \frac{S_{ax} S_{bx}}{S_{Ix}} = \sum_c N_{ab}^{c} S_{cx}. \]

Again this is generally true for any braided anyon theory, i.e., any theory which satisfies the pentagon and hexagon relations.

17.3.1 Unitary \( S = \) Modular

When \( S \) is unitary, we say the theory is modular\(^9\). It turns out that \( S \) is unitary if and only if the identity is the only transparent particles in the theory\(^10\). A particle \( a \) is said to be transparent if braiding \( a \) all the way around any other particle accumulates no phase. Equivalently, in

\(^8\) Here again we are using rule 0 from section 14.5 so that a single loop always evaluates to a positive number \( d_a = |d_a| \) independent of the sign of \( d_a \). If we do not apply rule 0, the right-hand side of Eq. 17.2 will instead give us \( S_{ab} \) sign(\( d_a d_b \)).

\(^9\) We will explain the meaning of the word "modular" in section 17.3.2.

\(^10\) We ran into this concept as far back as section 4.3.2.
17.3 The (modular) S-matrix

In other words, a full braiding of $a$ with any particle is trivial\(^\text{11}\).

It is clear that an $S$ matrix cannot be unitary if there is any transparent particle $a$ besides the identity (since the row $S_{ax}$ and the row $S_{Ix}$ would be proportional to each other). What is not obvious is that the absence of any transparent particle guarantees $S$ is unitary. This statement can be proven using our axiomatic diagrammatic principles (i.e., not invoking any of the topological discussion of section \[7.3.1\] above). However, the proof is a bit complicated and we refer the reader to Kitaev [2006]; Etingof et al. [2015] for details.

As mentioned a number of times, in some sense all “well-behaved” anyon theories are modular (we say they are modular tensor categories). Unfortunately, there are common theories which are not modular. For example, a simple theory of a single fermion which obtains a minus sign under exchange. A full braiding gives a plus sign exactly like the vacuum, and is hence non-modular (although it may have a well defined braiding). It is not unusual to have fermions in a theory that braid identically to the vacuum and thus prevent the theory from being modular.

Let us assume for the remainder of this section that we have a modular theory, i.e., that $S$ is unitary, as we had stated in section \[7.3.1\]. First, unitarity implies that

$$\sum_{a} |S_{aI}|^2 = \frac{1}{D^2} \sum_{a} d_a^2$$

which thus allows us to identify\(^\text{12}\)

$$D = \sqrt{\sum_{a} |d_a|^2}$$

which is usually called the total quantum dimension. Note in particular that this implies

$$S_{II} = 1/D$$

Again assuming a unitary $S$ matrix we can multiply Eq. \[17.9\] by $S^{-1} = S^\dagger$ on the right to obtain the often quoted Verlinde formula\(^\text{13}\)

$$N_{ab}^c = \sum_x S_{ax} S_{bx} [S^{-1}]_{xc} S_{Ix} = \sum_x S_{ax} S_{bx} S_{cx}^\dagger / S_{Ix}$$

which tells us that all the information about the fusion algebra is contained entirely within the $S$ matrix!

Alternatively, one can multiply Eq. \[17.9\] by $S^{-1} = S^\dagger$ on the left to

\[^{11}\text{An equivalent statement using the ribbon identity Eq. \[15.3\] is that } a \text{ is transparent if and only if } \theta_{c/(b \theta_a b)} \neq 1 \text{ for all } b \text{ and } c \text{ whenever } N_{ab}^c \neq 0.\]

\[^{12}\text{We conventionally choose the positive square root. Choosing the negative square root describes a theory whose central charge is different by 4 from the positive square root case, which we can see from Eq. \[17.16\] below.}\]

\[^{13}\text{Verlinde [1988] derived this in the context of conformal field theories. In different context it was derived earlier by Pasquier [1987].}\]
obtain (with $S$ and $N$ treated as matrices on the left)

$$[S^TN_aS]_{xy} = \delta_{xy} \left( \frac{S_{ax}}{S_{Ix}} \right)$$

This means that the $S$ matrix (at least for modular theories) is the unitary matrix that simultaneously diagonalizes all of the $N_a$ fusion multiplicity matrices (We mentioned this previously after Eq. 8.13).

A useful quick application of the Verlinde formula, Eq. 17.13, is to write an expression for the conjugation\textsuperscript{14} matrix

$$C_{ab} \equiv \delta_{ab}$$

This is simply a permutation matrix that permutes each particle with its antiparticle. Obviously $C^2 = 1$ is the identity.

We can find a relationship between $C$ and the $S$-matrix by writing $C$ as the fusion multiplicity matrix of two particles fusing to the identity

$$C_{ab} = N_{ab}^I = \sum_x S_{ax} S_{bx} = [SS^T]_{ab}$$

where we have used the Verlinde formula to evaluate $N_{ab}^I$ along with $S_{Ix}$ being real. Finally using the fact $S$ is symmetric we obtain

$$C = S^2$$

### 17.3.2 Modular Group and Torus Diffeomorphisms

Let us define one more matrix, which is the diagonal matrix of the twist factors

$$T_{ab} = \theta_a \delta_{ab}$$

It turns out to be more useful to absorb an additional complex phase into this matrix, so let us also define\textsuperscript{15}

$$\tilde{T} = Te^{-2\pi ic/24}$$

where $c$ is a real constant, known as the \textit{chiral central charge} which is an important piece of data for an anyon theory. The central charge modulo 8 can be calculated from the twist factors and quantum dimensions via (Fröhlich and Gabbiani\textsuperscript{1990}; Rehren\textsuperscript{1990}),

$$e^{2\pi ic/8} = \frac{1}{D} \sum_a d_a^2 \theta_a$$

We will discuss the central charge further in section \textsuperscript{17.3.3} below.

The set of operations generated by $\tilde{T}$ and $S$ form\textsuperscript{16} a group known as the \textit{modular group}.\textsuperscript{17}

$$S^2 = C \quad C^2 = 1 \quad (S\tilde{T})^3 = C$$

\textsuperscript{14}The word “conjugation” is meant to evoke charge conjugation which changes positive charges to negative charges.

\textsuperscript{15}Most references mean $\tilde{T}$ when they say “$T$-matrix”. However, a few mean $T$.

\textsuperscript{16}We do not need $C$ as an independent generator since $C = S^2$.

\textsuperscript{17}Be warned there are several closely related groups that are sometimes known as the modular group.
For proof of the last identity we again refer the reader to Kitaev [2006]; Etingof et al. [2013]. These relations are equivalent to the group \( SL(2, \mathbb{Z}) \), the group of two-by-two matrices with integer coefficients and unit determinant which has generators

\[
S = \begin{pmatrix} 0 & -1 \\ 1 & 0 \end{pmatrix} \quad T = \begin{pmatrix} 1 & 1 \\ 0 & 1 \end{pmatrix}
\]  

(17.18)

It is easy to check that Eq. (17.17) are satisfied by these two by two matrices.

The modular group has a beautiful topological interpretation: it is the group of topologically distinct orientation preserving diffeomorphisms of the torus surface. To see how this works we consider a torus to be a plane \( \mathbb{R}^2 \) with the lattice of integers \( \mathbb{Z}^2 \) modded out so that each lattice point is identified with every other lattice point.

Any transformation on the plane that one-to-one maps lattice-points to lattice points gives a representative diffeomorphism of the torus. Such transformations are given by the elements of the group \( SL(2, \mathbb{Z}) \) just by mapping points \( \vec{r} \) in the plane to \( A\vec{r} \) where \( A \) is an member of \( SL(2, \mathbb{Z}) \).

Graphical description of the \( S \) and \( T \) transformations are shown on the image in Fig. 17.6.

![Fig. 17.6 The S and T transformations on the surface of a torus. Here one interprets the picture as being put on the surface of a torus with opposite sides of the picture identified. The matrices from Eq. (17.18) are applied to the coordinates in the plane.](image)

This analogy with the diffeomorphisms of the torus is certainly not coincidental! Let think back to the discussion of section 7.3 and 7.4. We considered the solid torus \( D^2 \times S^1 \) with a particle world line of type \( a \) around the handle. We wrote the wavefunction on the surface as

\[ |\psi_a\rangle = |Z(D^2 \times S^1; a)\rangle \]

which forms an orthonormal basis \( \langle \psi_a | \psi_b \rangle = \delta_{ab} \). As mentioned there, this inner product corresponds to sewing together the two solid tori to create \( S^2 \times S^1 \). However, we could also have sewn the tori together after exchanging meridian and longitude to create \( S^1 \), and the inner product then becomes (See Eq. (7.6))

\[
\langle Z(S^1 \times D^2; b) | Z(D^2 \times S^1, a) \rangle = S_{ab}
\]

A different way of thinking of this is that we make a diffeomorphism on...
the surface of the torus before gluing the two halves back together. Thus we could have said
\[ \langle \psi_b | \hat{S} | \psi_a \rangle = S_{ab} \]
where \( \hat{S} \) is the operator that makes the diffeomorphism on the surface (the diffeomorphism precisely exchanges meridian and longitude).

Note further that \( \hat{C} \equiv \hat{S}^2 \) exchanges the meridian and longitude twice, giving a net effect of rotating the torus by 180 degrees. If we think back to the solid torus with a world line of type \( a \) around the handle, the rotation of the torus surface by 180 degrees changes the relative direction of the embedded world line and thus changes \( a \) to \( \bar{a} \), implementing the conjugation operation \( C \).

Finally, we can define a \( \hat{T} \) operation on the torus surface which implements an elementary Dehn twist as described in section 7.4. Analogously the matrix elements of the \( \hat{T} \) operation will give the matrix \( \tilde{T} \). The fact that the \( \tilde{T} \) operation on the state \( | \psi_a \rangle \) corresponds to a twist factor \( \theta_a \) is fairly obvious from looking at Fig. 7.15. The presence of the additional complex factor related to the central charge in Eq. 17.15 is a subtle point and stems from a change in the 2-framing discussed in section 5.3.4.

17.3.3 Central Charge

In Eq. 17.15 and 17.16 we introduced the idea of the central charge of a TQFT. The central charge stems from the relationship between 2+1 dimensional TQFTs and 1+1 dimensional conformal field theories which we will explore more in chapters *** below in the context of the fractional quantum Hall effect. However, this connection has some deep physics buried in it.

Roughly, 1+1 dimensional conformal field theories describe the physics of gapless one dimensional systems\(^{20}\). For a chiral system (meaning that flow only goes in one direction) the central charge \( c \) is related to (i.e., can be defined by) the heat capacity \( c_v \) of the system via \( c_v = \pi k_B^2 T c / (3v) \)
where \( v \) is the speed of light (the velocity of flow at low energy), \( T \) is temperature, \( k_B \) is Boltzmann’s constant\(^{21}\). If one makes a small change in temperature, the change in heat flowing along the edge is then
\[ \delta J^I = (\pi k_B^2 T c / 3) \delta T \]  \hspace{1cm} (17.19)

TQFT are gapped systems: there are no arbitrarily low energy excitations in the bulk. Thus the low temperature heat capacity must be zero. However, in a finite system (say on a disk geometry) the edge of a TQFT system may have gapless modes which can carry heat, and indeed this heat transport defines the central charge of the TQFT as well (Cappelli et al. [2002]). For example, one can consider a finite rectangular strip of the TQFT and attach two opposite ends to two thermal reservoirs differing in temperature by \( \delta T \). The total heat flowing between the reservoirs is then given by Eq. 17.19 where \( c \) is the central charge of the bulk. Note that net heat always flows from the hot reservoir to the cold reservoir\(^{22}\) even though the central charge can have either sign. The sign of

---

\(^{20}\)“Gapless” means having arbitrarily low energy excitations in the large system size limit.

\(^{21}\)\(\pi \) is pi and 3 is three.

\(^{22}\)One must obey the second law of thermodynamics!
the central charge simply determines in which direction (which chirality, clockwise or counterclockwise) around the edges the heat is flowing.

The coefficient in parenthesis in Eq. 17.19 in the context of two dimensional systems is known as the thermal Hall coefficient\(^{23}\) \(\kappa_{xy}\) in analogy with the electrical Hall coefficient \(\sigma_{xy}\) which we will study in chapters ***.

Given two different TQFTs, say \(T\) and \(t\), as described in section 8.5, we can construct a product theory \(T \times t\) which can roughly be thought of as just putting the two theories in the same place at the same time. The central charge of the two theories necessarily add since the total thermal current will just be the sum of the two thermal currents of the two theories.

The fact that the central charge in Eq. 17.16 is only defined modulo 8 has an interesting explanation: There exists a particular CFT, the Chern-Simons theory \((E_8)_1\) (meaning we use the exceptional Lie group \(E_8\) at level 1) which has \textit{no particles} in it except the identity particle \(I\), but has central charge \(c = 8\). Given any TQFT, call it \(T\), we can construct the product theory \(T \times (E_8)_1\) and it would have exactly the same particle content, and hence the same result on the right of Eq. 17.16 but the central charge would be increased by 8 compared to the theory \(T\). (Similarly the opposite chirality theory \((E_8)_{-1} = (E_8)_1\) has central charge \(c = -8\)).

### 17.4 Periodic Tables of TQFTs

As mentioned in sections 9.3 and 13.3, anyon theories (or TQFTs) are extremely constrained by the pentagon and hexagon equations. Indeed, for any given set of fusion rules there are only a finite number of possible solutions up to gauge equivalence \cite{etingof2005}. Once one includes additional conditions, such as the theory being modular, the number of possible solutions drops even more. This makes it possible to consider building a “periodic table” of possible TQFTs — i.e., a complete list of all consistent modular solutions of pentagon and hexagon. The procedure for building this table is to hypothesize that there are only \(n\) different particle types, with \(n\) a small number. With fixed \(n\) one can constrain the possible fusion rules, then pentagon solutions, and finally hexagon solutions. For modular tensor categories this program has been carried out by Rowell et al. \cite{rowell2009} for up to four particle types\(^{24}\) Some of the key results from this periodic table is presented in table 17.1. In chapter 18 we will see a few of the basic principles used for compiling such a table.

Extensions of the idea of a periodic table have been made in a number of directions. Of note, the periodic table has been extended to include all possible braided theories (solutions of pentagon and hexagon) without imposing the modularity condition. Further since theories with fermions are such an important special case, theories that only fail to be modular due to a fermion that braids like the vacuum, have been put in periodic

\(^{23}\) It is also known as the Righi-Leduc coefficient, after Augusto Righi and Sylvestre Anatole Leduc who independently discovered the thermal Hall effect in 1888.

\(^{24}\) See also earlier work by Gepner and Kapustin \cite{gepner1994} as well as Bonderson \cite{bonderson2007}.
Further Structure

Table 17.1: Rank ≤ 4 modular tensor categories. From [Rowell et al. 2009]

<table>
<thead>
<tr>
<th>particle types</th>
<th>Fusion rules</th>
<th># solutions</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>$I$</td>
<td>$X^2 = I$</td>
<td>1</td>
<td>$(E_8)_1$</td>
</tr>
<tr>
<td>$I,X$</td>
<td>$X^2 = I + X$</td>
<td>2</td>
<td>$\mathbb{Z}_2 = SU(2)_1 =$ Semion</td>
</tr>
<tr>
<td>$I,X$</td>
<td>$X^2 = I + X$</td>
<td>2</td>
<td>$(G_2)_1$ = Fibonacci</td>
</tr>
<tr>
<td>$I,X,Y$</td>
<td>$X^2 = Y, XY = I, Y^2 = X$</td>
<td>2</td>
<td>$\mathbb{Z}_2 = SU(3)_1$</td>
</tr>
<tr>
<td>$I,X,Y$</td>
<td>$X^2 = I + Y, XY = X, Y^2 = I$</td>
<td>8</td>
<td>$SU(2)_2$, Ising, $(E_8)_2$</td>
</tr>
<tr>
<td>$I,X,Y$</td>
<td>$X^2 = I + X + Y, XY = X + Y, Y^2 = I + X$</td>
<td>2</td>
<td>$SU(2)_3/\mathbb{Z}_2$</td>
</tr>
<tr>
<td>$I,X,Y,Z$</td>
<td>$X^2 = Y = Z^2, XZ = I = Y^2, XY = Z, ZY = X$</td>
<td>4</td>
<td>$\mathbb{Z}_4 = SU(4)_1$</td>
</tr>
<tr>
<td>$I,X,Y,Z$</td>
<td>$X^2 = I, XY = Z, XZ = Y, Y^2 = I, YZ = X, Z^2 = I$</td>
<td>1</td>
<td>of type $D(\mathbb{Z}_2)$ = Toric Code</td>
</tr>
<tr>
<td>$I,X,Y,Z$</td>
<td>$X^2 = I + X, XY = Z, XZ = Y^2 = I + Y, YZ = X + Z, Z^2 = I + X + Y + Z$</td>
<td>3</td>
<td>of type $(D_4)_1$</td>
</tr>
<tr>
<td>$I,X,Y,Z$</td>
<td>$X^2 = I + X + Y, XY = X + Y + Z, XZ = Y, Y^2 = I + X + Y + Z, YZ = X + Y, Z^2 = I + X$</td>
<td>2</td>
<td>of type $\mathbb{Z}_2 \times \mathbb{Z}_2$</td>
</tr>
</tbody>
</table>

Some comments on this table: $G_2$, $E_8$, $D_4$ and $SU(N)$ are Lie groups. Theories listed, such as $(G_2)_2$ are Chern-Simons theories with the last subscript indicating the level of the theory. $SU(2)_1/\mathbb{Z}_2$ means start with the Chern-Simons theory $SU(2)_1$ and consider only the “even” subset of anyons, which form a complete theory by themselves. (Similar for $SU(2)_2/\mathbb{Z}_2$). $D(\mathbb{Z}_2)$ is the quantum double of $\mathbb{Z}_2$ which we will discuss in section ***. The counting of the number of solutions in most cases is obvious. For $SU(2)_1, (G_2)_1, SU(2)_3/\mathbb{Z}_2$ and $(G_2)_2$ we have the right ($R$) and left ($L$) handed versions of the theory (The left and right version of $(D_4)_1$ turn out to be the same). The 3 cases of Fib $\times$ Fib and $\mathbb{Z}_2 \times \mathbb{Z}_2$ correspond to taking $L \times L$, $R \times R$, and $L \times R$. Note that $R \times L$ is equivalent to $L \times R$ so does not count as a different theory. However, for Fib $\times \mathbb{Z}_2$, all four $L \times L$, $R \times R$, $R \times L$ and $L \times R$ are different. We will discuss the 8 versions of anyon theories related to the Ising or $SU(2)_2$ in section [15.3]. Note that the $(E_8)_1$ theories need not be entirely trivial – even though it has no nontrivial particles – since it can have a central charge which is any integer multiple of 8. Note that nowhere in this table do we have fusion multiplicity $N_{ab} > 1$.  

218 Further Structure
tables as well. References to these works are given in the further reading below.

17.5 Ω Strand (Kirby Color)

A particularly useful object to consider is a weighted sum of particle types particular sum of particle types known as an Ω-strand\(^{25}\), or sometimes “Kirby color” strand\(^{26, 27}\), as shown (purple) in Fig. 17.7.

\[
\Omega = \frac{1}{D} \sum_a d_a \delta_a = \sum_a S_{Ia} \delta_a
\]

Fig. 17.7 A String of Kirby color (Ω-strand) is a weighted superposition of all anyon string types\(^{25}\). Note that the Kirby color string does not have an arrow on it since it is an equal sum over all particles and their antiparticles. Here \(d\) is the quantum dimension, \(S\) is the modular \(S\)-matrix, and \(I\) is the identity, or vacuum, particle.

This weighted sum will occur in several different contexts, including sections\(^{22, 23, 24, 25}\) below. The Ω-strand has some interesting properties. For example, in Fig. 17.8 we show the so-called “killing property” of a loop of Ω for modular theories — a loop of Ω allows only the identity to go through it: all other particle types are “killed.”

\[
\Omega \overset{x}{=} \sum_a S_{Ia} \delta_a = \sum_a S_{Ia} \tilde{S}_{ax} = D \delta_{lx} I
\]

Fig. 17.8 The killing property of the Ω-strand for modular theories. A loop of Kirby color (Ω) allows no particles through it except the identity. In the final step we use unitarity of \(S\) (and the fact that \(S_{Ia} = \tilde{S}_{Ia}\)).

A useful corollary of the Killing property is given in Fig. 17.9.

\[
\Omega = \sum_c \sqrt{\frac{d_c}{d_a d_b}} \delta_{ac} \delta_{bd} = \delta_{ab} \frac{D}{d_a}
\]

Fig. 17.9 The Ω-strand joins two lines due to the killing property which we use in the second step to force \(c\) to be the identity.

Given the factors of \(D\) that occur in these expressions, it is sometimes useful to define an Ω strand with a different normalization

A loop of Ω strand with a twist, shown in Fig. 17.11, interestingly reproduces our expression for the central charge modulo 8, as in Eq. 17.16

---

\(^{25}\)The Ω strand can be defined for any planar diagram algebra, even without a well defined braiding.

\(^{26}\)Sometimes the Ω strand is normalized with an additional factor of \(D\) or \(D^{-1}\) out front. We define this below as \(\tilde{\Omega}\) in Fig. 17.10.

\(^{27}\)One can also define Ω with a prefactor of \(d_a / D\) instead of \(d_a / D\). This is appropriate if one is not applying rule 0, i.e., if one is making a non-unitary evaluation (as discussed in section 14.5).
Further Structure

\[ \tilde{\Omega} = \frac{1}{D^2} \sum_a d_a \frac{\theta_a}{\theta} = \sum_a \frac{S I_a}{D} \]

Fig. 17.10 A different normalization of the Ω-strand. Compare Fig. 17.7
A loop of \( \tilde{\Omega} \) is normalized to have value unity.

\[ \Omega = \frac{1}{D} \sum_a d_a \frac{\theta_a}{\theta} = \frac{1}{D} \sum_a d_a^2 \theta_a = e^{2\pi ic/8} \]

\[ \Omega = \frac{1}{D} \sum_a d_a \frac{\theta_a}{\theta} = \frac{1}{D} \sum_a d_a^2 \theta_a = e^{-2\pi ic/8} \]

Fig. 17.11 For modular theories, a twisted Ω-strand gives \( e^{\pm 2\pi ic/8} \) with \( c \) the (real) central charge. See Eq. 17.16. (See also exercise 22.5)

17.6 Still Further Structure

Modular (and many non-modular) theories have a great deal of extra structure that we have not even touched on. The theories are obviously very highly constrained, so it is rather natural to expect that there will be many nontrivial relationships between the quantities we have discussed.

A useful relationship which is assigned as exercise 17.4 is

\[ S_{ab} = \frac{1}{D} \sum_c N^c_{ab} \theta_a \theta_b d_c \quad . \] (17.20)

Another interesting result is a theorem by Bantay [1997] which gives us the following nontrivial relationship between the Frobenius-Schur indicator \( \kappa_k \) of a particle \( k \) and the modular \( S \) matrix

\[ \sum_{i,j} N^k_{ij} S_{0i} S_{0j} \frac{\theta^2_i}{\theta^2_j} = \left\{ \begin{array}{ll} \kappa_k = \epsilon_k \text{sign}(d_k) & k = \bar{k} \\ 0 & k \neq \bar{k} \end{array} \right. \] (17.21)

A beautiful theorem by Vafa [1988] tells us that for any braided unitary theory (modular or not) all the spin factors \( \theta_a \) must be an \( n^{th} \) root of unity so that \( \theta^n_a = 1 \) where the integer \( n \) is determined only by the fusion multiplicity matrices \( N^c_{ab} \). For example, we have

\[ \prod_b \theta_b^{N_{ab}} = 1 \] (17.22)

with

\[ X_{ab} = -2N^b_{a\bar{a}} N^\bar{a}_{a\bar{b}} - N^b_{aa} N^\bar{a}_{a\bar{b}} + 4\delta_{ab} \sum_q N^q_{aa} N^q_{a\bar{a}} \]
17.7 Appendix: Perron-Frobenius Theorem

The Perron-Frobenius theorem states that for a matrix with all positive entries, there is a unique eigenvector with all positive entries (up to multiplication by an overall constant) known as the Perron-Frobenius eigenvector. The corresponding eigenvalue is the largest magnitude eigenvalue, and is positive. Further for a unitarily diagonalizable nonzero matrix with all non-negative entries, if there exists an eigenvector with all positive entries (up to multiplication by an overall constant), its eigenvalue is positive and is of magnitude greater or equal to all other eigenvalues. The application of the Perron-Frobenius theorem to the fusion matrix $N_a$ is a bit tricky since the theorem is stronger when the matrix being considered has strictly positive entries and the $N_a$ matrices are only guaranteed to have nonnegative entries. To avoid this problem, construct an arbitrary sum of the $N_a$ matrices $M = \sum a \alpha_a N_a$ where all the coefficients $\alpha_a$ are positive. Since all the $N'_a$'s have common eigenvectors (since they are normal matrices and they commute with each other, see section 8.3.1), these are also the eigenvectors of the matrix $M$. Further, all the elements of $M$ are strictly positive, so we may apply the Perron-Frobenius theorem for positive definite matrices to $M$. We thus obtain a Perron-Frobenius eigenvector of $M$ with strictly positive entries (up to a multiplicative constant). But the eigenvectors of $N_a$ match those of $M$ so we have a positive definite eigenvector of $N_a$ which then must be the Perron-Frobenius eigenvectors whose eigenvalue is greater or equal to any eigenvalue of $N_a$.

17.8 Appendix: Algebraic Derivation of the Verlinde Form

In this section we show that we do not need the structure of braiding in order to derive an equation of the Verlinde form, analogous to Eq. 17.13. Let us begin by recalling from section 8.3.1 that for any topological theory\textsuperscript{28,29} the fusion rules are described by fusion multiplicity matrices $N_{ab}^c$, which can be viewed as a set of square normal matrices $N_a$ with indices $b$ and $c$. As discussed in section 8.3.1 these normal matrices commute with each other and therefore can be simultaneously diagonalized by a matrix which we will call $U$ (See Eq. 8.13)

$$N_a = U \lambda^{(a)} U^\dagger$$ \hspace{1cm} (17.23)

where $\lambda^{(a)}$ is a diagonal matrix for each $a$. We note again that we will discover below that for a so-called modular braided theory we will find that $S$ is the modular $S$-matrix. More generally we call $U$ the mock $S$-matrix.

From Eq. (17.23) the columns of $U$ are the simultaneous eigenvectors

\textsuperscript{28}Here we can mean any planar algebra (unitary fusion category) or any 2+1 dimensional topological theory with a braiding (unitary braided fusion category).

\textsuperscript{29}All we actually need is a commutative fusion ring with a unique identity and inverses.
of the $N$ matrices which we can make explicit as

$$
\sum_c [N_a]_c^d U_{cd} = U_{bd}\lambda_d^{(a)}
$$

(17.24)

and no sum on $d$ implied. Note, at this point, the columns of $U$ may be multiplied by an arbitrary phase (i.e., a phase redefinition of the eigenvectors).

Since there is a particle type called the vacuum $I$ (or identity) which fuses trivially with all other particles, we have $[N_a]_I^c = \delta^c_a$ so we have

$$
U_{ad} = \sum_c [N_a]_I^c U_{cd} = U_{Id}\lambda_d^{(a)}
$$

so that

$$
\lambda_d^{(a)} = \frac{U_{ad}}{U_{Id}}
$$

(17.25)

substituting back into Eq. 17.23 we get the Verlinde formula

$$
[N_a]_b^c = \sum_x U_{bx} U_{ax}^{I^*_x} U_{cz} U_{Iz}^{I^*_x} .
$$

(17.26)

This result is extremely general.

17.9 Appendix: Algebraic Derivation that Quantum Dimensions Form a Representation of the Fusion Algebra

Recall that the columns of $U$ are the simultaneous eigenvectors of the $N_a$ matrices. Invoking the Perron-Frobenius theorem, there must be a particular index $z$ such that the eigenvector $U_{bz}$ has all positive entries with fixed $z$ (up to an overall multiplicative constant). This is a common eigenvector of all the $N_a$ matrices, and the corresponding eigenvalues are $\lambda_z^{(a)}$. By the Perron-Frobenius theorem, since the eigenvector is all positive $\lambda_z^{(a)}$ must be the largest eigenvalue of $N_a$. Recalling (Eq. 8.10) that the quantum dimension can also be defined as the largest eigenvalue of $N_a$, we have

$$
d_a = \lambda_z^{(a)}
$$

Now let us multiply the Verlinde relation Eq. 17.26 on both sides by $d_c = \lambda_z^{(c)} = U_{cz}/U_{Iz}$ (see Eq. 17.25) and sum over $c$. We have

$$
\sum_c N_a^c d_c = \sum_{x,c} U_{bx} U_{ax}^{I^*_x} U_{cz} U_{Iz}^{I^*_x} = d_a d_b
$$
where we have used the fact that $U$ is unitary. Thus we conclude

$$d_a d_b = \sum_c N_{ab}^c d_c$$

**Further Reading**

The table from [Rowell et al. 2009] was extended to five particle types in [Bruillard et al. 2013; Hong and Rowell 2010] and to six in [Creamer 2019]. The modularity condition was relaxed to build a periodic table of all unitary braided theories for up to four particles in [Bruillard 2016] and five in [Bruillard and Ortiz-Marrero 2018]. Periodic table for theories with fermions (which are a special subset of non-modular theories) have been built in [Bruillard et al. 2020] (See also [Bruillard et al. 2017]).

**Exercises**

**Exercise 17.1 Fibonacci $S$-matrix** Recall the Fibonacci theory which we introduced in sections 10.2.1 and 10.2.2.

(a) First let us pretend that we have not calculated the $R$-matrices or $\theta$, i.e., we do not know the braiding phases or the twist factors. We only know the fusion rules $\tau \times \tau = I + \tau$. Using the quantum dimensions, we can obtain three out of four elements of the 2 by 2 $S$-matrix. Determine the remaining element of the $S$-matrix by enforcing unitarity.

(b) Given the twist factor $\theta_s = e^{\pm 4\pi i/5}$ (With $\pm$ being for right or left-handed theories), calculate the $S$-matrix explicitly by using Eq. 17.20.

**Exercise 17.2 Using the pivotal property**

Use the pivotal property (Section 14.8.1) to demonstrate the identity shown in Fig. 17.12. You should not assume full isotopy invariance. Nor should you assume $\epsilon = +1$ for any of the particles.

**Exercise 17.3 Symmetries of $S$**

Use isotopy of diagrams and Hermitian conjugation of diagrams to show the identities in Eq. 17.21.

**Exercise 17.4 Evaluation of the $S$-link**

(a) Use the $R$-matrices and Eq. 15.3 to derive the value of the matrix $\tilde{S}_{ax}$ (See Fig. 17.2) in terms of fusion multiplicities, twist factors $\theta$, and the quantum dimensions $d_a$.

(b) From your result show that

$$\begin{array}{c}
\begin{array}{c}
\tau
\end{array}
\end{array} = \sum_c N_{ab}^c \frac{\theta_c}{\theta_a \theta_b} d_c$$

Note that this diagram differs from $S_{ab}$ by a factor of $Z(S^3) = 1/D$.

**Exercise 17.5 Theories with one nontrivial particle**

Consider an anyon theory with only the identity and one nontrivial particle type $a$ having twist factor $\theta_a$. The only possible fusion rules are $s \times s = I + ms$.
for some integer \( m \) (the semion model is \( m = 0 \) the Fibonacci model is \( m = 1 \)).

Calculate \( d_s \) and \( D \) from the fusion rules. Use Eq. [17.20] to calculate the \( S \) matrix in terms of \( \theta_s \). Show that this matrix cannot be unitary for any \( m > 1 \).

This justifies that on table [17.1] there are only two types of theory with one nontrivial particle.

**Exercise 17.6 Product theories** [Easy]

Given two anyon theories \( A \) and \( B \) with corresponding \( S \)-matrices \( S_A \) and \( S_B \).

(a) Show that the product theory \( A \times B \) has \( S \)-matrix \( S_A \otimes S_B \).

(b) Show that \( A \times B \) is modular if and only if both \( A \) and \( B \) are modular.

(c) Show that the central charge of the product theory is the sum of the central charges of the constituent theories. I.e.,

\[
c_{A \times B} = (c_A + c_B) \mod 8 \quad (17.27)
\]

In fact, central charges strictly add in product theories. However, we have only defined the central charge mod 8 so far!

**Exercise 17.7 Probability of Fusion Channels**

Consider a modular anyon theory on a sphere with a very large number of quasiparticles of all types.

(a) Divide these anyons randomly into two large groups. Show that the probability that the two groups have overall fusion channels \( a \) and \( \bar{a} \) is given by

\[
p(a, \bar{a}) = \frac{d_a^2}{D^2}.
\]

Hint: You are counting the total number of fusion trees. Use the strategy of section 8.3 along with the Verlinde formula, and the knowledge that \( |S_0b/S_00| \geq |S_{xb}/S_{xb}| \) for all \( x \). (You may start by assuming this is strictly > and worry about the \( \geq \) case later).

(b) Instead divide the anyons randomly into three large groups. Show that the probability that the groups have overall fusion channels \( a, b, c \) is given by

\[
p(a, b, c) = \frac{N_{abc}d_ad_bd_c}{D^4}
\]

(c) Finally try four large groups. Show that the probability that the groups have overall fusion channels \( a, b, c, f \) is given by

\[
p(a, b, c, f) = \frac{N_{abcf}d_ad_bd_c}{D^6}
\]

where \( N_{abcf} \) is the number of ways \( a, b, c \) and \( f \) can fuse to the vacuum.
Part IV

Some Examples: Planar Diagrams and Anyon Theories
Some Simple Examples

In this chapter we consider a few simple examples of anyon theories. Our strategy will be the same in each case. First, we decide on a set of fusion rules. From this we examine the possible planar planar diagram algebras. To a mathematician these are known as spherical tensor categories (See section 14.5.2). Once we have found the possible planar algebras we will look for possible braidings to build full anyon theories.

All of the examples given here will enjoy full isotopy invariance (with $\epsilon_a = +1$, although if there are nontrivial Frobenius-Schur indicators we will need negative $d$'s) so we will use the notation of chapter 16.

18.1 $\mathbb{Z}_2$ Fusion Rules

Let us start with the simplest system of particles we can imagine, an identity 0 and a nontrivial particle 1. The simplest fusion rules we can have are\(^1\)

$$1 \times 1 = 0$$

which tells us that 1 is its own antiparticle $1 = \bar{1}$ so we do not draw arrows on the corresponding line. This is known as $\mathbb{Z}_2$ fusion rules and is shown in Fig. 18.1 The corresponding fusion multiplicity matrix is

$$N_{11} = N_{00} = N_{10} = N_{01} \text{ and } N_{11} = N_{10} = N_{01} = N_{00} = 0.$$ 

With 0 being the identity, the only nontrivial vertices we can have with these fusion rules is where one particle 1 comes in and one particle 1 also goes out as shown in Fig. 18.2. If one does not draw the identity particle, diagrams must then be just a so-called loop gas as shown in Fig. 18.3. The constraint $N_{01} = N_{10} = N_{10} = 0$ means that loops cannot end, and $N_{11} = 0$ means that loops cannot intersect.

---

\(^1\)We have switched notations — here the vacuum is 0 not $I$, and the nontrivial particle is 1. I hope this does not cause confusion!

Fig. 18.1 Fusing two 1-particles to the vacuum, shown in two notations.

Fig. 18.2 Examples of allowed vertices for the $\mathbb{Z}_2$ fusion rules. A 1 particle (drawn solid) comes into the vertex and the 1-particle must also go out of the same vertex. The 0 particle, the identity, is drawn dotted, but it need not be drawn at all.

Fig. 18.3 A loop gas has $\mathbb{Z}_2$ fusion rules. The loop gas drawn here is planar — there are no over- or under-crossings.
Some Simple Examples

Since the largest eigenvalue of $[N_{ab}]$ is 1, we have quantum dimensions $d_1 = d_0 = 1$ (See Eq. 8.10).

The $\mathbb{Z}_2$ loop gases were studied in Exercise 2.2 (where we allowed over and undercrossings in addition to just planar diagrams), and we will consider them again in section 19.1 below.

With these fusion rules, there are two sets of $F$-matrices that give a consistent fully isotopy invariant planar algebras. These moves two solutions correspond to $d_1 = \pm 1$, which are the only two options given $d_1 = 1$.

18.1.1 $d = +1$ Loop Gas

Here we choose $d_1 = +1$ for the nontrivial particle, in which case every $F$ which is nonzero is +1. (I.e., every $F$ diagram where all vertices are consistent with the fusion rules. See Fig 16.3). In other words, $F_{\text{bad}} = 1$ for every case where $N_{abd} = N_{cde} = N_{bef} = N_{ae} = 1$ and $F$ is zero otherwise. We can write out explicitly the nonzero elements

\[
\begin{align*}
F_{000}^{000} &= 1/d_0 = 1 \\
F_{110}^{110} &= 1/d_1 = 1 \\
F_{101}^{101} &= F_{011}^{011} = 1 \\
F_{101}^{101} &= F_{011}^{011} = 1
\end{align*}
\]

The first two lines are required from Eq. 16.4. Eq. 18.3 is from Eq. 16.7. Eq. 18.4 and Eq. 18.5 can be derived from Eq. 18.2 and Eq. 18.3 by the tetrahedral symmetry equation Eq. 16.17. Examples of these $F$-moves are shown in Fig. 18.4 and 18.5.

The $d = 1$ planar loop gas turns out to be a relatively trivial diagrammatic algebra. The value of every allowed diagram is unity! (or is zero if there is anything disallowed in the diagram, such as the intersection of loops.)

We now turn to consider the possible braidings that we can impose on this planar algebra. The only nontrivial $R$ matrix element is $R_{01}^{11}$. Using the hexagon equation 16.21 and setting $a = b = c = d = 1$ and $e = c = g = 0$ we obtain (the only allowed value of $f$ is 0)

\[
[R_{01}^{11}]^2 F_{110}^{110} = [F_{110}^{110}]^2 R_{110}^{010}
\]

The $R$ on the right is unity, and the $F$’s are all unity. Thus

\[
[R_{01}^{11}]^2 = 1
\]

This limits us to two possible anyon theories for the $d = +1$ loop gas:
Bosons

We choose the $R_0^{11} = +1$ case. This gives us no phases or signs with $F$ moves or braiding. The corresponding twist factor (via Eq. 15.1) is \( \theta_a = +1 \) which corresponds to bosons.

Fermions

We choose the $R_0^{11} = -1$ case. This gives us minus sign under exchange of identical particles. The corresponding twist factor (via Eq. 15.1) is \( \theta_a = -1 \) which corresponds to fermions.

For both bosons and fermions the $S$ matrix describing the braiding is

\[
S = \frac{1}{\sqrt{2}} \begin{pmatrix} 1 & 1 \\ 1 & 1 \end{pmatrix}
\]

which is not unitary, so neither of these two cases are modular\(^2\).

18.1.2 $d = -1$ Loop Gas

Here we choose $d_1 = -1$ for the nontrivial particle, in which case every $F$ which is consistent with the fusion rules is $\pm 1$. The signs of the nonzero elements of $F$ are given as follows

\[
F_{000}^{000} = 1/d_0 = 1 \\
F_{110}^{110} = 1/d_1 = -1 \\
F_{010}^{110} = F_{000}^{111} = 1 \\
F_{101}^{101} = F_{011}^{011} = 1 \\
F_{010}^{101} = F_{100}^{011} = 1
\]

As with the $d = +1$ loop gas, the first two lines are required from Eq. 16.4. Eq. 18.3 is from Eq. 16.6. Eq. 18.3 and Eq. 18.5 can be derived from Eq. 18.2 and Eq. 18.3 by the tetrahedral symmetry equation Eq. 16.17. Note in particular how the signs work in Fig. 16.16 in the definition of the tetrahedral diagram.

It is worth looking at the two different signs that $F$ can take. (If necessary, refer back to Fig. 16.3 for details of how the $F$-matrix is defined). Moves such as shown in Fig. 18.4 simply deform the path of the particle and do not incur a sign. However, the move shown in Fig. 18.5 perform “surgery” on the parths and reconnect loops and does change the sign. Such a surgery always changes the number of loops in the diagram by one. The value of any loop diagram is thus given by

\[
\text{Value of } (d = -1) \text{ loop diagram} = (-1)^{\text{number of loops}}
\]

As discussed in detail in sections 14.2 (See also 16.3), while this is a perfectly consistent planar diagrammatic algebra, it has non-positive definite inner products and therefore is not appropriate for describing

\[\text{Eq. 18.16}\]
quantum mechanics. I.e., this is the non-unitary evaluation of the diagram in the language of section 14.5.

However as discussed in 14.5 we can make a proper unitary theory out of the $d = -1$ loop gas just by implementing rule 0. So the unitary evaluation of a diagram is

$$\text{Value of } (d = -1) \text{\ loop diagram } = (-1)^\text{number of loops} \times \text{number of caps including rule 0} \quad (18.15)$$

For example, in Fig. 18.3 there are 10 loops and 14 caps, so the full value of the diagram is $+1$.

The nontrivial particle here has a negative Frobenius-Schur indicator. Here we have chosen to do our bookkeeping by working with a negative $d$, by maintaining isotopy invariance of diagrams (with $\epsilon = +1$) and implementing rule 0 to obtain a unitary theory. There are other possible ways to account for this sign, but in the end we will always have to pay the price of a minus sign for a space-time zig-zag such as Fig. 14.5.

**Semions**

We now consider possible braidings for the $d = -1$ loop gas. As in the case $d = +1$, we can apply the hexagon to obtain Eq. 18.6. In this case, however, $F_{110}^{110} = -1$ so we obtain

$$[R_0^{111}]^2 = -1$$

Again there are two solutions $R_0^{111} = \pm i$ corresponding to right- and left-handed semions. In either case wrapping a semion all the way around another gives $-1$, so the $S$ matrix is given by

$$S = \frac{1}{\sqrt{2}} \begin{pmatrix} +1 & +1 \\ +1 & -1 \end{pmatrix} \quad (18.16)$$

This is unitary, telling us that the theory is modular. The right handed semion theory $R_0^{111} = \theta_1 = i$ (See Eq. 15.2) is equivalent to the Chern-Simons theory $SU(2)_1$ or $U(1)_2$.

### 18.2 Fibonacci Fusion Rules: The Branching Loop Gas

We now consider Fibonacci fusion rules as discussed in sections 8.2.1 and 9.1 above. Here the nontrivial fusion rule is

$$\tau \times \tau = I + \tau$$

Again $\tau = \bar{\tau}$ is self-dual. These fusion rules allow vertices with three $\tau$ particles (one coming from each direction as shown in Fig. 18.6) so the loop gas can have branches as shown in Fig. 18.7.
The fusion multiplicity matrix $N_{ab}$ is zero if exactly one of the indices is $\tau$ and the other two are $I$. Otherwise $N_{ab} = 1$. We can establish the nonzero components of the $F$-matrices for these fusion rules:

\begin{align*}
F_{\tau I \tau}^{\tau I} &= 1/d_\tau = 1 \\
F_{\tau I \tau}^{\tau I} &= 1/d_\tau \\
F_{\tau \tau I}^{\tau \tau I} &= F_{\tau I \tau}^{\tau I} = 1 \\
F_{\tau \tau I}^{\tau \tau I} &= 1/\sqrt{d_\tau} \\
F_{I \tau I}^{\tau I} &= F_{I \tau \tau}^{\tau I} = 1 \\
F_{I \tau I}^{\tau I} &= 1/\sqrt{d_\tau} \\
F_{I \tau I}^{\tau I} &= F_{I \tau \tau}^{\tau I} = F_{I \tau \tau}^{\tau I} = 1 \\
F_{\tau \tau \tau}^{\tau \tau \tau} &= -1/d_\tau.
\end{align*}

As with the case of the $Z_2$ loop gases, the first two lines are required from Eq. 16.4. Eq. 18.19 and Eq. 18.20 are from Eq. 16.7. Eq. 18.21 comes from Eq. 18.20 and Eq. 18.19. Eqs. 18.22, 18.23, and 18.24 can be derived from Eqs. 18.18, 18.19, 18.20 and 18.21 by the tetrahedral symmetry equation Eq. 16.17. Finally, Eq. 18.22 comes from the requirement that the two by two matrix $[F_{\tau \tau}^{\tau \tau}]$ is a unitary matrix (See Fig. 16.3) which we write out as

\[ F_{\tau \tau}^{\tau \tau} = \begin{pmatrix} 1/d_\tau & 1/\sqrt{d_\tau} \\ 1/\sqrt{d_\tau} & -1/d_\tau \end{pmatrix} \]

The unitarity requirement on this matrix also gives us

\[ \frac{1}{d_\tau^2} + \frac{1}{d_\tau} = 1 \]

The solution to this is

\[ d_\tau = \frac{1 + \sqrt{5}}{2} \]

which matches the expected quantum dimension $d_\tau$ given in Eq. 8.2 as it must, given the considerations of section 17.1. Eq. 18.27 also has a solution with $d_\tau < 0$. However, we cannot accept this solution because it would violate Eq. 14.7.

As in the case of the $Z_2$ loop gases, many of the $F$-matrix elements correspond to simple deformations of paths (isotopy) as in Fig. 18.3.

\[ \text{Fig. 18.7} \quad \text{A Fibonacci branching loop diagram allows intersections of loops, but no loop ends} \]

\[ \text{This } F\text{-matrix matches our previous claim in Eq. 16.3. With any proposed } F\text{-matrix, one should always check that one has a valid solution of the pentagon equation Eq. 9.7 (or Eq. 16.3). See exercise 10.4 for the Fibonacci case!} \]

\[ \text{Since } \tau \times \tau = \tau + \ldots \text{ we need to have } \text{sign}[d_\tau] \times \text{sign}[d_\tau] = \text{sign}[d_\tau]. \]
Some Simple Examples

\[ \frac{1}{d_r} + \sqrt{\frac{1}{d_r}} = \frac{1}{d_r} - \frac{1}{d_r} \]

Fig. 18.8 The F-moves for the Fibonacci branching loop gas. Note that the first line is actually the insertion of a complete set of states as in Fig. 16.8

The nontrivial F-moves (corresponding to the matrix \( F \) in Eq. 18.26) are summarized in Fig. 18.8.

18.2.1 Braiding for Fibonacci Anyons

To determine the possible braiding for Fibonacci fusion rules, we must solve the hexagon equation given the \( F \) matrices we just derived. This is assigned as exercise 13.1. There are two possible solutions, a right-handed solution given in Eq. 10.2, and a left-handed solutions which is the complex conjugate of the right handed solution. These are the only solutions of the hexagon equations for the Fibonacci fusion rules. The solution with \((R_{	au	au})^* = \theta_\tau = e^{\frac{4\pi i}{5}}\) matches the Chern-Simons theory \((G_2)_1\) whereas the case with \((R_{	au	au})^* = \theta_\tau = e^{-\frac{4\pi i}{5}}\) matches \((F_4)_1\).

18.3 Ising Fusion Rules: A Two Species Loop Gas

As discussed in section 8.2.2, the Ising fusion rules (also known as \( SU(2)_2 \) fusion rules) are given by

\[
\begin{align*}
\psi \times \psi &= I \\
\psi \times \sigma &= \sigma \\
\sigma \times \sigma &= I + \psi
\end{align*}
\]

with both particle types being self dual \( \psi = \bar{\psi} \) and \( \sigma = \bar{\sigma} \). These rules describe a loop gas with two non-vacuum particles \( \psi \) (which we draw as blue lines and loops in Fig. 18.9) and \( \sigma \) (which we draw red loops in Fig. 18.9). The rule of this loop gas is that one may have a vertex with two sigmas and one \( \psi \), which appears as a blue line splitting off from a red loop.

Looking at the first fusion rule, \( \psi \times \psi = I \), we realize this rule alone, is simply a \( \mathbb{Z}_2 \) fusion rule. Indeed, this tells us immediately that we have

\[
1/d_I = 1/d_\psi = 1
\]

\[
F_{III}^{II} = F_{\psi I}^{\psi I} = F_{\psi I}^{\psi \psi} = F_{\psi I}^{\psi \psi} = F_{\psi I}^{\psi I} = F_{\psi I}^{\psi I} = F_{\psi I}^{\psi I} = F_{\psi I}^{\psi I}
\]
as given in Eqs. 18.1-18.5. One might wonder why we do not consider
d_{\psi} = -1. This is for the same reason why we could not consider negative
d_{\tau} in the Fibonacci case. Here we must have \text{sign}[d_{\psi}][\text{sign}[d_{\sigma}] = \text{sign}[d_{\psi}]
so we must have d_{\psi} positive.

Very similarly we have

\begin{align*}
F_{\sigma\sigma I}^{\sigma\sigma I} &= 1/d_{\sigma} \\
F_{\tau I\sigma}^{\tau I\sigma} &= F_{\sigma I\sigma}^{\sigma I\sigma} = 1 \\
F_{\sigma I\psi}^{\sigma I\psi} &= F_{\psi I\sigma}^{\psi I\sigma} = 1 \\
F_{\sigma I\tau}^{\sigma I\tau} &= F_{\tau I\sigma}^{\tau I\sigma} = 1 \\
F_{\sigma I\psi}^{\sigma I\psi} &= F_{\tau I\sigma}^{\tau I\sigma} = 1
\end{align*}

The first equation is from Eq. 16.4 and the second from Eq. 16.7. The
last two are derived from the first two via the tetrahedral symmetry
Eq. 16.17.

Further using Eqs. 16.7 and 16.16 we obtain

\begin{align*}
F_{\sigma\sigma I}^{\sigma\sigma I} &= F_{\psi\psi I}^{\psi\psi I} = 1/d_{\sigma} \\
F_{\psi\psi I}^{\psi\psi I} &= F_{\sigma\sigma I}^{\sigma\sigma I} = 1
\end{align*}

Enforcing unitarity on the two by two matrix \( F_{\sigma\sigma} \) we get

\begin{equation}
F_{\sigma\psi}^{\sigma\psi} = -1/d_{\sigma}
\end{equation}

giving the two by two matrix the form

\begin{equation}
[F_{\sigma\sigma}^{\sigma\sigma}] = \begin{pmatrix} 1/d_{\sigma} & 1/d_{\sigma} \\ 1/d_{\sigma} & -1/d_{\sigma} \end{pmatrix}
\end{equation}

The unitarity condition also gives us the condition that

\begin{equation}
d_{\sigma} = \pm \sqrt{2}
\end{equation}

which is expected from section 8.2.2, since the fusion rules give us \( d_{\sigma} = |d_{\sigma}| = \sqrt{2} \). Both of these roots are viable solutions of the pentagon.

The remaining nonzero elements of \( F \) are obtained from Eq. 18.28
18.30 by using tetrahedral symmetry Eq. 16.17 to obtain

\begin{align*}
1 &= F_{\sigma\psi\psi}^{\sigma\psi\psi} = F_{\psi\sigma\psi}^{\psi\sigma\psi} = F_{\psi\psi\sigma}^{\psi\psi\sigma} = F_{\psi\psi\psi}^{\psi\psi\psi} \\
&\quad = F_{\psi\sigma\sigma}^{\psi\sigma\sigma} = F_{\sigma\psi\sigma}^{\sigma\psi\sigma} = F_{\sigma\sigma\psi}^{\sigma\sigma\psi} = F_{\sigma\sigma\sigma}^{\sigma\sigma\sigma}
\end{align*}
234 Some Simple Examples

\[ F_{\sigma \psi \sigma} = F_{\psi \sigma \psi} = F_{\psi \sigma \psi} = F_{\sigma \psi \sigma} \]
\[ -1 = F_{\sigma \psi \sigma} = F_{\psi \sigma \sigma} \]

(18.34)  

(18.35)

The nontrivial \( F \)-moves corresponding to the matrix Eq. (18.31) are shown in Fig. 18.10.

\[ \frac{1}{d_\sigma} \quad + \quad \frac{1}{d_\sigma} \]

\[ \frac{1}{d_\sigma} \quad - \quad \frac{1}{d_\sigma} \]

Fig. 18.10 The nontrivial \( F \)-moves for the Ising fusion rules. Note that the first line is actually the insertion of a complete set of states as in Fig. 16.8.

18.3.1 Braiding For Ising Fusion Rules

The most straightforward way to find all the possible braidings for the Ising fusion rules is to explicitly solve the hexagon equations (13.1) or equivalently Eq. (16.21). We here outline how we proceed (Exercise 18.1 asks you to work out the details!). For each possible settings of the variables in the hexagon equation, we derive a different identity. For each of the following cases, the \( F \)-matrices are simple scalars (1 and \(-1\) only) so we derive

\[ a = \psi, b = c = \sigma, d = I \Rightarrow R_\sigma^\psi R_\sigma^\sigma = R_\sigma^\psi \]
\[ b = \psi, a = c = \sigma, d = I \Rightarrow R_\sigma^\psi R_\sigma^\sigma = R_\sigma^\psi \]
\[ a = b = \sigma, c = \psi, d = I \Rightarrow [R_\sigma^\psi]^2 = R_\sigma^\psi \]
\[ a = b = \sigma, c = d = \psi \Rightarrow [R_\sigma^\psi]^2 = -1 \]

(18.36)  

(18.37)  

(18.38)  

(18.39)

For the following case one uses the two-by-two \( F \) matrix meaning we are working with a two dimensional vector space and the hexagon gives us two identities

\[ a = b = c = d = \sigma, e = I \Rightarrow \begin{cases} [R_\sigma^\psi]^2 = \frac{1}{d_\psi} (1 + R_\sigma^\psi) \\ R_\sigma^\psi R_\sigma^\psi = \frac{1}{d_\psi} (1 - R_\sigma^\psi) \end{cases} \]

(18.40)

These equations are enough to pin down all of the possible solutions for the \( R \)-matrix. From Eq. (18.38) and (18.39) we obtain

\[ R_\psi^\psi = -1 \]

which also implies \( \theta_\psi = -1 \) from Eq. (15.2). Note that since \( \psi \) is a \( \mathbb{Z}_2 \) field with \( d_\psi \) comparing to our above discussion of the \( \mathbb{Z}_2 \) fusion rules we already knew that we had to have \( \psi \) be either a fermion or a boson (we could re-establish this by looking at the hexagon with only \( \psi \) and
is $I$ fields. The hexagon including the $\sigma$ field now establishes $\psi$ to be a fermion!

From Eqs. (18.36) and (18.37) and (18.39) we establish

$$R_{\sigma}^{\sigma \psi} = R_{\psi}^{\psi \sigma} = \pm i$$

This sign is an additional free choice we can make (in addition to the choice of $d_{\sigma} = \pm \sqrt{2}$). To keep these independent choices straight we will use the notation

$$d_{\sigma} = \pm \sqrt{2} \quad (18.41)$$

$$R_{\psi}^{\psi \sigma} = \frac{2}{\pm i} \quad (18.42)$$

We now plug in our choices for $d_{\sigma}$ and $R_{\psi}^{\psi \sigma}$ into the first of Eq. (18.40) to solve for $R_{i}^{\sigma \sigma}$. This gives yet another independent choice of sign for a square root which we label with $\pm$. We thus obtain

$$R_{i}^{\sigma \sigma} = \exp \left[ 2\pi i \left( \frac{3}{8} \right) \right] \exp \left[ 2\pi i \left( \frac{3}{4} \frac{1}{8} \frac{1}{16} \right) \right]$$

Note now that

$$\theta_{i}^{\sigma} = R_{i}^{\sigma \sigma} \text{sign}[d_{\sigma}] = \exp \left[ 2\pi i \left( \frac{5}{8} \right) \right] \exp \left[ 2\pi i \left( \frac{3}{4} \frac{1}{8} \frac{3}{16} \right) \right] \quad (18.43)$$

from which we see there are a total of eight possible choices, and they are all of the possible odd-sixteenth roots of unity which is what we would predict from the fusion rules given Eq. (17.22).

For the record, from Eq. (18.36) we also have

$$R_{\psi}^{\psi \sigma} = \frac{2}{\pm i} R_{i}^{\sigma \sigma} = \exp \left[ 2\pi i \left( \frac{3}{8} \right) \right] \exp \left[ 2\pi i \left( \frac{3}{4} \frac{1}{8} \frac{3}{16} \right) \right] \quad (18.44)$$

We should also check that none of the other hexagon relations are violated for any of these eight solutions (We could for example, evaluate the hexagon with $a = d = \psi$, $b = c = \sigma$ or any of the other possible combinations). Remarkably, perhaps, all eight solutions solve all the hexagon relations with no violations (see exercise 18.1).

The eight possible solutions all have the same $S$-matrix (see exercise 18.2)

$$S = \frac{1}{2} \begin{pmatrix} 1 & \sqrt{2} & 1 \\ \sqrt{2} & 0 & -\sqrt{2} \\ 1 & -\sqrt{2} & 1 \end{pmatrix} \quad (18.44)$$

The two possibilities

$$\begin{align*}
1 &= + \\
2 &= - \\
3 &= - \\
1 &= + \\
2 &= + \\
3 &= -
\end{align*}$$
Exercises

Exercise 18.1 Using the Hexagon for Ising Fusion Rules
Use the hexagon relations to derive Eqs. [18.36] [18.40]. Confirm that the eight solutions we find give no violations of any hexagon relations.

Exercise 18.2 S-matrix for Ising Fusion Rules
Explicitly derive the S-matrix for all eight solutions of the hexagon equation for the Ising fusion rules and confirm that they all give Eq. [18.44]. Thus confirm that all eight solutions are modular. Hint: It might be easiest to use Eq. [17.20].

Exercise 18.3 Frobenius-Schur Indicator for Ising Fusion Rules
Use Eq. [17.21] to calculate the Frobenius-Schur indicator for the \( \sigma \) particle in each of the eight possible solutions of the hexagon equation for the Ising fusion rules. Show that the Frobenius-Schur indicator is negative exactly when \( d_\sigma \) is negative.

Exercise 18.4 Evaluating Diagrams I
Show that evaluation of the diagram in Fig. [18.7] gives \( -d_\sigma^{0/2} \).

Exercise 18.5 Evaluating Diagrams II
Show that evaluation of the diagram in Fig. [18.9] gives \( d_\psi^2 d_\kappa^3 \).

Exercise 18.6 Deriving an \( F \)-matrix
(Easy) Consider a theory containing three particle types, \( I, A, B \) where \( I \) is the identity. Let the nontrivial fusion rules be given by
\[
\begin{align*}
B \times B &= I \\
A \times A &= I + A \\
A \times B &= A
\end{align*}
\]
Let us assume we have a theory with full isotopy invariance and full tetrahedral symmetry. There is only one set of \( F \) matrices for these fusion rules. Find these \( F \)-matrices and convince yourself that it satisfies the pentagon equation.

Exercise 18.7 Deriving an \( F \)-matrix
(Hard) Consider a theory containing three particle types, $I, S, V$ where $I$ is the identity. Let the nontrivial fusion rules be given by
\[
\begin{align*}
S \times S &= I \\
S \times V &= V \\
V \times V &= I + S + V
\end{align*}
\]
Let us assume we have a theory with full isotopy invariance and full tetrahedral symmetry. There is only one set of $F$-matrices for these fusion rules. Find these $F$-matrices. (Hint: applying the techniques of this chapter will give the correct solution as well as a spurious solution which you can eliminate by showing it does not satisfy the pentagon equation.) Show that the hexagon equation is solved with $R_{ab}^c = 1$ for all $a, b, c$. 
Temperly-Lieb Algebra and Jones-Kauffman Anyons

Let us look back at the Kauffman bracket invariant that we introduced in chapter 2. In the current chapter we want to make use of these rules and determine some of the properties of the corresponding anyons in the language we have been developing since chapter 8. Our strategy will be to first consider a planar diagram algebra in detail before considering braiding properties in section 19.5 below.

So we start by considering a planar version of the Kauffman bracket. I.e., we only consider diagrams with no over- and under-crossings. Our diagrams are isotopically invariant in the plane and the only additional rule then is that the a loop is given a value $d$ as shown in Fig. 19.1. As compared to the diagrammatic algebra we have constructed over the last few chapters (roughly starting in chapter 8 and continuing through chapter 16), one things that was missing in the discussion of the Kauffman bracket invariant is the idea of multiple particle types and fusion rules. In this chapter we will try to construct particle types, fusion rules, and $F$-matrices given only the rule 19.1 as a starting point. The planar algebra of loops that we will construct is known as the Temperly-Lieb algebra. (When we reintroduce braiding to our theory the resulting theory is called called Jones-Kauffman, or Temperley-Lieb-Jones-Kauffman.)

Let us start by thinking a bit about what kind of particle types we already have in our theory. Certainly we have the simple string which we will call “1”; and we always have a vacuum particles, which we will call “0”. Now we would like to ask whether we can fuse two of these 1-strings together to make another particle.

Several things are immediately obvious. First consider the fact that two 1-particles can fuse to the vacuum, or in other words, a 1-string can go up and then turn down, as shown in Fig. 19.2. This tells us immediately that

$$1 = 1.$$  

The fact that 1 is its own antiparticle is why we do not draw arrows on the 1-string. For simplicity, if a string is not labeled we will assume it is a 1-string. Given that loop of 1-string is assigned the value $d$, we identify this $d_1$ (which is often called the quantum dimension, although we have been reserving the words “quantum dimension” for $d_1 = |d_1|$).

We might also consider the possibility that two of these 1-particles can fuse to something besides the vacuum, in a way similar to that shown in Fig. 19.3. This is a good idea, but it isn’t yet quite right. If  

---

1 It is admittedly confusing that 1 is not identity, but this is the usual notation! It is (not coincidentally!) similar to spins where spin 0 is the identity (no spin), and spin 1 is nontrivial.
the two strings fuse to some object besides the vacuum 0, we have to make sure that this new object is appropriately “orthogonal” to 0. This orthogonality must be in the sense of the locality, or no-transmutation rule (see Fig. 3.7): a particle type must not be able to spontaneously turn into another particle type (without fusing with some other particle or splitting). In Fig. 19.3 it looks like the two strings brought together could just fuse together to form the vacuum as in Fig. 19.2 and this would then turn the collection of two strings into the vacuum. To prevent such transmutation, we will work with operators known as Jones-Wenzl projectors.

19.1 Jones-Wenzl Projectors

The general definition of a projector is an operator $P$ such that $P^2 = P$. This means that $P$ has eigenvalues 0 and 1. Let us think of a string diagram as an operator that takes as an input strings coming from the bottom of the page, and gives as an output strings going towards the top of the page (compare Fig. 12.2). Now consider a set of $n$-strings traveling together in the same direction (in what is often called a cable). The Jones-Wenzl projection operator $P_n$ operates on a set of $n$ such strings — it takes $n$-strings in and gives $n$-strings out — and it is defined such that attaching a cup or a cap to the bottom or top of the operator gives a zero result (See Fig. 19.4). The $n$-particle Jones-Wenzl projector $P_n$ acting on a cable of $n$-strings should be interpreted as the $n^{th}$ particle species.

The purpose of the Jones-Wenzl projector is to fix the problem we discovered with Fig. 19.3. That is, if a cable of two strings forms a nontrivial particle (the particle we will call 2), we should not be able to put a cap on the top of these two strings and transmute the 2-particle to the vacuum. I.e., adding a cap should make the entire diagram vanish, and this is the property we are looking for in the 2-string Jones-Wenzl projector.

Let us now try to construct the 2-string Jones-Wenzl projector $P_2$ out of two incoming 1-particles\(^2\)(two elementary strings). To do this we first construct a different projector $\bar{P}_2$ that forces the two incoming particles to fuse to the vacuum as shown in Fig. 19.5.

\[
\bar{P}_2 = \frac{1}{d}
\]

Fig. 19.5 The projector of two strings to the vacuum $P_2$. This figure should be thought of as an operator that takes as an input two strings coming in from the bottom, and gives as an output two strings going out the top. Sometimes the operator is represented as a labeled box as shown on the right.

To establish that this $\bar{P}_2$ operator is a projector we need to check that $[\bar{P}_2]^2 = \bar{P}_2$. To apply the $\bar{P}_2$ operator twice we connect the two strings coming out the top of the first operator to two strings coming in the
bottom of the second operator. As shown in Fig. [19.6] using the fact that a loop gets value $d$ we see that $[\bar{P}_2]^2 = \bar{P}_2$ meaning that $\bar{P}_2$ is indeed a projector.

$$[\bar{P}_2]^2 = \frac{1}{d} \qquad \rightarrow \qquad \frac{1}{d^2} = \frac{1}{d} = \bar{P}_2$$

**Fig. 19.6** Checking that $[\bar{P}_2]^2 = \bar{P}_2$. In the second step we have used the fact that a loop gets the value $d$.

The Jones-Wenzl projector $P_2$ for two strings is the complement of the operator $\bar{P}_2$ we just found, meaning $P_2 = I - \bar{P}_2$ where $I$ is the identity operator, or just two parallel strings. Diagrammatically we have Fig. [19.7]. Since the $\bar{P}_2$ operator projects the two strings onto the vacuum, the $P_2$ operator projects the two strings to a different orthogonal particle type which we call 2.

$$P_2 = \left[ -\frac{1}{d} \right] = \bar{P}_2$$

**Fig. 19.7** The projector of two strings to the nontrivial particle made of two strings $P_2 = I - \bar{P}_2$. Sometimes this projector is drawn as a labeled box, as on the right.

We can algebraically check that $P_2$ is indeed a projector

$$P_2^2 = (I - \bar{P}_2)(I - \bar{P}_2) = I - 2\bar{P}_2 + \bar{P}_2^2 = I - \bar{P}_2 = P_2$$

and also we can check that $P_2$ is orthogonal to $\bar{P}_2$, by

$$\bar{P}_2P_2 = \bar{P}_2(I - \bar{P}_2) = \bar{P}_2 - \bar{P}_2^2 = 0$$

and similarly $P_2\bar{P}_2 = 0$.

Often it is convenient to draw these projection operators as a labeled box, as shown on the right of Figs. [19.5] and [19.7]. Sometimes instead of drawing two lines with a projector $\bar{P}_2$ or $P_2$ inserted, we simply draw a single line with a label, 0 or 2 respectively as in the right of Fig. [19.10] or the left of Fig. [19.8].

It is useful to calculate the value of the of the 2-string loop$^4$. This is shown in Fig. [19.8]

$$d_2 = \frac{1}{d} = \bar{P}_2 = \frac{1}{d} \frac{1}{d} = d^2 - 1$$

**Fig. 19.8** Evaluating the 2-string loop.

$^4$In many references $d_2$ is called $\Delta_2$ (and similarly $d_n$ is called $\Delta_n$). We will stick with $d$ to fit with the notation in the rest of this book.
Abelian Case

In the case where \( d = \pm 1 \) it is easy to prove (see Exercise 2.2 and ***) that two horizontal strings equals \( d \) times two vertical strings as shown in Fig. 19.9. In this case, notice that the projector \( P_2 = 0 \) since the two terms in the projector in Fig. 19.7 are equal with opposite signs. Correspondingly note that for \( d = \pm 1 \) (and only for these values), the value of the 2-string loop is \( d_2 = 0 \) as shown in Fig. 19.8 meaning that no such 2-particle exists. Thus the only possible outcome of fusion of two 1-strings is the vacuum as shown in Fig. 19.2. Thus the entire fusion rules of these theories are

\[
1 \times 1 = 0
\]

where again 0 is the identity or vacuum. These abelian fusion rules result in abelian braiding statistics.

Two Strands in the General Case

For values of \( d \) not equal to \( \pm 1 \), the projector \( P_2 \) does not vanish. This means that two 1-strands can fuse to either 0 or 2 as shown in Fig. 19.10. We can write the fusion rule as

\[
1 \times 1 = 0 + 2
\]

We might ask whether it is possible to assemble a third type of particle with two strands. It is obvious this is not possible since \( P_2 + P_2 = I \), which means these two particle types form a complete set (\( P_2 \) projects the two particles to the vacuum, and \( P_2 \) projects to the 2-particle type).

Three Strands in the General Case

We can move on and ask what kind of particles we can make if we are allowed to fuse three strands together. We want to try to construct a three leg projector. The most general three legged operator we can construct is of the form in Fig. 19.11.

\[
\begin{align*}
P_3 &= \alpha \bigg[ 1 + \beta \bigg( \bigg. \bigg) + \gamma \bigg( \bigg) + \delta \bigg( \bigg) + \epsilon \bigg( \bigg) \bigg]
\end{align*}
\]

Fig. 19.11 The form of the most general three legged operator we can construct. Where \( \alpha, \beta, \gamma, \delta, \epsilon \) are arbitrary constants.
We would like to find the three-string operator which is a projector. So we should enforce \( P_3^2 = P_3 \). However, there are other things we want to enforce as well. Since 0 is the identity, we want \( 0 \times 1 = 1 \) which means we should not be able to fuse \( P_2 \) (the projector of two strings onto the vacuum) with a single strand to get \( P_3 \). Diagrammatically this means we must insist on relations like Fig. 19.12

\[
\begin{align*}
1 & \quad 3 \\
0 & \quad = & \quad P_3 \\
& \quad P_2 \\
& \quad = 0
\end{align*}
\]

Fig. 19.12 Insisting that \( 0 \times 1 \) does not give 3

This and analogous constraints allow us to insist on the conditions shown in Fig. 19.13

\[
\begin{align*}
P_3 & = P_3 \\
P_3 & = P_3 \\
P_3 & = P_3 \\
P_3 & = 0
\end{align*}
\]

Fig. 19.13 Four conditions that come from the fusion condition shown in Fig. 19.12

However, we should allow fusions of the form \( 1 \times 2 = 3 \) as shown in Fig. 19.14. Enforcing the condition in Fig. 19.13 along with \( P_3^2 = P_3 \) gives the form of \( P_3 \) shown in Fig. 19.11 with the results that (see Exercise 19.1)

\[
\begin{align*}
\alpha & = 1 \\
\beta & = \gamma = -\frac{d}{d^2 - 1} \\
\delta & = \epsilon = \frac{1}{d^2 - 1}
\end{align*}
\]

Fig. 19.14 We allow \( 1 \times 2 = 3 \)

We can do a short calculation in the spirit of Fig. 19.8 to obtain the value of a loop of 3-string, giving the result (See exercise 19.1) shown in Fig. 19.15

\[
d_3 = d(d^2 - 2)
\]

Fig. 19.15 Evaluating the 3-string loop.

**Ising Anyons**

In the case where \( d = \pm \sqrt{2} \) (and only in these cases) the three string loop has \( d_3 = 0 \) meaning that there is no 3-string particle. Equivalently it is possible to show that \( P_3 \) vanishes when evaluated in any diagram
Temperly-Lieb Algebra and Jones-Kauffman Anyons

(See exercise 19.1). It is similarly possible to show that \( P_1 = 0 \) and so forth. Thus, for the case of \( d = \pm \sqrt{2} \) there are only three particle types 0, 1, and 2. In addition to the fusions we have already determined, we have \( 2 \times 2 = 0 \) as shown in Fig. 19.16 and \( 2 \times 1 = 1 \) as shown in Fig. 19.17 (Note that showing \( 2 \not\in 2 \times 2 \) requires another explicit calculation, not shown here! See exercise 19.1)

We thus have the full set of nontrivial fusion rules

\[
\begin{align*}
1 \times 1 & = 0 + 2 \\
2 \times 2 & = 0 \\
1 \times 2 & = 1 \\
\end{align*}
\]

which we recognize as Ising fusion rules (see sections 8.2.2 and 18.3) where 1 = \( \sigma \) and 2 = \( \psi \) and 0 is the vacuum \( I \).

Recall in our discussion of Ising anyons in sections 8.2.2 and 18.3. There we found that \( d_\sigma = \pm \sqrt{2} \) and \( d_\psi = 1 \). This indeed agrees with the present discussion: We obtain Ising fusion rules for \( d_1 = d_\sigma = \pm \sqrt{2} \) and evaluating using Fig. 19.8 we also have \( d_\psi = d_2 = 1 \). Thus our string algebra recovers details of the Ising fusion algebra.

19.2 General Values of \( d \)

The generalization of the above discussions for \( d = \pm 1 \) and \( d = \pm \sqrt{2} \) is fairly straightforward. One can generally show the following properties (See Kauffman and Lins [1994] and exercise 19.2). First, the Jones-Wenzl projector for \( n + 1 \) strands can always be written in terms of the projector for \( n \) strands as shown in Fig. 19.19 (See exercise 19.2).

Note in particular that if \( P_n \) vanishes, we can conclude that \( P_m \) vanishes for all \( m > n \) as well. We define \( d_n \) of particle type \( n \) by connecting \( n \) strings coming from the bottom of projector \( P_n \) to those coming from the top as shown in Fig. 19.20.

Using the recursion shown in Fig. 19.19 and the definition of \( d_n \) in Fig. 19.20 we obtain the recursion relation (you can do this in your head!)

\[
d_{n+1} = dd_n - d_{n-1}
\]

(19.1)

where we define \( d_{-1} \equiv 0 \) and \( d_0 = 1 \) and hence \( d_1 = d \). This recursion has the general solution

\[
d_n = U_n(d/2)
\]

(19.2)
where $U_n$ is the $n^{th}$ Chebyshev polynomial of the second kind. These are defined by (See exercise 19.2)

$$U_n(\cos \theta) \sin \theta = \sin[(n + 1)\theta]$$  \hspace{1cm} (19.3)

A theory has a finite number of particle types if $d_n = 0$ for some $n$ (Such that $P_n$ vanishes for all $p \geq n$). This situation occurs precisely when (See exercise 19.2)

$$d = 2 \cos \left(\frac{k\pi}{n + 1}\right)$$  \hspace{1cm} (19.4)

for some $k \in 1, \ldots, n$. For values of $d$ that are not of this form, one can construct an infinite number of orthogonal particle types ($n$-strand projectors with different values of $n$), which indicates a badly behaved theory. (I.e., the algebra never “closes”).

Once one constructs the appropriate $n$-strand projectors, the general vertex between three different particle types can be constructed analogous to that shown in Fig. 19.21. Consider a vertex between particle types $(a, b, c)$ as in with $a, b, c, \geq 0$ as in Fig. 19.22. The number of strings going between the projectors (as in the fight of Fig. 19.21) is given by

$$m = \frac{(a + b - c)}{2} = \text{strings between } a \text{ and } b$$  \hspace{1cm} (19.5)

$$n = \frac{(a + c - b)}{2} = \text{strings between } a \text{ and } c$$  \hspace{1cm} (19.6)

$$p = \frac{(b + c - a)}{2} = \text{strings between } b \text{ and } c$$  \hspace{1cm} (19.7)

these quantities must be non-negative, and we must have all of these

$$\text{Fig. 19.20} \text{ Evaluating the quantum dimension of the } n\text{-string particle. We connect the } n\text{ strings coming from the top of the projector } P_n \text{ to those coming from the bottom. Often this quantity is notated as } \Delta_n.$$

By odd $n$ the $k = (n+1)/2$ case corresponds to the unphysical case of $d = 0$. 

$$\text{Fig. 19.21} \text{ The general vertex in the Temperly-Lieb algebra. Here the vertex is shown for 4 and 5 fusing to 3.}$$
Temperly-Lieb Algebra and Jones-Kauffman Anyons

quantities integer, which is assured if

\[(a + b + c) \text{ is even} \quad (19.8)\]

Note that \(a, b, \text{ or } c\) are allowed to have the value 0, meaning no strings come out that edge. These variables are also allowed to have the value 1 meaning a single string comes out the edge (and no projector is needed, see note \(2\)).

One can show that a vertex between particle types \((a, b, c)\) can be nonzero only if further if the projector

\[P_{(a+b+c)/2} \text{ is nonzero} \quad (19.9)\]

This final condition is nontrivial and we will not prove it in all generality here (See for example, Kauffman and Lins [1994], for a proof). However, Fig. 19.18 is an example of this condition: When \(d = \pm \sqrt{2}\), we’ve shown that \(P_3\) vanishes and this implies the vertex \((2, 2, 2)\) must also vanish.

The conditions we have just described for a vertex \((m, n, p)\) non-negative integers and \(P_{(a+b+c)/2}\) nonzero gives us the fusion relations for the theory which are given by

\[a \times b = |a - b|, |a - b| + 2, \ldots, \min(a + b, 2k - a - b) \quad (19.10)\]

where we have made the identification that \(a\) in the Temperly-Lieb-Jones-Kauffman theory is \(2j\). Note further that in the case where \(k\) is infinitely large (so that the final term in the series on the right of Eq. 19.10 is always \(j_1 + j_2\)), these fusion rules match the angular momentum addition rules of regular \(SU(2)\).

This value, Eq. 19.11, of the Theta diagram does not match what we would have expected given the rules in chapter 10. Comparing to Fig. 16.15 we would have expected the Theta diagram in Fig. 19.23 to have a value \(\sqrt{d_1d_2d_3}\) which in general it does not here. We will now fix this problem.

---

Note: This is entirely equivalent to the general \(SU(2)\) Chern-Simons fusion rules where particles \(j\) take integer and half-integer values and

\[j_1 \times j_2 = |j_1 - j_2|, |j_1 - j_2| + 1, \ldots, \min(j_1 + j_2, k - j_1 - j_2) \quad (19.10)\]

where we have made the identification that \(a\) in the Temperly-Lieb-Jones-Kauffman theory is \(2j\). Note further that in the case where \(k\) is infinitely large (so that the final term in the series on the right of Eq. 19.10 is always \(j_1 + j_2\)), these fusion rules match the angular momentum addition rules of regular \(SU(2)\).

With this definition of a vertex we can evaluate any planar diagram. A particularly useful diagram is the version of the Theta diagram shown in Fig. 19.23. The value of this diagram can be derived generally and is given by

\[\Delta(a, b, c) = \frac{(d_{(a+b+c)/2})!}{(d_{a-1})!(d_{b-1})!(d_{c-1})!} \quad (19.11)\]

where we have defined

\[(d_n)! \equiv d_n d_{n-1} d_{n-2} \ldots d_2 d_1\]

with \(d_1 = d\) and \(d_0 = d_{-1} = 1\). From Eq. 19.11 we see that \(\Delta(a, b, c)\) is symmetric in exchanging any of its arguments. Further we see that the quantity vanishes when \(d_{(a+b+c)/2}\) vanishes which agrees with the condition Eq. 19.9.

While the most general derivation of Eq. 19.11 is somewhat complicated (See Kauffman and Lins [1994]), it is easy enough to confirm it is correct for a few examples (See exercise 19.3).
19.3 Unitarization

The diagrammatic algebra we have constructed so far in this chapter is a perfectly self-consistent algebra (See Kauffman and Lins [1994] for a large amount of detail of this algebra). However, this algebra does not fit the rules we have established in prior chapters. In section 19.2 we just found that the value of the Theta diagram does not match the expectation from chapter 16. If we tried to work out further details of the diagrammatic algebra, we would find other failures as well — for example, we would find the $F$-matrices to be non-unitary! Fortunately, it is not hard to modify the theory a small amount so that it fits within our existing framework from chapter 16.

Let us define a new vertex which is a constant multiple of the old vertex as shown in Fig. 19.24. We define the rescaling factor as

$$v(a, b, c) = \sqrt{\frac{\Delta (a, b, c)}{d_a d_b d_c}}$$

such that the value of the Theta diagram in Fig. 19.25 is now $\Theta(a, b, c) = \sqrt{d_a d_b d_c}$ as we expect from Fig. 16.15. It turns out that this simple modification is sufficient to make the theory fit into the framework developed in chapter 16.

19.4 $F$-matrices

We can now determine the $F$-matrices directly from the graphical algebra. As a simple example, consider the $F$-matrices $F^{11 \alpha}_{1 \beta}$ (which we abbreviate as $F_2^{\alpha}$) as shown in Fig. 19.26. Note that for this equation we use renormalized vertices as defined in Eq. 19.24 and notated by dots on the vertices.

This $F$-matrix equation is that of Fig. 16.3 for four incoming 1-string particles. The $F$ matrix is nontrivial since there is more than one fusion channel when we fuse the 1’s together: $1 \times 1 = 0 + 2$, so long as $d \neq \pm 1$ (in which case the 2-string particle vanishes). We can now rewrite the $F$-matrix equation in terms of string diagrams as in Fig. 19.27. Note that in Fig. 19.27 the prefactors of $d / \sqrt{d^2}$ come from the vertex renormalization factors $v(1, 1, 2)^2$, and the quantities in brackets are $P_2$ projectors which force the two strings to fuse to the 2-particle.
Temperly-Lieb Algebra and Jones-Kauffman Anyons

\[ F_{11}^{11} = F_{0}^{0} + \frac{d}{\sqrt{d^2}} F_{2}^{0} \]

\[ \frac{d}{\sqrt{d^2}} \left[ \left( -\frac{1}{d} \right) \right] = F_{2}^{0} + \frac{d}{\sqrt{d^2}} F_{2}^{0} \]

\[ = F_{11}^{11} \]

Fig. 19.26 The F-matrix in the Temperly-Lieb-Jones-Kauffman theory is unitary when use renormalized vertices, indicated by dots. Here we have abbreviated \( F_{\alpha\beta}^{11} \) as \( F_{\beta}^{\alpha} \) for brevity.

\[ \frac{d}{\sqrt{d^2}} \left[ \left( -\frac{1}{d} \right) \right] = F_{2}^{0} + \frac{d}{\sqrt{d^2}} F_{2}^{0} \]

\[ = F_{11}^{11} \]

\[ \left( \begin{array}{cc} \frac{1}{d} & \frac{\sqrt{d^2 - 1}}{d} \\ \frac{\sqrt{d^2 - 1}}{d} & -\frac{1}{d} \end{array} \right) \]

(19.12)

Note that this matrix is properly unitary for any value of \( d \). For \( d = \pm \sqrt{2} \) the matrix matches our expectation for the Ising fusion rules given in Eq. [18.31]

With similar diagrammatic calculations, we can work out the F-
matrices for any incoming and outgoing $n$-string particles. Detailed calculations are given in [Kauffman and Lins 1994]. However, note that the results given there are nonunitary expressions due to the use of unrenormalized vertices.

### 19.5 Twisting and Braiding

So far we have not yet used the braiding rules of the Kauffman bracket invariant, we have only used the loop rule and we have only considered planar diagrams. We finally can reintroduce the braiding rules for the Kauffman invariant for evaluating crossings as in Fig. 2.3 and thus we are now considering a full anyon theory. As shown in Fig. 2.6 comparing to Fig. 15.3 we see that the twist factor of the single strand is

$$\theta_1^* = -A^{-3}.$$ 

It is a reasonably straightforward exercise to use these crossing rules to evaluate the twist factors for other particles in the theory (See exercise 19.5), as well as the $R$-matrices. Just to do a simple example, let us evaluate $R_{11}^{21}$ as shown in Fig. 19.28.

$$R_{21}^{11} = A^{-1}$$

We have already seen that in order to have a well-behaved theory with a finite number of particle types, $d$ must take some very special values (Eq. 19.4). Also recall from Fig. 2.3 that $d$ is related to $A$ via

$$d = -A^2 - A^{-2}$$

This means that we must have (with $k = 1, \ldots, n$)

$$-A^2 = \exp \left( \pm i\pi k/(n+1) \right)$$
or

\[ A = \exp \left( \pm 2\pi i \left[ \frac{1}{4} \pm \frac{k}{4(n + 1)} \right] \right) \]  

where we have labeled the two different ± with two different superscripts to keep track of them. Note that the first ± (with the superscript 1), just changes the overall chirality of the theory.

Let us take, for example, the \( n = 3 \) case, meaning \( P_3 \) vanishes and we have Ising fusion rules as discussed in section 19.1. There are two possible values of \( k \) given by \( k = 1 \) corresponding to \( d = \sqrt{2} \) and \( k = 3 \) corresponding to \( d = -\sqrt{2} \). Each of these has four possible choices of the signs in Eq. 19.13, thus resulting in eight possible anyon theories. This matches the eight theories with Ising fusion rules that we found in section 18.3 above.

Just for completeness, let us determine the twist factors for the \( \sigma \) particle for each of these anyon theories. We have

\[ \theta_a^* = -A^3 = \exp \left( \pm 2\pi i \left[ -\frac{1}{4} \pm \frac{k}{16} \right] \right) \]

with \( k = 1 \) or 3. This gives all the possible odd-sixteenth roots of unity as in Eq. 18.43.

Further Reading

- some of the ideas date back to [Penrose 1971]

Exercises

Exercise 19.1 Jones-Wenzl projectors \( P_0, P_2, \) and \( P_3 \)

For two strands one can construct two Jones-Wenzl projectors \( P_0 \) and \( P_2 \) as shown in Fig. 19.5.\(^{1,7}\)

(a) Show that these projectors satisfy \( P^2 = P \), so their eigenvalues are 0 and 1. Further show that the two projectors are orthogonal \( P_0P_2 = P_2P_0 = 0 \). (should be easy, we did this in lecture)

(b) Show that for \( d = \pm 1 \) we have \( P_2 = 0 \) in the evaluation of any diagram. The result means that in these models there is no new particle which can be described as the fusion of two elementary anyons. Why should this be obvious? Hint: Look back at the exercise 2.2.

(c) The three strand Jones-Wenzl projector must be of the form shown in the figure 19.11.
The coefficients $\alpha, \beta, \gamma, \delta, \epsilon$ are defined by the projector condition $P_3^2 = P_3$ and also by the condition that $P_3$ is orthogonal to $P_0$ which is shown in the Figs. 19.12 and 19.13.

Calculate the coefficients $\alpha, \beta, \gamma, \delta$ in $P_3$. Calculate the quantum dimension $d_3$ shown in Fig. 19.15.

(d) Choosing $d = \pm \sqrt{2}$ show that $P_3 = 0$ in the evaluation of any diagram. We can then conclude that in this model there is no new particle that is the fusion of three elementary strands. Hint: Try putting $P_3$ within a some simple diagrams and calculate the results.

(e) For the case of $d = \pm \sqrt{2}$ show that, when evaluated in any diagram, $2 \times 2 \notin \mathbb{R}$. In other words, prove Fig. 19.18.

Exercise 19.2 More General Jones-Wenzl Projectors

(a) A Jones-Wenzl projector for $n$ strands is defined both by $P_n^2 = P_{n+1}$ as well as by being orthogonal to $P_0$ analogous to Fig. 19.13. Assuming these properties are satisfied for $P_n$ show that they are satisfied for $P_{n+1}$ given by Fig. 19.19. Hint: Use the fact that connecting up a single string from $P_{n+1}$ from top to bottom as in Fig. 19.29 must give something proportional to $P_n$ (Why?).

(b) Using Fig. 19.19 derive Eq. 19.1. Show that the solution to this equation is given by Eqs. 19.2 and 19.3. Confirm the condition for $d_n$ to vanish given in Eq. 19.4.

Exercise 19.3 Theta Diagram

(a) Show $\Delta(a + 1, a, 1) = d_{a+1}$. Hint: Use Fig. 19.29.

(b) More generally show $\Delta(a + k, a, k) = d_{a+1}$. Hint: Generalize Fig. 19.29 to the case where $k$ strands are connected in a loop from the top to the bottom.

Exercise 19.4 $F$-matrix diagrammatics

Using the diagrammatic algebra, determine $F_{12}^{1a}$ and $F_{21}^{2a}$ for arbitrary $d$. Confirm that your results are unitary matrices.

Exercise 19.5 Twists of Kauffman Anyons

Use the Kauffman bracket rules to calculate $\theta_a$ for the $P_a$ kauffman anyon. Show that

$$\theta_a^* = (-1)^a A_a^{-a(a+2)}$$

Hint: Try $a = 2$ then $a = 3$ to figure out the pattern.
Anyons from Groups

In this chapter we will use the structure of discrete groups to build rules for anyon theories\(^1\). There are two general approaches to consider. In section 20.1 we will label the lines in our diagrams with group elements whereas in section 20.2 we will label lines in our diagrams with group representations. As we have done previously, in each case we will consider planar diagrammatic algebras first before considering braiding.

\(^1\)In chapter *** (also section ***), we discuss another construction of an anyon theory from a discrete group, known as the quantum double construction. We defer discussion of that construction for now.

20.1 Fusion as Group Multiplication

One way to construct a wide variety of consistent planar diagrammatic algebras is to construct our fusion rule based on the structure of a group. In this approach we consider a discrete group \(G\), and each element \(g \in G\) is a particle type with the identity element \(I\) of the group being the vacuum.

Fusion rules follows the rules for group multiplication. That is, for \(g, h \in G\)

\[ g \times h = gh \]

which we draw as shown in Fig. 20.1.

Since \(gg^{-1} = g^{-1}g = I\), antiparticles are given by the inverse elements in the group, or \(\bar{g} = g^{-1}\). This means that in a diagram we may reverse an arrow if we invert the group element as shown in Fig. 20.2.

Let us consider diagrams where each line is labeled by a group element \(g \in G\). Reversal of a line corresponds to inversion of the group element as shown in Fig. 20.2, analogous to reversing an arrow in order to turn a particle into its antiparticle.

In cases where the group is abelian so that \(g \times h = gh = hg = h \times g\) which is what we required for fusion of particle types in section 8.1 above.

In section 20.1.3 we will consider the possibility of using nonabelian groups, but for now we will assume the group is abelian. We thus have fusion rules given by group multiplication

\[ N_{g,h}^{\alpha} = \delta_{\alpha,gh} = \delta_{\alpha,hg} \]

Since the result of any fusion is always uniquely defined by group multiplication (one never has a sum on the right hand side, such as \(g \times h = a + b\)), the quantum dimension of every particle is \(d_{g} = 1\) meaning the Hilbert space size does not grow with the number of particles.

An example of a planar diagram with this type of group multiplication is shown in Fig. 20.3.
20.1.1 Group Cohomology

We now have the task of trying to construct consistent $F$-matrices for our planar diagram algebra. This is an extremely well studied problem in the field of group cohomology.

Consider a general group $G$. A so-called 3-cocyle of the group is given by a function of three variables $\omega(a, b, c)$ where $a, b, c \in G$ that satisfies

$$\omega(a, b, c)\omega(a, bc, d)\omega(b, c, d) = \omega(ab, c, d)\omega(a, b, cd)$$

(20.1)

Generally we will consider cases of $\omega$ being a $U(1)$ valued complex phase. In group cohomology notation we say that

$$\omega \in H^3(G, U(1))$$

(20.2)

Eq. (20.1) may look obscure, but it is actually just a translation of the pentagon equation! Let us make the identification, in the notation of chapter 9

$$[F_{(abc)}^{a,b,c}]_{(ab),(bc)} = \omega(a, b, c)$$

So that we have diagrammatically

Examinining the pentagon equation Eq. 9.7 and Fig. 9.7 we see that this is precisely the same as Eq. 20.1 in a different language. Note that
there is no sum over indices here (like the sum over possible elements \( h \) in Eq. 9.7) since the fusion of any two group elements always gives a unique group element as an outcome.

As with \( F \)-matrices, it is possible to choose different gauges (See section 9.4). In particular given a 3-cocycle (ie., a solution of the pentagon equation) we can multiply each \( a, b \) vertex by a phase \( u(a, b) \) as shown in Fig. 20.5 to transform the cocycle by

\[
\omega(a, b, c) \rightarrow \frac{u(a, bc)u(b, c)}{u(a, b)u(ab, c)} \omega(a, b, c).
\]

By making such a gauge transform we generate additional solutions of the pentagon equation. We view different solutions which are gauge transforms of each other as being physically equivalent, We will typically work with just one representative 3-cocycle for each equivalence class by choosing a convenient gauge. It is useful to always work with a so-called normalized gauge, where \( \omega(a, b, c) = 1 \) whenever \( a = I \) or \( b = I \) or \( c = I \). (I.e, fusing with the vacuum gives no phase). Further we want to only consider gauge transforms that maintain this normalized gauge, so we must insist on \( u(I, g) = u(g, I) = u(I, I) = 1 \). Given this restriction to normalized gauge, however, one still has a large additional gauge freedom.

The 3-cocycle (pentagon) equation Eq. 20.1 typically will have more than one gauge-inequivalent solution. Further, if we have two different 3-cocycles \( \omega \) and \( \omega' \), we may multiply these together to generate another solution \( \omega \omega' \) and we may invert \( \omega \) to generate another solution. Thus, the space of 3-cocycles \( H^3(G, U(1)) \) in Eq. 20.2 is itself a group, known as the third cohomology group of \( G \) with coefficients in \( U(1) \).

A trivial 3-cocycle \( \omega(a, b, c) = 1 \) for all \( a, b, c \in G \) is always possible. In this case all diagrams have value 1. However, for any group (beyond the trivial group with only one element), there are always other possible 3-cocycles as well. Such 3-cocycles and group cohomology in general have been studied extensively in the mathematics and physics communities and it is possible to simply look up the form of the possible 3-cocycles. (See the end of the chapter for good references).

While all 3-cocycles provide a solution to the pentagon equation, they do not always allow for full isotopy invariance as discussed in chapter 16. Indeed, for any 3-cocycle \( \omega \), we will need to check whether it satisfies all the requirements for full isotopy invariance. For example, we want to be able to freely turn up and down legs of a vertex as shown in Fig. 20.6.

Thus for full isotopy invariance (and allowing for \( d \) both +1 and −1) we need to have

\[
\begin{align*}
\text{s}(a, b)\omega(a, a^{-1}, b) &= 1, \\
\text{s}(a, b)\omega(a, b^{-1}, b) &= 1
\end{align*}
\]
for all $a, b$ in the group with

$$s(a, b) = \begin{cases} 
-1 & d_a = d_b = -1 \\
+1 & \text{otherwise}
\end{cases} \quad (20.6)$$

While this condition seems quite restrictive, the gauge freedom Eq. 20.3 allows us often to achieve this.

\[a \xrightarrow{a^{-1}} b\]

\[ab \xrightarrow{b^{-1}} b\]

\[a \xrightarrow{a^{-1}b} a\]

\[ab^{-1} \xrightarrow{b^{-1}} b\]

\[s(a, b)\omega(a, a^{-1}, b)\]

\[s(a, b)\omega(a, b^{-1}, b)^{-1}\]

A further item to note is that

$$d_a = \omega(a, a^{-1}, a)$$

and if $a$ is self-dual ($a = a^{-1}$) this is the gauge invariant Frobenius-Schur indicator. If we can do so, we will try to choose a gauge for our 3-cocycles such that we have full isotopy invariance (this is not always possible) by choosing negative $d_a$ but keeping $\epsilon_a$ positive as discussed in section 14.5.

20.1.2 Simple Examples with $G = \mathbb{Z}_N$

For example, let us take a simple case of the group $G = \mathbb{Z}_N$, the group of integers modulo $N$ with the group operation being addition modulo $N$. Since this group is abelian, we have $gh = hg$ as we want for fusion of particle types as described in chapter 8.

The inequivalent 3-cocycles of the group $\mathbb{Z}_N$ can be written as (See references at the end of the chapter)

$$\omega(a, b, c) = \exp \left( \frac{2\pi ip}{N^2} a(b + c - [b + c]_N) \right) \quad (20.7)$$

where here $a, b, c \in 0, \ldots, N - 1$, and the brackets $[b + c]_N$ means $b + c$ modulo $N$ where the result is chosen to lie in the range $0, \ldots, N - 1$. Here the index $p$ is an integer in the range $0, \ldots, N - 1$ describing the $N$ different gauge-inequivalent 3-cocycles.

\[\text{Fig. 20.6 Turning up and down relations (analogous to Fig. 14.17). The prefactor } s\text{ comes from the proper interpretation of the sign of the } \sqrt{d}\text{ factors in 14.17. See section 14.5.}\]

\[\text{A further item to note is that}\]

$\text{d}_a = \omega(a, a^{-1}, a)$

\[\text{and if } a\text{ is self-dual } (a = a^{-1}) \text{ this is the gauge invariant Frobenius-Schur indicator. If we can do so, we will try to choose a gauge for our 3-cocycles such that we have full isotopy invariance (this is not always possible) by choosing negative } d_a \text{ but keeping } \epsilon_a \text{ positive as discussed in section 14.5.}\]

\[\text{Confusingly } gh = g \times h \text{ here mean addition of } g \text{ and } h \text{ modulo } N.\]
The trivial 3-cocyle is given by \( p = 0 \) which gives \( \omega = 1 \) always. The nontrivial 3-cocycles are more interesting.

\( \mathbb{Z}_2 \)

Let's consider the simple case of \( \mathbb{Z}_2 \) fusion rules. Here the group elements are \( g = 0, 1 \) and the group operation is addition modulo 2. One has the trivial 3-cocycle \( p = 0 \) in Eq. 20.7 giving \( \omega = 1 \), or all \( F \)-matrix elements equal to 1, which we identify as being exactly the same as the \( d = 1 \) loop gas from section 18.1.1.

The only nontrivial 3-cocycle is the \( p = 1 \) case. Here, using Eq. 20.7 we determine the 3-cocycle is of the form

\[
\omega(a, b, c) = \begin{cases} 
-1 & \text{if } a = b = c = 1 \\
+1 & \text{otherwise}
\end{cases}
\]

We recognize this as being exactly the case of the \( d = -1 \) loop gas from section 18.1.2 (This translates to saying that the \( F \)-matrix is \(-1\) if and only if all four incoming legs \( a, b, c \) and \( abc \) are in the 1 state as in Eq. 18.10, and note that \( abc \) here means multiplication with the group operation so is really \((a + b + c) \mod 2\).

\( \mathbb{Z}_3 \) and beyond

Generalizing the \( \mathbb{Z}_2 \) fusion to \( \mathbb{Z}_3 \), we now have \( g = 0, 1, 2 \) with the group operation being addition modulo three. In this case we have three different 3-cocycles: the trivial 3-cocycle \( p = 0 \) in Eq. 20.7 and two nontrivial 3-cocycles \( p = 1 \) and \( p = 2 \) in Eq. 20.7.

While these nontrivial cocycles provide a valid solution to the pentagon equation 20.1 (or Eq. 9.7) they are not in a form where they enjoy full isotopy invariance. One can use gauge transforms Eq. 20.3 to try to put the cocycles in different forms, but it is not possible to find a gauge where both Eq. 20.4 and Eq. 20.5 are satisfied at the same time\(^4\). Nonetheless they still provide a consistent planar diagrammatic algebra, although not a fully isotopy invariant one. Thus the only isotopy invariant case is the trivial cocycle \( p = 0 \). We will discuss the possible braidings that are consistent with this planar algebra when we discuss the trivial cocycle below.

One can further show that \( \mathbb{Z}_N \) for any odd \( N \) is similar to the case of \( \mathbb{Z}_3 \): the only fully isotopy invariant case is the trivial cocycle \( p = 0 \). See exercise 20.2. And this theory has \( N \) solutions of the hexagon equation. These are given in detail in appendix 20.4.

For \( \mathbb{Z}_N \) with \( N \) even, the situation is slightly more interesting. Here, there are two possible values of \( p \) which can give an isotopy invariant solution: \( p = 0 \) (the trivial cocycle similar to what we found in the \( N \) odd case) and \( p = N/2 \), which is analogous to the nontrivial cocycle we found for the \( \mathbb{Z}_2 \) case. (See exercise 20.2). For each of these values of \( p \) there are \( N \) solutions of the hexagon equation. All of these cases are discussed in detail in appendix 20.4. Again, for other values of \( p \) we

\(^{4}\)To see that it is not possible to achieve full isotopy invariance note that from Eq. 20.4 and 20.5 isotopy invariance requires \( \omega(1, 1, 2) = \pm 1 \) and \( \omega(1, 2, 2) = \pm 1 \). However, for \( N = 3 \), the product \( \omega(1, 1, 2)\omega(1, 2, 2) \) is gauge invariant, and it is only \( \pm 1 \) for the case of \( p = 0 \). See exercise 20.2.
Anyons from Groups

have a perfectly consistent planar diagrammatic algebra, although not a fully isotopy invariant one (no matter what gauge we choose).

Other abelian groups

Abelian groups are always\(^5\) of the form \(\mathbb{Z}_{N_1} \times \mathbb{Z}_{N_2} \times \ldots\) with some number of factors of cyclic groups \(\mathbb{Z}_N\). We can look up the cocycles for such groups in, for example, de Wild Propitius [1995] or Hu et al. [2013b] or a book on group cohomology! Note that the variety of different possible cocycles grows when there are multiple \(\mathbb{Z}_N\) factors. We will not pursue these theories further here\(^6\).

It looks like a fun exercise to explore this!

We have a bit of a language difficulty here. Here we use the word nonabelian to mean when \(g \times h \neq h \times g\) whereas previously (See section 8.2) we used non-abelian to describe fusion rules where there is more than one fusion channel, such as \(g \times h = a + b + \ldots\).

20.1.3 Using Nonabelian Groups?

In the case where the group is nonabelian we deviate from what was done when we discussed fusion of particle types in section 8.1 above. In the discussion of fusion of particle types, we have always assumed \(g \times h = h \times g\) and with a nonabelian\(^7\) group \(gh\) may not be the same as \(hg\).

Why did we insist in chapter 8 that particle fusion should satisfy \(g \times h = h \times g\)? If we think about particles living in three dimensions, when we bring two particles, \(g\) and \(h\) together, looking at the system from one angle it looks like \(g\) is to the right of \(h\) but looking at the two particles from another angle, it looks like \(h\) is to the right of \(g\). Thus there is no way to decide whether the pair fuses to \(gh\) or \(hg\).

However, if we are only concerned with a planar diagram algebra (or a diagram algebra on the surface of sphere) then there is no ambiguity! The surface we are considering is assumed to be oriented so we can always unambiguously decide which particle is clockwise of which other particle at a vertex. Thus we can make the general rule that for a vertex to be an allowed fusion, the three particles leaving the vertex must multiply in clockwise order to the identity as shown in the right of Fig. 20.1. Thus, at least for planar diagrams we can generalize our rules for particle fusion to allow non-commutative fusions.

All of the figures we have drawn in this section (Fig. 20.1 – Fig. 20.6) have been drawn so as to be consistent with our rule for nonabelian groups — that is, if all of the arrows are outgoing, when you multiply the group elements clockwise around the vertex you obtain the identity.

For definiteness we discuss the example of the nonabelian group \(S_3\) in the appendix section 20.5.

20.2 Fusion of Group Representations: \(\text{Rep}(G)\)

Another way to construct a consistent planar diagrammatic algebra is to work with representations of discrete groups\(^8\). Suppose we have irreducible representations \(R_i\) of a group \(G\). A tensor product of two of these irreducible representations will necessarily decompose into a direct sum of irreducible representations. I.e., we have\(^9\)

\(\text{If we write } M \otimes N = P \text{ we mean the following. If } M_{ab} \text{ is a matrix of dimension } m \text{ and } N_{cd} \text{ is a matrix of dimension } n \text{ then } P \text{ is defined as } P_{(ac),(bd)} = M_{ab}N_{cd} \text{ and is of dimension } mn. \text{ If we write } P = N \oplus M \text{ we mean that } P \text{ is block diagonal with blocks } N \text{ and } M. \text{ Finally note that the relation in Eq. 20.9 is an isomorphism not an equality. One can choose a basis such that the right hand side is block diagonal, however, this is not the natural basis for the left.} \)
\[ R_a \otimes R_b \simeq R_c \oplus R_d \oplus \ldots \]  (20.9)

with the sum on the right hand side being finite. We thus propose to label a particle type for our diagrammatic algebra with an irreducible group representation, and have the fusion relations be given by these tensor product decompositions. Thus we interpret the tensor product equation Eq. 20.9 as a particle fusion relation

\[ a \times b = c + d + \ldots \]

and accordingly particle \( a \)'s corresponding to representation \( R_a \) has an antiparticle \( \bar{a} \) corresponding to the dual representation which we write as \( \bar{R}_a = R_a^* \). This fusion category (this set of fusion rules with the associated \( F \)-matrices) using the representations of the group \( G \) is known as \( \text{Rep}(G) \).

It is fairly easy using some tricks of group theory to determine the fusion rules for discrete group representations. Recall that a representation \( R \) is a homomorphism from each group element \( g \) to a matrix \( \rho_R(g) \). Meaning a mapping where the group operation is preserved:

\[ \rho_R(g_1) \rho_R(g_2) = \rho_R(g_1 g_2) \]  (See section 41.2.4). The trace of the representation matrix is known as its character \( \chi_R(g) = \text{Tr}[\rho_R(g)] \).

One can either work out the characters of a group explicitly or (much more commonly) just look them up on character tables, which can be found in any group theory book or on the web.

Since \( \text{Tr}(ab) = \text{Tr}(ba) \) we have \( \chi_R(g) = \chi_R(hgh^{-1}) \) meaning that the character depends only on the so-called conjugacy class of the group element \( g \).

Characters combine in fairly simple ways under both direct product and direct sum

\[ \chi_{R_a \oplus R_b}(g) = \chi_{R_a}(g) + \chi_{R_b}(g) \]  (20.10)
\[ \chi_{R_a \otimes R_b}(g) = \chi_{R_a}(g) \chi_{R_b}(g) \]  (20.11)

Further we have orthonormality relations for irreducible representations:\n
\[ \frac{1}{|G|} \sum_{g \in G} [\chi_{R_a}(g)]^* \chi_{R_b}(g) = \delta_{R_a,R_b} \]  (20.12)

where the sum is over all elements \( g \) of the group \( G \) and \(|G|\) is the total number of elements in the group. We can thus deduce the tensor product decomposition:\n
\[ R_a \otimes R_b \simeq \bigoplus_{c \in \text{irreps}} N_{a \tilde{b}}^{c} R_c \]  (20.13)

\(^{10}\) Meaning a mapping where the group operation is preserved: \( \rho^R(g_1)\rho^R(g_2) = \rho^R(g_1g_2) \).

\(^{11}\) This orthonormality is derived trivially from the grand orthogonality theorem, Eq. 41.6. Since the character \( \chi(g) \) is a function of only the conjugacy class of \( g \) it is sometimes more convenient to replace the sum over all elements with a sum over classes where we then also include a factor of the number of elements in the class. So the left hand side would read instead

\[ \sum_{\text{classes } C} \frac{|C|}{|G|} [\chi_{R_a}(g \in C)]^* \chi_{R_b}(g \in C) \]

with \(|C|\) meaning the number of elements in class \( C \).

\(^{12}\) The \( \bigoplus \) symbol here means a direct sum of all the arguments. The prefactor \( N_{a \tilde{b}}^{c} \), here means the \( R_c \) representation occurs \( N_{a \tilde{b}}^{c} \) times in the direct sum.

\(^{13}\) We have \( R_a \otimes R_b \simeq R_b \otimes R_a \) meaning the two tensor products are isomorphic, but they are not equal. The two matrices have their entries in different places. See the definition in note 4 of this chapter above.
Anyons from Groups

Table 20.1. Character table for the group $S_3$. Notice the orthogonality of rows as defined by Eq. 20.12

<table>
<thead>
<tr>
<th></th>
<th>identity 1 element</th>
<th>rotations 2 elements</th>
<th>reflections 3 elements</th>
</tr>
</thead>
<tbody>
<tr>
<td>trivial rep ($I$)</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>sign rep ($S$)</td>
<td>1</td>
<td>1</td>
<td>-1</td>
</tr>
<tr>
<td>2d rep ($V$)</td>
<td>2</td>
<td>-1</td>
<td>0</td>
</tr>
</tbody>
</table>

where

$$N_{ab}^c = \frac{1}{|G|} \sum_{g \in G} [\chi^R_c(g)]^* \chi^R_a(g) \chi^R_b(g)$$  \hspace{1cm} (20.14)

or in our fusion product language

$$a \times b = b \times a = \sum_c N_{ab}^c$$

Note that in the case where the group is abelian, the representations themselves are also an abelian group (meaning $N_{ab}^c = N_{ba}^c \in \{0, 1\}$ only.)

It is not hard to show (See exercise **) that the quantum dimension of a representation $R_a$ is given by

$$d_a = \chi^{R_a}(e)$$  \hspace{1cm} (20.15)

where $e$ is the identity element of the group.

20.2.1 Example: Representations of $S_3$

As a simple example, let us consider the representations of the group $S_3$ which can also be thought of as the symmetries of a triangle. To remind the reader this group has 6 elements which can be written in terms of two generators $X$ (a reflection) and $R$ (a rotation) with multiplication rules $X^2 = R^3 = e$ and $XR = R^{-1}X$ with $e$ the identity. The 6 elements can be written as $e, R, R^{-1}, X, XR, XR^{-1}$. There are three conjugacy classes, which we will call the identity ($e$), the rotations ($R, R^{-1}$), and the reflections ($X, XR, XR^{-1}$).

There are also three irreducible representations. The group has a character table as given in table 20.1. It is then easy to use Eq. 20.14 to determine the fusion laws for the representations, which are given by

$$I \times I = I, \hspace{1cm} I \times S = S, \hspace{1cm} I \times V = V$$  \hspace{1cm} (20.16)
$$S \times S = I, \hspace{1cm} S \times V = V$$  \hspace{1cm} (20.17)
$$V \times V = I + S + V$$  \hspace{1cm} (20.18)

from which we see that $I$ plays the role of the vacuum particle. Just as an example, let us consider Eq. 20.18. From the character table we have $\chi^V = (2, -1, 0)$ and so $\chi^V \otimes ^V \chi^V = \chi^V \chi^V = (4, 1, 0) = (1, 1, 1) + (1, 1, -1) +$
Table 20.2 Character table for the group $Q_8$. Notice the orthogonality of rows as defined by Eq. 20.12.

$$(2, -1, 0) = \chi^I + \chi^S + \chi^V.$$ 

### 20.2.2 Example: Quaternion Group $Q_8$

The quaternion group $Q_8$ can be defined as the eight two-by-two matrices

\[
\begin{bmatrix}
\pm 1, \pm i\sigma_x, \pm i\sigma_y, \pm i\sigma_z
\end{bmatrix}
\]

The group has five conjugacy classes 1, $-1$, \{$\pm i\sigma_x$\}, \{$\pm i\sigma_y$\}, \{$\pm i\sigma_z$\}, and correspondingly five representations. The character table is given in table 20.2. From the character table it is easy to use Eq. 20.14 to derive the nontrivial fusion rules (Again $I$ plays the role of the vacuum particle and we do not write its fusions)

\[
R_i \times R_i = I \quad i = x, y, z \quad (20.19)
\]

\[
S \times R_i = S \quad i = x, y, z \quad (20.20)
\]

\[
R_x \times R_y = R_z \quad \text{(and cyclic permutations)} \quad (20.21)
\]

\[
S \times S = I + R_x + R_y + R_z \quad (20.22)
\]

Note that $S$ is the 2-dimensional representation given by the defining two-by-two matrices.

### 20.2.3 F-Matrices

With a bit of work, the $F$-matrices (often known as $6j$ symbols in this context) can also be derived using group theoretic methods. In general this can be a bit complicated but the principle is straightforward group theory. As usual we should think of $F^{{bad}_{eff}}$ as a basis transform (See Fig. 16.3). In this case it is convenient to think of the process of $b, a, e$ and $c$ fusing to the identity in different ways or equivalently, the tensor product of $R_b, R_a, R_e$ and $R_c$ fusing to the identity representation.

1. Consider $R_b \otimes R_b \simeq \bigoplus_d N^d_{ab} R_d$ and fuse with $R_c \otimes R_c \simeq \bigoplus_d N^d_{cc} R_d$. The resulting representations, $R_d$ and $R_d$ then fuse together to form the identity representation. Such a process corresponds to the diagram in the top of Fig. 20.7 (same as the left of Fig. 16.3).

2. Consider instead $R_b \otimes R_c \simeq \bigoplus_f N^f_{bc} R_f$ and fuse with $R_a \otimes R_c \simeq \bigoplus_f N^f_{ac} R_f$, and finally fuse $R_f$ and $R_f$ to form the identity representation. Such a process corresponds to the diagram on bottom

$16$ The quaternions were famously discovered by Hamilton. He was so excited by this discovery that he carved them into the stone of Brougham (Broom) Bridge in Dublin. There is a plaque there today to commemorate this event.
essence representing the braid operator by $B$
paths through the hexagon must give the same result. Realizing that the
as minus the identity. $\rho$
pared to the fully bosonic solution
section 18.3.1, are not consistent with a trivial braiding. The fact that $R_{a} \otimes R_{b}$
we are in
the hexagon is called $\text{Rep}(G)$,

$\rho_{a,b} = 1$ (20.23)
provides a solution$^{18}$ of the hexagon for $\text{Rep}(G)$ for any group $G$. If we
choose this braiding we are describing particles that are bosons — i.e.,
having trivial braiding$^{19}$ and trivial spin $\theta_{a} = 1$, with internal quantum
numbers given by the representations of the group $G$.

Fermions and Bosons
If it so happens that $G$ contains an central element$^{20}$ $z$ such that $z^{2} = e$
with $e$ the identity of the group, then each representation $R_{a}$ can be
assigned a degree $p(a)$ which is 1 or 0 depending on whether $z$ acts as
the identity in the representation or acts as $-1$ in the representation$^{21}$. A
consistent braiding is then given by (See exercise 20.5)

$R_{a \times b}^{a,b} = (-1)^{p(a)p(b)}$ (20.24)

In other words, we have declared a particle $a$ to be if bosonic if $p(a) = 0$
(if $R_{a}(z)$ acts as 1) or fermionic $p(a) = 1$ (if $R_{a}(z)$ acts as $-1$).$^{22}$

Both of these processes correspond to fusion of the four representa-
tions to the identity. The first, we might say is the identity component
of $(R_{b} \otimes R_{a}) \otimes (R_{c} \otimes R_{e})$ whereas the second is the identity component
of $(R_{b} \otimes R_{c}) \otimes (R_{a} \otimes R_{e})$. While these two tensor products are isomor-
phic, they are expressed in a different basis (see note$^{13}$ above). To find
the $F$-matrix $F_{abcd}^{\text{bos}}$ matrix relating these bases we simply have to find
the overlap between the $d$ contribution to the overall identity representa-
tion in case (1) above with the $f$ contribution to the overall identity
representation in case (2).

20.2.4 Some Simple Braidings for $\text{Rep}(G)$
So far we have only discussed consistent fusion of representations — i.e., fusion rules that will satisfy the pentagon equation. Given a set of
$F$-matrices, we can then look for braidings, or $R$-matrices that satisfy the
hexagon. We already know that there will typically be multiple solutions$^{17}$ of the hexagon. Some of the braidings, however, can be
stated easily for any group $G$, as we now discuss.

Trivial Braidings: Bosons
It is always the case that the trivial braiding

$R_{a \times b}^{a,b} = 1$ (20.23)

provides a solution$^{18}$ of the hexagon for $\text{Rep}(G)$ for any group $G$. If we
choose this braiding we are describing particles that are bosons — i.e.,
having trivial braiding$^{19}$ and trivial spin $\theta_{a} = 1$, with internal quantum
numbers given by the representations of the group $G$.

Fermions and Bosons
If it so happens that $G$ contains an central element$^{20}$ $z$ such that $z^{2} = e$
with $e$ the identity of the group, then each representation $R_{a}$ can be
assigned a degree $p(a)$ which is 1 or 0 depending on whether $z$ acts as
the identity in the representation or acts as $-1$ in the representation$^{21}$. A
consistent braiding is then given by (See exercise 20.5)

$R_{a \times b}^{a,b} = (-1)^{p(a)p(b)}$ (20.24)

In other words, we have declared a particle $a$ to be if bosonic if $p(a) = 0$
(if $R_{a}(z)$ acts as 1) or fermionic $p(a) = 1$ (if $R_{a}(z)$ acts as $-1$).$^{22}$

sometimes this braided solution to
the hexagon is called $\text{Rep}(G,z)$ com-
pared to the fully bosonic solution
which can then be called $\text{Rep}(G,e)$. I.e., if you set $z = e$, the identity, then
you set all particles to be bosons.

$^{18}$Not every set of fusion rules is consistent with the trivial braiding. (For example, the Ising fusion rules, as discussed in
section 14.3.4) are not consistent with a trivial braiding). The fact that $\text{Rep}(G)$ always allows a trivial braiding is a subtle,
but straightforward statement about using tensor products of representations to represent fusion of particle types. We are in
essence representing the braid operator by $B(R_{a} \otimes R_{b}) = R_{a} \otimes R_{b}$, i.e., we just re-order the tensor factors without adding any
phase. Realizing that the $F$-matrices are also just relating different re-orderings of tensor factors, this tells us that the two
paths through the hexagon must give the same result.

Fig. 20.7 Fusion of representations. Compare to the diagrams in Fig. 16.3.

The top figure can be thought of as the
$d$ component of $R_{b} \otimes R_{e} \simeq \bigoplus_{j} N_{ab}^{j} R_{j}$
fused with $R_{a} \otimes R_{e} \simeq \bigoplus_{j} N_{ab}^{j} R_{j}$ to form the identity. The bottom figure
can be thought of as the $f$ component of $R_{b} \otimes R_{e} \simeq \bigoplus_{j} N_{ab}^{j} R_{j}$
fused with $R_{a} \otimes R_{e} \simeq \bigoplus_{j} N_{ab}^{j} R_{j}$ to form the identity.

Recall that if $G$ is abelian, then the
representations of $G$ are isomorphic to
the group $G$ itself. In which case
$\text{Rep}(G)$ is just the trivial cocycle of $G$.
As we discussed in section 20.1.4 and
20.24 the trivial cocycle of $Z_{N}$ has $N$
different braidings.

Note that $R_{ab}^{b}$ is gauge dependent,
so Eqs. 20.23 and 20.24 imply a gauge
choice. However, $R_{ab}^{b} R_{bc}^{b}$ is gauge
independent, and $\theta_{a}$ is gauge independent.

$^{19}$“Central” means that $z$ commutes
with all the elements of the group. I.e.,
$zg = gz$ for all $g \in G$.

$^{20}$To see this consider applying $z$ in
the representation twice $p^{R}(z)p^{R}(z) = p^{R}(z^{2}) = p^{R}(e)$. This means that
$p^{R}(z)$ can only act as the identity or
as minus the identity.

$^{21}$Sometimes this braided solution to
the hexagon is called $\text{Rep}(G,z)$ com-
pared to the fully bosonic solution
which can then be called $\text{Rep}(G,e)$. I.e., if you set $z = e$, the identity, then
you set all particles to be bosons.

$^{22}$Not every set of fusion rules is consistent with the trivial braiding. (For example, the Ising fusion rules, as discussed in
section 14.3.4) are not consistent with a trivial braiding). The fact that $\text{Rep}(G)$ always allows a trivial braiding is a subtle,
The fermionic particles have \( \theta_a = -1 \) whereas the bosonic particles have \( \theta_a = +1 \).

An nontrivial example of \( \text{Rep}(G, z) \) is given by the quaternion group \( \mathbb{Q}_8 \) where we choose \( z = -1 \). It is easy to check that this makes the \( R_i \) representations bosons, but the \( S \) representation is a fermion (note the negative sign on the character \( \chi^S(-1) \)).

### 20.2.5 Continuous (Lie) Group Representations?

One can imagine that instead of looking at the representations of discrete groups, one considers instead the representations of Lie groups (See section 41.2.3). For example, the different representations of the group \( SU(2) \) are the different values of the spin quantum number \( j \), and these fuse together with the usual angular momentum addition rules. Further, the \( F \)-matrices are (up to a normalization) precisely what we call \( 6j \) symbols of angular momentum addition.

While such a scheme makes a perfectly good planar diagrammatic algebra, the problem is that there are an infinite number of different representations (For the case of \( SU(2) \) for example, the angular momentum \( j \) can infinitely large) and this violates our rule of having a finite number of “particle types” for our diagrammatic algebra. Such algebras can be problematic when used for physical purposes (For example, as we will see in section 21.3 using a diagrammatic algebra with an infinite number of representations for construction of a TQFT results in divergences). Schemes have been constructed to regularize such a diagrammatic algebra and arrange that only a finite number of representations ever occur; such schemes are often known as “deformations” of the Lie algebra representation\(^\text{23}\). The most common such deformations correspond precisely to the particle types of a corresponding Chern-Simons theory at some finite level. For example, in the case of \( SU(2) \), one can consider \( SU(2)_k \) Chern-Simons theory which has deformed \( F \)-matrices such that angular momentum \( j = 0, 1/2, \ldots k/2 \) can occur, but one never gets any higher angular momenta.

### 20.3 Parastatistics Revisited

Way back in section 3.5.1 we asked why we could not have exotic statistics in 3+1 dimensions. While there are nontrivial representations of the permutation group that would satisfy the quantum mechanical composition rule, we stated that additional constraints — such as particle creation and annihilation and locality — limits us to just bosons and fermions. We are now at the point where we can discuss exactly what we mean by this.

The structure we have built up for anyons in 2+1 dimensions is that of a braided unitary category: a set of particles with fusions, \( F \)-matrices satisfying the pentagon, and \( R \)-matrices satisfying the hexagon. If we try to do something similar in 3+1 dimensions we will no longer have nontrivial braiding of world lines since, as discussed in section 3.3.2 in

\[ b \quad a \]

\[ \quad = \quad \]

![Fig. 20.8](image) In 3+1 dimensions these two pictures are topologically equivalent. Thus all particles are transparent. This implies Eq. 20.23

\(^\text{23}\)The term “quantum group” is often used. Be warned that a quantum group is not a group.
264 Anyons from Groups

3+1 dimension no knots can be formed in one-dimensional world lines. Thus, we must impose the restriction shown in Fig. 20.8 that all particles are transparent. In equations this can be stated as

$$R_{c}^{ab} R_{c}^{ba} = 1$$  \hspace{1cm} (20.25)$$

for all $a,b,c$ such that $N_{ab}^{c} > 0$. If the condition Eq. 20.25 holds and yet we have a solution of the hexagon equation, we say we have a \textit{symmetric tensor category}\textsuperscript{24}. Thus if we are to construct an anyon theory with point particles in 3+1 dimensions, it must be described by a symmetric tensor category.

In fact we have already given two examples of symmetric tensor categories, in section 20.2.4: (a) the theory $\text{Rep}(G)$ (or $\text{Rep}(G,e)$) which describes bosons having internal quantum numbers given by the representations of the group $G$, and (b) the theory $\text{Rep}(G,z)$ where some of the particles are instead declared fermions depending on how their corresponding representation transforms under the action of the element $z$. The crucial theorem we mentioned in section 3.5.1, originally due to Doplicher and Roberts (See also M"{u}ger [2007]; Deligne [2002]), is that there are no other possibilities: Any symmetric tensor category is equivalent to $\text{Rep}(G) \equiv \text{Rep}(G,e)$ if it has no fermions, or $\text{Rep}(G,z)$ if it has fermions. In other words for point particles in 3+1 dimensions, there are only bosons and fermions; nothing else!

20.4 Appendix: Isotopy Invariant Planar Algebras and Anyon Theories from $G = \mathbb{Z}_N$ Cohomology

Here we start with the cocycles for $\mathbb{Z}_N$ given in Eq. 20.7 and we look for isotopy invariant cases. We have two general types of isotopy invariant solutions: the $p = 0$ (trivial cocycle) solution, which exists for any $N$ and the $p = N/2$ case which exists for even $N$ only. Let us discuss each of these in a bit more detail.

20.4.1 Trivial Cocycle: $\mathbb{Z}_N^{(n)}$ Anyons

For any $\mathbb{Z}_N$ one can always choose $p = 0$ in Eq. 20.7 which gives the trivial cocycle $\omega(a,b,c) = 1$ for all $a,b,c$ and we correspondingly have $d_{a} = 1$ for all $a$.

Recall that these cocycles are really $F$-matrices, which have now all been set to unity. We now want to determine the possible braidings for this theory by using the hexagon Eq. 13.1. Plugging in $F = 1$ into the hexagon, we obtain\textsuperscript{25}

$$R_{[c + a]_{N}}^{c,a} R_{[c + b]_{N}}^{c,b} = R_{[c + a + b]_{N}}^{c,[a + b]_{N}}$$  \hspace{1cm} (20.26)$$

There are exactly $N$ solutions\textsuperscript{26} of this system which we label $n = \ldots$
0, \ldots, N - 1 which are given by
\[ R_{[a+b]}^{a,b} = \exp \left[ \frac{2\pi in}{N} ab \right] \]
where \( ab \) on the right is actual multiplication, not the group operation which is addition modulo \( N \). The twist factors are (using Eq. [15.2])
\[ \theta_a = e^{2\pi ina^2/N} \]
and the corresponding \( S \) matrix is (perhaps easiest derived with Eq. [17.20])
\[ S_{a,b} = \frac{1}{\sqrt{N}} \exp \left[ \frac{4\pi in}{N} ab \right] \]
This is a modular theory only for \( N \) odd with \( n \) and \( N \) mutually prime.

These theories are sometimes known as \( Z_N^{(n)} \) anyons (See Bonderson [2007]).

### 20.4.2 Nontrivial Cocycle: \( Z_{N=2p}^{(n)} \)

Here we consider \( N = 2p \). The cocycle in Eq. [20.7] again has \( N \) consistent solutions of the hexagon equations, given by \( n = 0, \ldots, (N - 1) \) in the equation (See Bonderson [2007])
\[ R_{[a+b]}^{a,b} = \exp \left[ \frac{2\pi i(n + \frac{1}{2})[a]_N [b]_N}{N} \right] \]
with all \( d_a = +1 \), resulting in (from Eq. [15.1])
\[ \theta_a = \exp \left[ \frac{2\pi i(n + \frac{1}{2})}{N} [a]_N^2 \right] \]
\[ S_{a,b} = \frac{1}{\sqrt{N}} \exp \left[ \frac{4\pi i(n + \frac{1}{2})}{N} [a]_N [b]_N \right] \]

These are known as \( Z_N^{(\frac{n}{2} + \frac{1}{2})} \) anyon theories for obvious reasons. They are modular when \( 2n + 1 \) and \( p \) are coprime.

These anyon theories, in the gauge given by Eq. [20.7] are not generally isotopy invariant. We can generally make transformations to put these results in potentially simpler forms.

#### Case I: \( p \)-odd

With \( N = 2p \) and \( p \) odd, the \( p^{th} \) particle is self dual particle, and has Frobenius-Schur indicator \(-1\), which is a gauge invariant quantity. We will thus need to push this sign onto \( d \) in order to have a fully isotopy invariant theory. As discussed in section [14.5] we choose a gauge where
\[ d_a = (-1)^a \]

27For \( n = (N - 1)/2 \) the modular theory matches the Chern-Simons theory \( SU(N)_1 \) with \( N \) odd.

28When \( n = (N/2 - 1) \) the modular theory matches \( SU(N)_1 \) with \( N \) even, and when \( n = 0 \) the modular theory matches \( U(1)_{N/2} \). Be cautioned that there is some disagreement in the literature as to how you label the level of a \( U(1) \) Chern-Simons theory.
If we try to use the same rule for $N = 2p$, with $p$ even where we still set $d_a = (-1)^a$, this actually gives us the trivial cocycle discussed above in a less convenient gauge. Note that the self-dual particle has $d = +1$ in the $p$ even case indicating that we do not need to push any signs onto $d$.

The hexagon equation now takes the form

$$R^{c,a}_{[c+a]N} R^{c,b}_{[c+b]N} = \omega(a, b, c) R^{c,[a+b]N}_{[a+b]N}$$

which compared to Eq. 20.26 introduces a minus sign if all $a, b, c$, are odd. This system of equations again has $N$ solutions which we index as $\bar{n} = 0, \ldots, N - 1$,

$$R^{a,b}_{[a+b]N} = \exp \left[ \frac{2\pi i \bar{n}}{N} [a]_N [b]_N \right] (i)^{r(a,b)}$$

where $r(a, b) = 1$ if both $a$ and $b$ are odd and equals zero otherwise. It is a short exercise to show that this recovers the correct $\theta_a$, Eq. 20.27, and $S$-matrix Eq. 20.28 with the mapping $\bar{n} = n - (p - 1)/2$. Again, the advantage of using this gauge for the description of $Z_{N^+}^{(n+1)/2}$ anyons is full isotopy invariance (with $N = 2p$ and $p$ odd).

**Case II: $p$-even**

In this case the Frobenius-Schur indicator of the self-dual particle is $+1$ so we can choose to work in a gauge where all $d_a = +1$. While it is possible to make a gauge transform that puts the theory into an isotopy invariant form, the transform is not particularly transparent. For this reason it is often convenient to stay with the gauge given in Eq. 20.7. However, it is not too hard to transform to an isotopy invariant gauge if we would like.

An example of this, let us consider the nontrivial cocycle for the group $Z_4$. Here we can make a gauge transform (See exercise 20.2) such that

$$\omega(a, b, c) = \begin{cases} 
-1 & (a, b, c) = (1, 1, 1); (1, 2, 3); (2, 1, 2); \\
(2, 3, 2); (3, 2, 1); (3, 3, 3) \\
1 & \text{otherwise} \end{cases}$$

and with $d = 1$ for all particles 0,1,2,3. This gives a fully isotopy invariant theory. There are correspondingly 4 solutions of the hexagon given by $n = 0, 1, 2, 3$ with

This $n$ is the same as that of Eq. 20.28
Appendix: Cocyles for $S_3$

To give an example of a non-abelian group, let us look at the case of the group $S_3$. To remind the reader this group has 6 elements which can be written in terms of two generators $X$ and $R$ with multiplication rules $X^2 = R^3 = e$ and $XR = R^{-1}X$ with $e$ the identity. The 6 elements can be written as $e, R, R^{-1}, X, XR, XR^{-1}$. Let us write them as $(A,a) = X^A R^a$ with $A = 0, 1$ and $a = -1, 0, 1$. There are 6 independent 3-cocycles described by $p = 0, \ldots, 5$ in the equation (See references at the end of the chapter)

$$\omega((A,a), (B,b), (C,c)) = \exp \left( \frac{i\pi p}{9} (B + C^{-1} a \{ (B + C^{-1} a)^3 \} \right)$$

where the bracket $[.]_3$ indicates modulo 3 where the result is assumed to be in the range $-1, 0, 1$.

Note that within $S_3$ there is a $\mathbb{Z}_2$ subgroup consisting of $e$ and $X$, or $a = 0$ with $A = 0, 1$. The first term on the right hands side, $\exp(i\pi pABC)$, matches the two possible 3-cocyles from the $\mathbb{Z}_2$ group. For even $p$ it is the trivial cocycle, whereas for odd $p$ we have a $\omega$ being $-1$ only when $A, B, C$ are all in the 1 state, equivalent to Eq. [20.8] The second factor looks similar to the $\mathbb{Z}_3$ cocycles but only when $C = 0$. Setting $C = 0$ for a moment, the same argument as in the $\mathbb{Z}_3$ case shows that we cannot have full isotopy invariance unless $p = 0$ or $p = 3$, in which case the second factor on the right hand side of Eq. [20.31] is trivial. Thus this case of $p = 3$ gives an isotopy invariant cocyle which essentially ignores the $a$ variable of $(A,a)$ and is equivalent to Eq. [20.8] for the $A$ variables with $d(A,a) = (-1)^A$.

Details of Working out an $F$-matrix

Here we give a few more details of how we work out the $F$-matrix for a group. Other, potentially more systematic schemes, can be found in Refs. Hamermesh [1989]; Buerschaper and Aguado [2009]; Wang et al. [2020] for example.

To implement our procedure, we work with explicit $D$ dimensional matrices $\rho^R_{ij}$ for each $D$ dimensional unitary representation $R$. We extract the identity component of the fusion of the four particles by writing

$$R^{a,b}_{[a+b]_4} = \exp \left[ \frac{2\pi i (n + \frac{1}{2})}{4} [a][b] \right] (-1)^{r(a,b)}$$

where $r(a,b) = 1$ for $(a, b) = (1, 2), (1, 3), (2, 1), (3, 1)$ only and is zero otherwise.
Anyons from Groups

\[
\sum_g \rho^R_a(g) \otimes \rho^R_b(g) \otimes \rho^R_c(g) \otimes \rho^R_e(g) = C \sum_d w_d w_d^\dagger (20.32)
\]

where \( w_d \) is a unit length orthogonal \( D_a D_b D_c D_e \) dimensional vector representing the process where \( a \) and \( b \) fuse to \( d \) and also \( c \) and \( e \) fuse to \( d \), and \( C \) is an unimportant normalization constant. (Note there is a gauge choice here in choosing the phase of \( w_d \)). Similarly we can write

\[
\sum_g \rho^R_a(g) \otimes \rho^R_c(g) \otimes \rho^R_b(g) \otimes \rho^R_e(g) = C \sum_f z_f z_f^\dagger (20.33)
\]

where \( z_f \) is a \( D_a D_b D_c D_e \) dimensional vector representing the process where \( a \) and \( c \) fuse to \( f \) and also \( b \) and \( e \) fuse to \( f \). In the second line of Eq. 20.33, the matrix \( P \) is simply a permutation matrix since the two tensor products are isomorphic and simply have rows and columns appropriately permuted.

The \( F \)-matrix is then just given by the overlap

\[
F_{ce \mid f} = w_d^\dagger \cdot z_f (20.34)
\]

The challenge is then simply to extract the correct vectors \( w_d \) and similarly \( z_f \) in Eq. 20.32 and 20.33.

We thus only need to build up the tensor product in Eq. 20.32 and 20.33 step by step. We have

\[
\rho^R_a(g) \otimes \rho^R_b(g) = \sum_{d \in a \times b} \sum_{\alpha, \beta = 1}^D \left( \sum_{\alpha} x^{\bar{d}}_\alpha \left[ \rho^R_{\bar{d}}(g) \right]_{\gamma \delta} [x^{\bar{d}}_\alpha]_{\gamma \delta}^\dagger \right) \otimes \left( \sum_{\beta} y^{d}_\beta \left[ \rho^R_d(g) \right]_{\gamma \delta} [y^{d}_\beta]_{\gamma \delta}^\dagger \right) (20.35)
\]

where the \( x \)'s are a set of orthonormal \( D_a D_b \) dimensional vectors (both sides of this equation are \( D_a D_b \) dimensional matrices)\(^{32} \). The particular form of the \( x \) vectors can be extracted using the grand orthogonality theorem Eq. 41.6. Performing the same decomposition for

\[
\rho^R_c(g) \otimes \rho^R_e(g) = \sum_{d \in c \times e} \sum_{\gamma, \delta = 1}^D \left( \sum_{\gamma} y^{\bar{d}}_\gamma \left[ \rho^R_{\bar{d}}(g) \right]_{\gamma \delta} [y^{\bar{d}}_\gamma]_{\gamma \delta}^\dagger \right) \otimes \left( \sum_{\delta} y^{d}_\delta \left[ \rho^R_d(g) \right]_{\gamma \delta} [y^{d}_\delta]_{\gamma \delta}^\dagger \right) (20.36)
\]

To find the identity element of the fusion between the tensors in Eq. 20.35 and 20.36 we simply match up the \( d \) representations with the \( \bar{d} \) representations. Thus we have

\[
w_d w_d^\dagger \sim \sum_{g} \sum_{\alpha, \beta, \gamma, \delta = 1}^{D_d} \left( x^{d}_\alpha \left[ \rho^R_d(g) \right]_{\gamma \delta} [x^{d}_\alpha]_{\gamma \delta}^\dagger \right) \otimes \left( y^{d}_\delta \left[ \rho^R_d(g) \right]_{\gamma \delta} [y^{d}_\delta]_{\gamma \delta}^\dagger \right) (20.37)
\]
where in going to the second line we have used the grand orthogonality theorem Eq. [41.6]. Both sides of this equation are \( D_a D_b D_c D_e \) dimensional matrices with a single nonzero eigenvalue, so it is then trivial to extract \( w_d \). Then extracting \( z_f \) by permuting rows of the vector \( w_d \) as noted in Eq. [20.33] we then determine the \( F \)-matrix using Eq. [20.34].

The procedure outlined here is fairly straightforward, although tedious. Equivalent schemes are outlined in Refs. [Buerschaper and Aguado 2009] and [Wang et al. 2020]. In exercise 20.6 we walk through calculation of \( F \)-matrices for the representations of the group \( S_3 \) whose fusion rules we worked out in Eqs. [20.10] [20.18] above.

As a final comment we note that there is a beautiful expression to calculate the Frobenius-Schur indicator \( \kappa \) for a group representation \( R \)

\[
\kappa_R = \frac{1}{|G|} \sum_g \chi^R(g^2)
\]

(20.38)

which gives \( \pm 1 \) if \( R \) is self-dual, and gives 0 for a non-self-dual representation. Recall that the Frobenius-Schur indicator is actually one of the elements of the \( F \)-matrix (See Eq. [14.3]). Derivation of this formula is a standard result in group theory (See, for example, [Hamermesh 1980]).

**Further Reading**

**Exercises**

**Exercise 20.1 Cocycle Equation**

(a) Show that the 3-cocycle given by Eq. [20.7] satisfies cocycle condition Eq. [20.1] and thus represents a valid cocycle.

(b) Show that Eq. [20.31] also satisfies Eq. [20.1].

**Exercise 20.2 Isotopy Invariance of Cocycles**

(a) Show that the cocycle Eq. [20.1] can only represent an isotopy invariant diagram algebra only in the following cases:

- For \( n \) an odd integer, only when \( p = 0 \).
- For \( n \) an even integer, only when \( p = 0 \) or \( p = n/2 \)

(b) For the case of \( n = 4 \) and \( p = 2 \), find the gauge transformation that transforms Eq. [20.1] into Eq. [20.18].

**Exercise 20.3 Quantum Dimension of a Representation**

Prove Eq. [20.15]. Hint: Remember that that quantum dimension \( d_a \) tells you how the Hilbert space dimension grows as you fuse together the particle \( a \) many times. Try fusing together many representations \( R^a \otimes R^a \otimes R^a \ldots \) and imagine decomposing the result into irreducible representations using the orthogonality theorem for characters. Note that for characters \( \chi(e) \geq \chi(g) \) for the identity representation.

**Exercise 20.4 Frobenius-Schur Indicators in \text{Rep}(G)**
It may be useful to use a computer to perform matrix manipulations in this problem and lab. octave, and python are all fairly easy to use.

**Exercise 20.5 Bosons and Fermions in \text{Rep}(G, z)**

Let $z$ be a central element of the group $G$ (i.e. $zg = gz$ for all $g \in G$) such that $z^2 = e$, the identity. As in section 20.2.4 for a representation $R_a$ set $p(a) = 0$ if $z$ acts as the identity in representation $R_a$ and set $p(a) = 1$ if $z$ acts as $-1$ in representation $R_a$.

(a) Show that if $R_a \otimes R_b = R_c \oplus \ldots$, then $p(a)p(b) = p(c)$. Hint, consider the characters $\chi^R(e)$ and $\chi^R(z)$.

(b) Given that setting all particles to bosons (i.e., Eq. 20.23) solves the hexagon equation, show that Eq. 20.24 also provides a solution to the hexagon equation.

**Exercise 20.6 Some $F$-matrix elements for representations of $S_3$ [Hard]**

Let us consider the simplest nonabelian group $S_3$, which we discuss in sections 20.3, 20.2.1 and 21.2.1.

We remind the reader that this group has 6 elements which can be written in terms of two generators $X$ and $R$ with multiplication rules $X^2 = R^3 = e$ and $XR = R^{-1}X$ with $e$ the identity. The 6 elements can be written as $e, R, R^2, X, XR, XR^2$ which are grouped into conjugacy classes $\{e\}, \{R, R^2\}, \{X, XR, XR^2\}$ (See Table 20.1).

The three representations are as follows: The trivial representation has $\rho^t(g) = 1$ for all $g$ in the group. The sign rep has $\rho^s(g) = 1$ for $g \in \{e, R, R^2\}$ and $\rho^s(g) = -1$ for $g \in \{X, XR, XR^2\}$. (Note that since both these reps are one dimensional, they are completely defined by the character table). We write the two dimensional representation in a unitary form as

$$
\rho^V(X) = \begin{pmatrix} -1 & 0 \\ 0 & 1 \end{pmatrix}, \quad \rho^V(R) = \frac{1}{2} \begin{pmatrix} 1 & \sqrt{3} \\ -\sqrt{3} & 1 \end{pmatrix}
$$

with $\rho^V(e)$ the identity matrix and all other matrices $\rho^V(g)$ for the other elements $g$ in the group can be generated by using the group multiplication properties. (Do this first, you will need it later) \footnote{It may be useful to use a computer to multiply matrices (Mathematica, mat lab, octave, and python are all fairly convenient), since there are a lot of matrix manipulations in this problem and a single error will destroy the result.}

Note that we already know the fusion rules for these representations as they are given in Eqs. 21.16, 21.18.

In this exercise we will calculate some $F$-matrix elements by focusing on the most interesting case, where all four incoming lines in Fig. 16.3 are in the two dimensional $V$ representation. Thus we are interested in the unitary matrix $F^VV_f$.

(a) Using the grand orthogonality theorem (Eq. 41.6) find the decomposition

$$
\rho^V(g) \otimes \rho^V(g) = x^f[x^f]^\dagger + \rho^S(g)x^S[x^S]^\dagger + \sum_{\alpha,\beta=1}^2 x^\alpha_{\alpha'\beta'} [\rho^{R_{\alpha'}(g)}]_{\alpha'\beta'} [x^{\text{V}_f}]_{\beta'}^\dagger
$$

HINT: The one dimensional representations are easy to obtain since they can be obtained by

$$(1/|G|) \sum_g \rho^R(g)^* \rho^V(g) \otimes \rho^V(g)$$
what remains is the two dimensional representation.

(b) Given two dimensional matrices $A, B, C, D$ find the permutation matrix such that

$$P(\mathbf{A} \otimes \mathbf{B} \otimes \mathbf{C} \otimes \mathbf{D})P^T = \mathbf{A} \otimes \mathbf{C} \otimes \mathbf{B} \otimes \mathbf{D}$$

(c) Use Eq. 20.34 and 20.37 to show that the $F$-matrix is given by

$$F^d \mathbf{V} = \frac{1}{2} \begin{pmatrix}
1 & 1 & \sqrt{2} \\
1 & 1 & -\sqrt{2} \\
\sqrt{2} & -\sqrt{2} & 0
\end{pmatrix}$$
State Sum TQFTs

Having learned about planar diagrammatic algebras we are now in a position to explicitly construct a real 3D TQFT\(^1\). There are several steps in this idea. We start by considering a closed 3D manifold \(\mathcal{M}\) which we discretize into tetrahedra (a so called simplicial decomposition of the manifold). Next we construct a model, similar in spirit to statistical mechanics, which sums a certain weight over all quantum numbers on all edges of all tetrahedra. The weights being summed are defined in terms of our planar diagrammatic algebra as we will see below. The result of this sum is the desired TQFT partition function \(Z(\mathcal{M})\) which we discussed extensively above, and particularly in chapter 7.

This discretization of a manifold into tetrahedra is very commonly used in certain approaches to quantum gravity, which we will discuss in section 21.3.

21.1 Simplicial Decomposition and Pachner Moves

We start by considering a so-called simplicial decomposition of our manifold. Such decompositions can be made of smooth manifolds in any number of dimensions\(^2\).

21.1.1 Two Dimensions

As a warm up let us think about two-dimensional manifolds. In two dimensions, the elementary 2-simplex is a triangle, so this decomposition is the familiar idea of triangulation shown in Fig. 21.1.

Since we are only concerned with the topology of the manifold, not the geometry, the precise position of vertex points we use is irrelevant — only

---

\(^1\)The input for the construction in chapter will be a planar diagram algebra — we do not have to specify any sort of \(R\)-matrix or braiding. It is a bit surprising that one only needs a planar algebra to make a 3D TQFT! In section 21.2 we will input a spherical tensor category whereas in section 21.4 we will input a group and a 3-cocycle.

\(^2\)It is interesting (but beyond the scope of this book) that manifolds exist in dimension \(d \geq 4\) that cannot be smoothed, and cannot be decomposed into simplicies.

---

Fig. 21.1 Some triangulations of 2-manifolds
The connectivity of the points is important, i.e., the topological structure of the triangulation network. Furthermore, a particular manifold, like a sphere, can be triangulated in many different ways. It turns out that any two different triangulations can be related to each other by a series of elementary “moves” known as two-dimensional Pachner moves\(^{3,4}\), which are shown in Figs. 21.2 and 21.3.

Thus if we want to construct a manifold invariant (like \(Z(M)\) we discussed in chapter 7) with a manifold represented in terms of a triangulation we only need to find some function of the triangulation that is invariant under these two Pachner moves.

### 21.1.2 Three Dimensions

The story is quite similar in three dimensions. Since we have been focused on 2+1 dimensional TQFTs we will mostly discuss three-dimensional manifolds. We discretize any closed three-dimensional manifold\(^{5}\) by breaking it up into tetrahedra (otherwise known as three-dimensional simplices). Any two discretizations are topologically equivalent to each other if they can be related to each other by a series of three-dimensional Pachner moves\(^{6}\), which are shown in Figs. 21.4 and 21.5. Again, the key point here is that if we can find some function of the the network structure that is invariant under the Pachner moves, we will have constructed a topological invariant of the manifold.

---

3 I encourage you to play with these two moves and see how you can restructure triangulations by a series of Pachner moves.

4 It is interesting to note that a Pachner moves can be thought of as viewing a 3D tetrahedron from two opposite directions. We can thus think of 2D Pachner moves as a *cobordism* (See chapter 7 in 3D between a surface triangulated with the initial triangulation and a topologically equivalent surface triangulated with the final triangulation.

5 For now let us focus on closed manifolds. We briefly discuss manifolds with boundary in section 21.2.2.

6 Analogous to the 2D case (see note 3 above), the 3D Pachner moves can be thought of as viewing a 4D-simplex (a so-called pentachoron) from two opposite directions.
The Turaev-Viro State Sum

The idea of the Turaev-Viro state sum is to build a 3D manifold invariant from one of the planar diagrammatic algebras we have been discussing in chapters 8-20.

First, let us choose any particular planar diagrammatic algebra. We take any decomposition of an orientable three dimensional manifold into tetrahedra. Let each edge of this decomposition be labeled with one of the quantum numbers (the particle labels) from the diagrammatic algebra. We then consider the following sum

\[ Z_{TV}(M) = D^{-2N_e} \sum_{\text{all edge labelings}} W(\text{labeling}) \]  

(21.1)

As we have been doing all along, when we label an edge with a quantum number we must put an arrow on the edge unless the particle type is self-dual.
where \( N_v \) is the number of vertices in the decomposition, and
\[
D = \sqrt{\sum_n |d_n|^2}
\]
is the total quantum dimension (See Eq. 17.11). In Eq. 21.1, \( W \) is a weight assigned to each labeling of all the edges. We consider the following definition of a weight assigned to a given labeling of edges
\[
W(\text{labeling}) = \prod_{\text{tetrahedra}} \tilde{G}(\text{tetrahedron}) \prod_{\text{edges}} d_{\text{edge}} \prod_{\text{triangles}} \tilde{\Theta}(\text{triangle})
\]
(21.2)
Thus each tetrahedron is given a weight \( \tilde{G} \), depending on its labeling, each edge labeled \( a \) is given a weight \( d_a \) and each triangle is given a weight \( \tilde{\Theta}^{-1} \) depending on its labeling.

The weights \( \tilde{G} \) and \( \tilde{\Theta} \) are very closely related to quantities \( G \) and \( \Theta \) we have already studied in chapter 16 for example. The functions \( \tilde{G} \) and \( \tilde{\Theta} \) are given by
\[
\tilde{\Theta}(c,b,a) = \Theta(a,b,c) = \sqrt{d_a d_b d_c}
\]
(21.3)
and
\[
\tilde{G}(d,e,c,f) = G_{bad} = \frac{F_{ecf} d_f}{\sqrt{d_e d_c}}\sqrt{d_a d_e d_f}
\]
(21.4)
Note that the tetrahedron shown here is different from the one shown in Eq. 21.4 on the other hand \( e,f,c \) meet at a point where \( f,e,\bar{a} \) form a loop. In Eq. 21.4 the three edges around any face must fuse together to the vacuum. I.e., we have the four conditions
\[
N_{bad} > 0 \quad N_{cde} > 0 \quad N_{fca} > 0 \quad N_{efb} > 0
\]
or else \( \tilde{G} \) will vanish. Note that, like \( G \), the value of \( \tilde{G} \) is unchanged under any rotation of the tetrahedron. It is important to note that \( Z_{TV} \) is gauge invariant. While we can make a gauge transformation on the \( F \)-symbols (and hence \( \tilde{G} \)) as in Eq. 16.23, for example, such gauge factors will cancel out in the overall weight Eq. 21.2 (See exercise 21.4).
21.2.1 Proof Turaev-Viro is a Manifold Invariant

The proof that \( Z_{TV}(\mathcal{M}) \) is a manifold invariant is not difficult — one only needs to show that it is unchanged under the 1-4 and 2-3 Pachner moves. This is basically an exercise in careful bookkeeping (see exercise 21.2). Roughly, however, it is easy to see how it is going to work.

Let us first examine the 2-3 Pachner move shown in Fig. 21.5. On the left we have two tetrahedra (call them 1 and 2) which are joined along a triangle (call it \( \alpha \)). On the right we have three tetrahedra (call them 3, 4 and 5 which are joined along three triangles (call them \( \beta, \gamma, \) and \( \delta \)) with the three triangles intersecting along a new edge down the middle (shown vertical in the figure) which we label with the quantum number \( n \). To show that the \( Z_{TV} \) remains invariant we need to show that

\[
\tilde{G}(1) \tilde{G}(2) \tilde{\Theta}(\alpha) = \sum_n \tilde{G}(3) \tilde{G}(4) \tilde{G}(5) \tilde{\Theta}(\beta) \tilde{\Theta}(\gamma) \tilde{\Theta}(\delta) d_n
\]

The factors of \( \tilde{\Theta} \) are simply factors of \( \sqrt{d_a} \) and these cancel some factors of \( \sqrt{d_a} \) in the definition of \( \tilde{G} \) in Eq. 21.3. After this cancellation what remains is a relationship between two \( F \)'s on the left and a sum over three \( F \)'s on the right. The relationship that remains is exactly the pentagon equation Eq. 16.3 (or Eq. 9.7)! Thus any diagrammatic algebra which satisfies the pentagon equation will result in a Turaev-Viro partition function (Eq. 21.1) that is invariant under the 2-3 Pachner move!

The case of the 1-4 Pachner move is only a bit harder and we will sketch the calculation here. The large tetrahedra on the left of Fig. 21.4 (lets call this large tetrahedron 1) needs to be equivalent to the four smaller tetrahedra on the right (lets call these small tetrahedra 3, 4, 5 and 6) once we sum over the quantum numbers on the four internal edges on the right. The three tetrahedra 3, 4 and 5 share a common edge, and this is entirely analogously to the three tetrahedra we considered in the case of the 2-3 Pachner move. Summing over the quantum number of this common edge, and using the same pentagon relation replaces the three tetrahedra 3, 4, 5 with two tetrahedra 1 and 2, where 1 is the large tetrahedron and 2 includes exactly the same edges as the remaining small tetrahedron 6. The tetrahedra 2 and 6 have 3 edges which are not shared with tetrahedron 1 — these are the remaining internal edges that need to be summed over. Summing over one of these internal edges, one invokes the consistency condition Eq. 16.5 to create a delta function which then kills one of the two remaining sums. The last remaining sum just yields a factor of \( D^2 = \sum_n d_n^2 \) which accounts for the prefactor in Eq. 21.1 being that we have removed one vertex from the lattice.

21.2.2 Some TQFT Properties

The Turaev-Viro state sum has all the properties we expect of a TQFT. Although we need to discretize our manifold, the resulting “partition function” \( Z_{TV}(\mathcal{M}) \) for a manifold \( \mathcal{M} \) is a complex number which is indeed independent of the discretization and depends on the topology.
of the manifold only.

As we discuss at length in section 7.1 we would also like $Z_{TV}(\mathcal{M})$ to represent a wavefunction if $\mathcal{M}$ is a manifold with boundary. To remind the reader, the point of this construction is that when we glue together two manifolds with boundary to get a closed manifold, this corresponds to taking the inner product between the two corresponding wavefunctions to get a complex number.

To see how this occurs let us consider discretizing a manifold with a boundary. Here the 3D bulk of the manifold $\mathcal{M}$ should be discretized into tetrahedra, and the 2D boundary surface $\Sigma = \partial \mathcal{M}$ should be discretized into triangles. We divide the edge degrees of freedom into bulk and boundary where a boundary edge is defined as an edge where both vertices are on the boundary and all other edges are defined to be bulk. We define $Z(\mathcal{M})$ of such a discretized manifold with boundary as a sum like Eq. 21.1 where the sum is only over the edges in the bulk, leaving fixed (un-summed) the quantum numbers for the edges that live entirely on the boundary (i.e., both vertices on the boundary). Thus for manifolds with boundary we more generally write

$$Z_{TV}(\mathcal{M}; a_1, \ldots, a_N) = \mathcal{D}^{-2N_v - n_v} W'(a_1, \ldots, a_N) \sum_{\text{bulk labelings}} W(\text{bulk labels})$$

where $N_v$ is the number of vertices in the bulk and $n_v$ the number of vertices on the boundary. The weight function $W$ is exactly the same as the weight function in Eq. 21.2 but only including edges, triangles, and tetrahedra in the bulk (All tetrahedra are considered bulk, and a triangle is considered boundary only when all three vertices are on the boundary). Here $a_1, \ldots, a_N$ are the quantum numbers of the edges on the boundary, and these are not included in the sum over bulk labels. An additional weight is included which is a function of these boundary edge labels

$$W'(a_1, \ldots, a_N) = \frac{\prod_{\text{boundary edges}} d_{\text{edge}}}{\prod_{\text{boundary triangles}} \Theta(\text{triangle})}$$

The partition function $Z_{TV}(\mathcal{M}; a_1, \ldots, a_N)$ is now a function of the edge variables and is interpreted as a wavefunction\(^{12}\) $|Z(\mathcal{M})\rangle$ that lives on the boundary $\Sigma = \partial \mathcal{M}$.

It is then quite natural to see how two manifolds can be glued together along a common boundary as in Fig. 7.3. In that figure we have a closed manifold $\mathcal{M} \cup_{\Sigma} \mathcal{M}'$ where $\mathcal{M}$ and $\mathcal{M}'$ are manifolds with boundary joined along their common boundary $\Sigma = \partial \mathcal{M} = [\partial \mathcal{M}']^*$. When we glue together $\mathcal{M}$ and $\mathcal{M}'$ we obtain the partition function for the full manifold as in Eq. 7.4 where we obtain the inner product by summing over the degrees of freedom of the wavefunction — which in this case means summing over the quantum numbers $a_1, \ldots, a_N$ of the edges on
21.3 Connections to Quantum Gravity Revisited

the boundaries. In other words, we have

\[ Z_{TV}(M \cup_S M') = \langle Z_{TV}(M') | Z_{TV}(M) \rangle \]
\[ = \sum_{a_1, \ldots, a_N} [Z_{TV}(M'; \bar{a}_1, \ldots, \bar{a}_N)]^* Z_{TV}(M; a_1, \ldots, a_N) \quad (21.5) \]
\[ = \sum_{j_1, \ldots, j_N} Z_{TV}(M'; a_1, \ldots, a_N) Z_{TV}(M; a_1, \ldots, a_N) \quad (21.6) \]

where in the second line the edge variables in the first term are inverted because the surface of \( M' \) has the opposite orientation from the surface of \( M \). Going from the second to third line is an easy exercise (See exercise 21.1). The final result is easily seen to be the correct expression for the Turaev-Viro invariant for the full manifold \( M \cup M' \). I.e., it now sums over all the quantum numbers in both bulks and on the common boundary.

As in section 7.2 on can generalize the idea of a TQFT to include particle world lines (labeled links) as well as the space-time manifold \( M \). As mentioned there we can roughly think of these world lines as internal boundaries, and we just fix the quantum number of edges along these hollow tubes to describe different world-line types. (See references at the end of the chapter).

21.2.3 Connection to Chern-Simons Theory

There is a remarkable connection between the Turaev-Viro partition function and Chern-Simons theory. If we build a Turaev-Viro theory from the \( F \)-matrices of a Chern-Simons theory, it turns out that the partition function of the Turaev-Viro theory is related to that of the Chern-Simons theory via\(^{13}\)

\[ Z_{TV}(M) = |Z_{CS}(M)|^2 \]

This is known as the Turaev-Walker theorem [Turaev 1992, 1994; Walker 1991], and we will outline its derivation in section 22.3.2 below. The TQFT resulting from Turaev-Viro is sometimes known as the quantum double of the input theory of \( F \)-matrices (the input category) and this formula gives at least one reason why the name is appropriate — Turaev-Viro is equivalent to two copies of the Chern-Simons theory (of opposite handedness).

21.3 Connections to Quantum Gravity Revisited

The Turaev-Viro invariant is a natural descendant of one of the very earliest approaches to quantum gravity pioneered by Penrose [1971] and Ponzano and Regge [1968]. Indeed much of the continued interest in Turaev-Viro and similar state-sum invariants is due to this relationship.
An interesting approach to macroscopic general relativity, used for example, in numerical simulation, is to discretize space-time into simplices — tetrahedra in three dimensions or four dimensional simplices (sometimes known as pentachora) in four dimensions. The curvature of the space-time manifold (the metric) is then determined by the lengths assigned to the edges.

If one then turns to quantum gravity, one wants to follow the Feynman prescription and perform a sum over all possible metrics as we discussed previously in section 6.1. We can write a quantum partition function as

$$Z = \int \mathcal{D}g \, e^{iS_{\text{Einstein}}[g]/\hbar} \quad (21.7)$$

We can imaging performing such a sum for a discretized system by integrating over all possible lengths of all possible edges. However, not all triangle edge lengths should be allowed — in Euclidean space one must obey the crucial constraint of the triangle inequality

$$|l_1 - l_2| \leq l_3 \leq (l_1 + l_2). \quad (21.8)$$

The key observation is that the triangle inequality is precisely the same as the required inequality for regular angular momentum addition

$$j_1 \otimes j_2 = |j_1 - j_2| \oplus |j_1 - j_2| + 1 \oplus \ldots \oplus |j_1 + j_2|. \quad (21.9)$$

Thus it is natural to label each edge of with a quantum mechanical spin, and sum over all possible spins. Such an approach is known as a spin network. We thus imagine building a Turaev-Viro model (Eq. 21.1) with a planar diagrammatic algebra built from angular momentum addition rules: quantum numbers are the angular momenta $j$, the fusion rules are as given in Eq. 21.9, and the $F$-matrices are given by the regular 6j symbols of angular momenta addition.

As mentioned in section 21.2.3 above, the Turaev-Viro model built from the $SU(2)_k$ diagrammatic rules is equivalent to Chern-Simons theory $SU(2)_k \otimes SU(2)_{-k}$. As we mentioned in section 6.2 above, such a Chern-Simons theory is equivalent to 2+1D gravity with a cosmological constant $\lambda = (4\pi/k)^2$. Taking the limit of large $k$ then gives the classical limit of simple $SU(2)$ angular momentum addition corresponding to a universe with no cosmological constant.
21.4 Dijkgraaf-Witten Model

Another state sum model of some interest is the so-called Dijkgraaf-Witten model\(^{20}\) (Dijkgraaf and Witten \(^{1990}\)). As with Turaev-Viro this model discretizes space into simplices and sums over possible labels of all the edges.

In the Dijkgraaf-Witten model we choose a group \(G\) and we label the edges of the simplices with elements from that group. The general idea is very similar to that of Turaev-Viro just using the multiplication properties of the group to give us a set of fusion rules as in section 20.1 and we use a 3-cocycle in place of the \(F\)-matrix\(^{21}\). These fusion rules require that multiplication of the group elements around every triangle must result in the identity as shown in Fig. 21.6. This is the analog of Eq. 21.3 where three quantum numbers around a triangle must fuse to the identity. This condition is known as a “flatness” condition, with the name coming from lattice gauge theory, which we will see in more detail in chapter ***.

\[
\begin{align*}
  c &\triangle b \\ 
  \Rightarrow \quad cba &= \text{identity}
\end{align*}
\]

Fig. 21.6 Multiplying group elements around a triangle in Dijkgraaf-Witten theory results in the identity. This is known as the “flatness” condition

As mentioned in section 20.1 when we use group multiplication for fusion rules, the quantum dimensions\(^{22}\) of all the particles are all \(d_a = 1\). This means that in Eq. 21.2 both the \(d_a\) factor and the \(\tilde{\Theta}\) factor are trivial. We are thus left with only the tetrahedron factor and the Dijkgraaf-Witten partition function looks like a simplified version of the Turaev-Viro case in Eqs. 21.1 and 21.2 given by\(^{23}\)

\[
Z_{DW}(\mathcal{M}) = |G|^{-N_v} \sum_{\text{labelings}} \prod_{\text{tetrahedra}} \tilde{G}(\text{tetrahedron}) \quad (21.10)
\]

where \(N_v\) is the number of vertices, \(|G|\) is the number of elements in the group \(G\), and the sum is only over labellings that satisfy the flatness condition (Fig. 21.6).

The tetrahedral symbol \(\tilde{G}\) is a bit more complicated than in the case of the Turaev-Viro invariant. We do not generally have full tetrahedral symmetry so it could matter which way we orient the tetrahedron when we evaluate \(\tilde{G}\). In order to define the tetrahedral symbol \(\tilde{G}\) properly we do the following: First we label each vertex in the system with a unique integer (it will not matter which vertex gets which label!). Given a tetrahedron with vertices \(i_1, i_2, i_3, i_4\) we sort these vertices in ascending

\(^{20}\) Robbert Dijkgraaf is a very prominent theoretical physicist and string theorist. His surname is likely to be difficult to properly pronounce for those who are not from the Netherlands because the “g” is a guttural sound that only exists in Dutch. However, those from the south of the Netherlands don’t use the guttural “g” and instead pronounce it as Dike-Hraff, which is probably about the closest most English speakers will get to the right result.

\(^{21}\) The word “Dijkgraaf” refers to an occupation: A Dijkgraaf is the person in charge of making sure that water stays in the ocean and does not flood the cities and the rest of the Netherlands.

\(^{22}\) In chapter 20 we considered also the possibility of \(d_a = -1\) but this is a gauge choice. We are always entitled to choose +1 instead at the cost of possibly losing isotopy invariance.

\(^{23}\) With apologies for using \(G\) and \(\tilde{G}\) in the same equation to mean completely different things!
order so that

\[ [j_1, j_2, j_3, j_4] = \text{sort}[i_1, i_2, i_3, i_4] \quad \text{such that} \quad j_1 < j_2 < j_3 < j_4 \]

we then define

\[
\tilde{G} \left( \begin{array}{c} i_1 \\ i_2 \\ i_3 \\ i_4 \end{array} \right) = \omega(g_{j_2, j_1}, g_{j_3, j_2}, g_{j_4, j_3})^{s(j_1, j_2, j_3, j_4)} \tag{21.11}
\]

Here \( g_{k,l} \) is the group element on the edge directed from vertex \( k \) to vertex \( l \), and \( \omega \) is the chosen 3-cocycle. The exponent \( s(j_1, j_2, j_3, j_4) \) is either +1 or −1 depending on whether the orientation of the tetrahedron defined by the ordered set of vertices \([j_1, j_2, j_3, j_4]\) has the same or opposite orientation as the manifold we are decomposing\(^{24}\). This prescription gives a manifold invariant (the Dijkgraaf-Witten invariant) for any choice of 3-cocycle even if the corresponding diagrammatic algebra does not have isotopy invariance. Analogous to the Turaev-Viro invariant, the partition function here is gauge invariant if we transform the cocycles with Eq. 20.5 (See exercise 21.4).

### 21.4.1 Other Dimensions

An interesting feature of Dijkgraaf-Witten theory is that essentially the same recipe builds a Dijkgraaf-Witten TQFT in any number of dimensions. One discretizes the \( D \)-dimensional manifold into \( D \)-dimensional simplices (segments in 1D, triangles in 2D, tetrahedra in 3D, pentachora in 4D) and labels each edge with a group element \( g \in G \) and each vertex is assigned an integer label. The flatness condition is always the same as that shown in Fig. 21.6 — multiplying the group elements around a closed loop must give the identity. In \( D \)-dimensions we build the partition function by multiplying a weight for each \( D \)-simplex, where the weight is given now by a so-called \( D \)-cocycle\(^{25}\), which we call \( \omega_D(g_1, g_2, \ldots, g_D) \) which is now a function of \( D \) arguments. Finally, one builds a partition function by summing over all possible labelings

\[
Z_{DW}(M_D) = |G|^{-N_0} \sum_{\text{labelings}} \prod_{\text{D-simplices}} \omega_D(g_{j_2, j_1}, \ldots, g_{j_{D+1}, j_D})^{s(j_1, \ldots, j_{D+1})} \tag{21.12}
\]

As with the 3D case, the arguments of the cocycle \( g_{k,l} \) are the group elements along the edges of the simplex from vertex \( k \) to vertex \( l \) and we always write them ordered such that \( j_1 < j_2 < \ldots < j_D \). Finally the exponent \( s \) is always ±1 depending on whether the orientation of simplex described by the ordered set \([j_1, \ldots, j_{D+1}]\) matches that of the underlying manifold or not.

As a quick example, let us consider the 2D case. The definition of a 2-cocycle \( \omega_2 \) is any function that satisfies the condition\(^{26}\)

---

\(^{24}\)To find the orientation of a tetrahedron, place \( j_1 \) closest to you and see if the triangle \([j_2, j_3, j_4]\) is oriented clockwise or counterclockwise.

\(^{25}\)I won’t give the most general definition of cocycle as this takes us too far afield into group cohomology. However, as with the 3-cocycle it is simply a function satisfying a particular cocycle condition. See Eq. 20.1 for the 3D case and Eq. 21.14 for the 2D case.

\(^{26}\)The 2-cocycle condition is equivalent to the consistency condition for a so-called “projective representation” of the group. For projective representations we have the multiplication rule \( \rho(g)\rho(h) = \omega_2(g, h)\rho(gh) \) whereas for regular group representations we have \( \omega_2 = 1 \). See section 41.2.4.
\[ \omega_2(g, h)\omega_2(gh, k) = \omega_2(h, k)\omega_2(g, hk) \]  

(21.13)

In the partition function, Eq. 21.12, each triangle gets a weight given by the cocycle. It is then easy to see that the cocycle condition is precisely the condition necessary to make the partition function invariant under the 2-2 Pachner move, as shown in Fig. 21.7. It is not a hard exercise to demonstrate invariance under the 3-1 Pachner move as well (See exercise 21.3).

![Fig. 21.7](image)

**Fig. 21.7** Each triangle satisfies the flatness condition Eq. 21.6 meaning multiplying all three edges in order gives the identity. In the partition function each triangle gets a weight given by the corresponding cocycle \( \omega_2 \) as written in black text. All of the triangles in the figure are oriented positively \( s = +1 \). The cocycle condition Eq. 21.13 guarantees that the product of the cocycles on the left equals the product of the cocycle on the right.

### 21.4.2 Further Comments

One particularly interesting special case of Dijkgraaf-Witten theory is the case of the trivial 3-cocycle where \( \omega \) is always unity. In this case, the argument of the sum in Eq. 21.10 (or more generally Eq. 21.12) is just unity so the partition function just counts the number of flat field configurations (See Fig. 21.6) and then divides by \( |G|^N_v \). This partition function is exactly that of lattice gauge theory, as we will see in chapter *** below, and the resulting topological quantum field theory is known as the quantum double of the group \( G \). The more general case, with a nontrivial cocycle is correspondingly sometimes known as “twisted” gauge theory, where the cocycle is thought of as some sort of twist to the otherwise simple theory.

A further interesting relationship is that Dijkgraaf-Witten theory can be thought of as result of symmetry breaking an appropriately chosen Chern-Simons theory (See for example Dijkgraaf and Witten [1990], de Wild Propitius [1993]). One might imagine, for example, breaking a compact \( U(1) \) Chern-Simons gauge theory into a discrete \( \mathbb{Z}_n \) group — like breaking the symmetry of a circle into an \( n \)-sided regular polygon. The particular cocycle one gets in the resulting Dijkgraaf-Witten theory depends on the choice of the coefficient (the “level”) of the Chern-Simons term.

Dijkgraaf-Witten theory has had extensive recent applications within quantum condensed matter physics where it turns out that a classification of so-called symmetry protected topological (SPT) phases is given in terms of Dijkgraaf-Witten theories. We will briefly discuss SPT phases...
Further Reading

- The Turaev-Viro invariant was introduced in [Turaev and Viro 1992]. Rather interestingly Turaev and Viro were apparently unaware of the earlier work by Penrose, Ponzano, Regge and others when they first discussed these state sums! The work was extended to include all spherical fusion categories by [Barrett and Westbury 1996]. A recent rather complete discussion, including looking at the possible world-lines and boundaries is given in the book [Turaev and Virelizier 2017]. Unfortunately, these references and many other works in the field are written in rather mathematical language that is not particularly transparent for most physicists!

- It is worth commenting that the state-sum approach to quantum gravity has been extended in a multitude of ways, and continues to be an active area of research. Among the key directions are extension to 3+1 dimensions (Ooguri [1992] and Crane and Yetter [1993] for example), and extensions to Lorentzian signature (Barrett and Crane [2000]). A nice general discussion of discrete approaches gravity is given by [Regge and Williams 2000].

- One very popular extension of the spin-network modes, known as a spin-foam, is to discretize space but allow the discretization to change as a function of time. A nice review of this direction is given by [Lorentz 2006].

Exercises

Exercise 21.1 Some More Facts about Turaev-Viro

Consider a manifold $\mathcal{M}$ with boundary $\Sigma$ which has been discretized into tetrahedra on in the bulk and triangles on the surface. Let the edges on the surface be labeled by $j_1, \ldots, j_N$. Assume tha the theory has reflection symmetry as in Eq. [16.14] show that

\[ Z_{TV}(\mathcal{M}; \bar{a}_1, \ldots, \bar{a}_N) = Z_{TV}(\mathcal{M}; a_1, \ldots, a_N) \]

And as a result show that for a closed manifold $Z(\mathcal{M})$ is real.

Exercise 21.2 Details of Turaev-Viro

Work carefully through the details of the proof that the Turaev-Viro partition function is invariant under Pachner moves.

Exercise 21.3 2D Dijkgraaf-Witten

The invariance of the two dimensional Dijkgraaf-Witten partition function under the 2-2 Pachner move is established in section [21.4.1]. Show that the partition function is also invariant under the 3-1 Pachner move.

Exercise 21.4 Gauge Invariance
(a) Show that $Z_{TV}$ is unchanged if we make a gauge transformation of the type of Eq. 16.23 (Hint: each plaquette appears in two $F$-matrices in the product Eq. 21.2).

(b) Show that $Z_{DW}$ is unchanged if we make a gauge transformation of the type of Eq. 20.5.

**Exercise 21.5 Lattice Gauge Theory on $S^3$**

The manifold $S^3$ has two very simple one-tetrahedron triangulations (See Jaco and Rubinstein [2003]). These are shown in Fig. 21.8. First convince yourself that these triangulations describe closed manifolds (i.e., check that each face occurs twice in opposite orientations).

Dijkgraaf-Witten theory with a trivial cocycle ($\omega = 1$) is equivalent to lattice gauge theory. By performing the sum in Eq. 21.10 calculate $Z_{DW}(S^3)$ in the case of the trivial cocycle for both triangulations of $S^3$ and check that they give the same result.

**Exercise 21.6 Turaev-Viro on $S^3$**

(a) Consider a Turaev-Viro model built from any $F$-matrices with the property that there is no particle $a$, except the identity, for which $a \times a \times a$ gives the identity. Using a simple triangulation of $S^3$, calculate $Z_{TV}(S^3)$. (Hint: this is much easier than it seems if you use the single vertex triangulation from Fig. 21.8). Does this result agree with the statements in section **.

(b) [Harder] Repeat this calculation without the special condition on $a \times a \times a$.

**Exercise 21.7 Dijkgraaf-Witten for $Z_2$ cocycle**

Let us calculate $Z_{DW}(M)$ for a $Z_2$ theory on the three-torus manifold $M = S^1 \times S^1 \times S^1$. We can represent the 3-torus manifold by identifying opposite faces of the cube shown in Fig. 21.9. Once faces are identified there is only a single vertex in the figure and only 3 edges. One can make a triangulation of this cube with 6 tetrahedra given by 1278, 2378, 3478, 4578, 5678, 6178 (this introduces four new diagonal edges which can be written as 18 = 74 and 38 = 76 and 58 = 72 and 78).

With this triangulation, sum over all possible assignments of group elements to the edges satisfying the flatness condition (Fig. 21.6) and explicitly determine the partition function $Z_{DW}$ when the group is $G = Z_2$. There are two possible 3-cocyles we can choose for this group (See section 20.1.2). Show that $Z_{DW}$ is the same for both of them.

Note that there are much easier methods for evaluating partition functions of this type which do not involve actually doing the whole sum. (See the original work by Dijkgraaf and Witten [1990].)

---

Fig. 21.8 The manifold $S^3$ has two very simple triangulations each involving only a single tetrahedron. On the left two vertices (1 and 2) are connected together with three different edges (single, double, and triple arrow). On the right a single vertex (1) is connected to itself with two different edges (single and double arrow).

Fig. 21.9 To represent a 3-torus, all 8 points are identified. There are 3 distinct edges (for example, the edges 28 and 16 and 34 and 75 are all identified).
Formal Construction of TQFTs from Diagrams: Surgery and More Complicated 3-Manifolds

Having constructed diagrammatic algebras in 2+1 dimension, we have almost all we need to define a TQFT based on these diagrams. As discussed in section 14.3.2 our diagrammatic algebra gave us a way to evaluate a partition function $Z(\text{labeled link in } S^3)/Z(S^3)$, or equivalently $Z(\text{labeled link in } S^2 \times S^1)$ with the caveat that no link goes around the cycle of $S^1$. However, a TQFT should be able to evaluate a partition function in any arbitrary manifold $M$. Indeed, in the simplest case we might dispense with the labeled link and want to find a partition function of the manifold $M$ alone.

In this chapter we develop a prescription for handling more complicated manifolds. One important thing this will achieve will be to give a formal definition to Chern-Simons theory, which we like to think of as being defined as some sort of functional integral, but as pointed out in section 5.3.4 is not really well defined in that language as such integrals do not actually converge.

The way we will handle more complicated manifolds is by sewing pieces of manifolds together with a procedure known as surgery.

### 22.1 Surgery

In chapter 7 we saw two examples of assembling manifolds by gluing together pieces. We found that we could assemble together two solid tori ($D^2 \times S^1$) into either $S^3$ or $S^2 \times S^1$ depending on how we glue together the $S^1 \times S^1$ surfaces. (In fact, one can consider gluing together the surfaces in yet other ways to get even more interesting results, but we will not need that for the moment). We would like to use this sort of trick to study much more complicated three dimensional manifolds.

The understanding of three dimensional manifolds is a very rich and beautiful problem. In order to describe complicated manifolds it is useful to think in terms of so-called surgery. Similar to what we were

---

1. Although this chapter is super interesting and fun, physicists can probably skip it on a first reading.
2. I.e., including braiding again, which we did not need to define the TQFTs in chapter 24.
3. See for example the discussion in section 7.4 as well as Rolfsen [1976] for example.

---

4. Many important results on three dimensional manifolds have been discovered recently. Perelman’s proof of the Poincaré Conjecture, along with the methods he used are apparently extremely revolutionary and powerful. But this is way outside the scope of our book!
5. Grigori Perelman is a brilliant, but startlingly puzzling character. He famously declined the million dollar Millenium Prize offered to him for proving the Poincaré conjecture in three dimensions. He turned down the Fields Medal as well.
Formal Construction of “Chern-Simons” TQFT: Surgery and More Complicated Manifolds

$S^2 = [S^2 - (D^2 \cup D^2)] \cup_{S^1 \cup S^1} (D^2 \cup D^2)$

Fig. 22.1 Writing a sphere $\mathcal{M} = S^2$ as the union of two manifolds glued along their boundaries. $\mathcal{M}_2$ is the union of two disks $D^2 \cup D^2$. $\mathcal{M}_1 = S^2 - (D^2 \cup D^2)$ is the remainder. The two manifolds are glued along their common boundary $S^1 \cup S^1$.

6 Prehensile toes could be useful I suppose!
7 We will only be concerned with orientable manifolds

discussing in section 7.3 — assembling a manifold by gluing pieces together — the idea of surgery is that we remove a part of a manifold and we glue back in something different. Imagine replacing someone’s foot with a hand!6 By using successive surgeries we will be able to construct any three-dimensional manifold7.

The general scheme of surgery is to first write a manifold as the union of two manifolds-with-boundary sewed along their common boundaries. If we have a closed manifold $\mathcal{M}$ that we would like to alter, we first split it into two pieces $\mathcal{M}_1$ and $\mathcal{M}_2$ such that they are sewed together along their common boundary $\partial \mathcal{M}_1 = \partial \mathcal{M}_2^*$. So we have

$\mathcal{M} = \mathcal{M}_1 \cup_{\partial \mathcal{M}_1} \mathcal{M}_2$

We then find another manifold with boundary $\mathcal{M}_2'$ whose boundary matches $\mathcal{M}_2$, i.e.,

$\partial \mathcal{M}_2 = \partial \mathcal{M}_2'$

We can then replace $\mathcal{M}_2$ with $\mathcal{M}_2'$, to construct a new closed manifold $\mathcal{M}'$ as

$\mathcal{M}' = \mathcal{M}_1 \cup_{\partial \mathcal{M}_1} \mathcal{M}_2'$

We say that we have performed surgery on $\mathcal{M}$ to obtain $\mathcal{M}'$. In other words, we have simply thrown out the $\mathcal{M}_2$ part of the manifold and replaced it with $\mathcal{M}_2'$.

22.1.1 Simple Example of Surgery on a 2-manifold

To give an example of surgery consider the sphere $\mathcal{M} = S^2$ as shown in Fig. 22.1. Here we write the sphere as the union of two disks $\mathcal{M}_2 = D^2 \cup D^2$ and the remainder of the sphere $\mathcal{M}_1 = S^2 - (D^2 \cup D^2)$. These are glued along their common boundary $S^1 \cup S^1$.

Now we ask the question of what other 2-manifolds have the same boundary $S^1 \cup S^1$. There is a very obvious one, the cylinder surface! Let us choose the cylinder surface $\mathcal{M}_2' = S^1 \times I$ where $I$ is the interval (or $D^1$). It also has boundary $\partial \mathcal{M}_2' = S^1 \cup S^1$ as shown in Fig. 22.2.

$$
\begin{align*}
S^2 &= \left[ S^2 - (D^2 \cup D^2) \right] \cup_{S^1 \cup S^1} (D^2 \cup D^2)
\end{align*}
$$
Fig. 22.2 The boundaries of the cylinder surface is the same as the boundary of the two disks. Both boundaries are two circles. This means that we can remove two disks from a manifold and sew in the cylinder.

Thus we can sew the cylinder surface in place where we removed $M_2 = D^2 \cup D^2$, as shown in Fig. 22.3. The resulting manifold $M'$ is the torus $T^2$.

Fig. 22.3 Gluing the cylinder surface $M_2' = S^1 \times I$ to the manifold $M_1 = S^2 - (D^2 \cup D^2)$ along their common boundary $S^1 \cup S^1$ gives the torus $T^2$. Note that the object on the right is topologically a torus.

Thus we have surgered a sphere and turned it into a torus. Note that there is another way to think of this procedure. If $M = \partial N$ then surgery on $M$ is the same as attaching a handle to $N$. In the case we just considered we would take $N = B^3$ the 3-ball (sometimes denoted $D^3$), and we attach a handle $D^2 \times I$, the solid cylinder. We obtain the new manifold $N'$ which is the solid torus, whose boundary is $T^2$ the torus surface. This is written out in the diagram Fig. 22.4.

Fig. 22.4 Handle attaching on the manifold $N$ is the same as surgery on a manifold $M = \partial N$. 

\[ \partial(S^1 \times I) = \partial(D^2 \cup D^2) = S^1 \cup S^1 \]

$\partial(\bigcup \cup \circ) = \circ \cup \circ$
22.1.2 Surgery on 3-manifolds

We can also perform surgery on three-dimensional manifolds\(^8\). Start with a simple closed 3-manifold \(\mathcal{M}\), such as \(S^3\) (or, even simpler to think about, consider \(\mathcal{M} = \mathbb{R}^3\) and let us not worry about the point at infinity). Now consider a solid torus

\[
\mathcal{M}_2 = D^2 \times S^1
\]

embedded in this manifold. The surface \(\partial \mathcal{M}_2 = S^1 \times S^1 = T^2\) is a torus surface. Now, there is another solid torus with exactly the same surface:

\[
\mathcal{M}'_2 = S^1 \times D^2
\]

These two solid tori differ in that they have opposite circles filled in. Both have the same \(S^1 \times S^1\) surface, but \(\mathcal{M}_2\) has the first \(S_1\) filled in whereas \(\mathcal{M}'_2\) has the second \(S_1\) filled in.

The idea of surgery is to remove \(\mathcal{M}_2\) and replace it with \(\mathcal{M}'_2\) to generate a new manifold \(\mathcal{M}'\) with no boundary\(^9\). The reason this is difficult to visualize is because if we start with a very simple space like \(\mathcal{M} = \mathbb{R}^3\) the new structure \(\mathcal{M}'\) is not embeddable within the original manifold \(\mathcal{M}\).

This procedure, torus surgery on a 3-manifold, is called Dehn surgery. Another way to describe what we have done is that we have removed a solid torus, switched the meridian and longitude (switched the filled-contractible and the unfilled-uncontractible) and then glued it back in. In fact, one can make more complicated transformations on the torus before gluing it back in (and it is still called Dehn surgery, see section \(7.4\)) but we will not need this.

It is worth noting that the solid torus we removed could be embedded in a very complicated way within the original manifold — i.e., it could follow a complicated, even knotted, path, as in the figure on the right of Fig. 7.10. As long as we have a closed loop \(S^1\) (possibly following a complicated path) and it is thickened to \(D^2\) in the direction transverse to the \(S^1\) path, it is still a solid-torus topologically.

22.2 Representing Manifolds with Knots

22.2.1 Lickorish-Wallace Theorem

An important theorem\(^{10}\) of topology is due to Lickorish [1962] and Wallace [1960].

**Theorem:** Starting with \(S^3\) one can obtain any closed connected orientable 3-manifold by performing successive torus surgeries, where these tori may be nontrivially embedded in the manifold (i.e., they may follow some knotted path).

One has the following procedure. We start with a link (some knot

\[^8\]This is the part that is guaranteed to make your head explode.

\[^9\]Stop here, think about what we have done. Collect the pieces of your exploded head.

\[^{10}\]In Witten’s groundbreaking paper on the Jones polynomial (Witten [1989]), he states the theorem without citation and just says “It is a not too deep result...”. Ha! Seriously though, the proof is actually not too difficult. Although some of the details are a bit tricky the main idea is fairly understandable. See the references at the end of the chapter.
possibly of several strands), embedded in $S^3$. Thicken each strand to a solid torus. Excise each of these solid tori, and replace them by tori with longitude and meridian switched\footnote{See also section 7.4.}. Any possible 3-manifold can be obtained in this way by surgering an appropriately chosen link. We summarize with the mapping

$$\text{Link in } S^3 \xrightarrow{\text{surger}} \text{Some } M^3$$

We can thus represent any three dimensional manifold as a link in $S^3$. If we think of a topological quantum field theory as being a way to assign a complex number to a three dimensional manifold, i.e., $Z(M)$ we realize that what we are now looking for is essentially a knot invariant — a way to assign a number to a knot. We exploit this connection further when we discuss the Witten-Reshitikhin-Turaev invariant below in section 22.3.

### 22.2.2 Kirby Calculus

It turns out that not all topologically different links, when surgered, give topologically different manifolds. Fortunately, the rules for which knots give the same manifolds have been worked out by Kirby \cite{Kirby1978}. These rules, known as Kirby calculus, are stated as a set of transformation moves on a link which change the link, but leave the resulting manifold unchanged. There are several different sets of moves that can be taken as “elementary” moves which can be combined together to make more complicated transformations. Perhaps the simplest set of two elementary basic moves are known as Kirby moves which we will present here\footnote{If one does not start with the knot embedded in $S^3$, one may need a third move known as “circumcision”. This says that if any string loops only once around another string (without twisting around itself and without looping around anything else), both strings may be removed. I.e., in Fig. 22.5 both strings may be removed (independent of how the string going off to the left forms any knot).}.

We will not rigorously prove that these moves leave the manifold unchanged, but we will give rough arguments instead.

**Kirby Move 1: Blow Up/ Blow Down:**

One can add or remove a loop with a single curl, as shown in Fig. 22.6, to a link and the manifold resulting after surgery remains unchanged.

**Argument:** First let us be a bit more precise about the surgery prescription. Given a link, we think of this link as being a ribbon (usually...
we draw it with blackboard framing, see section 2.2.2. Thicken each strand into a solid torus, and draw a line around the surface of this torus that follows one of the edges of the ribbon. Remove this solid torus, but the torus surface that remains still has the line drawn around it. Reattach a new solid torus where the new meridian (the circle surrounding the contractible direction) follows precisely this line.

Now consider a curled loop as in Fig. 22.6 embedded in $S^3$. As shown in Fig. 2.7 a string with a small curl as in Fig. 22.6 can be thought of as a ribbon with a twist (but no curl) in it. Let us use this description instead. Thicken the loop to a torus, and then the edge of the ribbon traces out a line as shown in Fig. 22.7 on the torus surface. We remove the solid torus and insert a new torus where the meridian follows the twisted line on the surface of the hole that is left behind. This is exactly the construction of $L(1,1) = S^3$ described in 7.4 above (it is $(1,1)$ since the blue line goes around each cycle once), thus showing an example of how surgery on the curled loops in Fig. 22.6 does nothing to the manifold.

Kirby Move 2: Handle-Slide:\footnote{The nomenclature “handle slide” comes from an interpretation of this move as sliding handles around on a manifold. Consider the example used in section 22.1 where we attached a handle to a ball and obtained a solid torus. We could also attach two handles and get a two-handled solid torus. Here it doesn’t matter where the handles are attached to the sphere – they can be slid around. Indeed, they can even be slid over each other (where one handle attaches to some point on the other handle). It is the sliding of a handle over another handle which gives this move its name.}

A string can be broken open and pulled along the full path of another string, and then reconnected, and the resulting manifold remains unchanged. See Fig. 22.8 or 22.9.

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{handle-slide.png}
\caption{A handle-slide move. (See Fig. 22.9 for another example.) Both left and right sides of this picture represent the same 3-manifold after surgery. Note that we should always view both strings as ribbons, and we need to keep track of how many self-twists the ribbon accumulates when it is slid over another string.}
\end{figure}

**Argument:** Consider the simple handle-slide shown in Fig. 22.9. Let us think about what happens when we surge the horizontal loop. First we thicken the horizontal loop into a torus (as shown), then we exchange the contractible and non-contractible directions. In this procedure, the longitudinal direction (The long direction) of the torus is made into something contractible. This means (after surgery) we can pull the far left vertical line through this torus without touching the three vertical blue lines. Thus the right and left pictures must describe the same manifold. While it is a bit harder to argue generally, this principle remains true even if the torus is embedded in the manifold in a complicated way,
Fig. 22.9 An example of a simple handle-slide move.

as in Fig. 22.8

Two links in $S^3$ describe the same 3-manifold if and only if one link can be turned into the other by a sequence of these Kirby moves as well as any smooth deformation of links (i.e., regular isotopy).

Note that if we have two disconnected links $L_1$ and $L_2$ which surgered give two manifolds $M_1$ and $M_2$ respectively,

$$L_1 \xrightarrow{\text{surger}} M_1$$
$$L_2 \xrightarrow{\text{surger}} M_1$$

then if we consider a link $L_1 \cup L_2$ which is the disconnected union of the two links (i.e., the two links totally separated from each other) it is fairly easy to see that we obtain the so-called connected sum of the two manifolds which we write as follows:

$$L_1 \cup L_2 \xrightarrow{\text{surger}} M_1 \# M_2$$

See the discussion of connected sums in section 7.3.3.

22.3 Witten-Reshitikhin-Turaev Invariant

By using the ideas of surgery, we are now in a position to use our diagrammatic algebra to handle complicated manifolds. Recall that one of the definitions of a TQFT is a mapping from a manifold $M$ to a complex number $Z(M)$ in a way that depends only on the topology of the manifold (for example, Eq. 5.17 or Fig. 7.1 but without the embedded link). By using surgery (Eq. 22.1) we can describe our manifolds as links in $S^3$. If we can then find a link invariant that is unchanged under Kirby moves, we will effectively have something we can use as a manifold invariant. Thus we are now looking to construct a link invariant, and our diagrammatic algebra will provide exactly what we need!

We want to have a link invariant which is fully isotopy invariant (since Kirby calculus is isotopy invariant). In the simplest case let us assume we have no negative Frobenius-Schur indicators$^{15}$ so we can take all $d_a = d_a$ positive and we have a fully isotopy invariant diagram algebra.

$^{15}$In the more general case where we have negative Frobenius-Schur indicators, we can use the scheme of section 14.3 to insure isotopy invariance. It is also possible to work work with negative $d_a$'s as discussed in section 14.3 and not apply rule 0, i.e., performing a non-unitary evaluation of the diagram. In many discussions of Reshitikhin-Turaev invariant, this is essentially what is done. Note that in this case the Kirby color Fig. 17.7 is then defined with $d_a/D$ rather than $d_a/D$. The identities of Fig. 17.8 and 17.11 hold for this non-unitary evaluation.
The key to this construction is to consider a link of the \( \Omega \) (Kirby) strands discussed above in section 17.5. This link made of \( \Omega \) represents the link to be surgered, and thus represents our manifold. Let us now consider a manifold invariant defined as

\[
Z_{WRT}(\mathcal{M}) = \frac{1}{D} \left[ e^{2\pi i c/8} \right]^\sigma \left( \text{Evaluate link made of } \Omega \text{-strands where surgery on link in } S^3 \text{ gives } \mathcal{M} \right)
\]

(22.2)

where \( \sigma \) is the so-called signature of the link, defined to be the number of positive eigenvalues minus the number of negative eigenvalues of the matrix of linking numbers \( \text{link}_{ij} \) between the (possibly multiple) strands of the link (The diagonal element \( \text{link}_{ii} \) is just the self-linking or writhe of strand \( i \). See section 2.6.2 for definition of linking number\(^{16} \)).

It is not so obvious that the definition in Eq. 22.2 should provide a manifold invariant. What we would need to show is that \( Z_{WRT}(\mathcal{M}) \) gives the same output for any link that describes the same \( \mathcal{M} \). In other words we have to show that the expression on the right hand side of Eq. 22.2 is unchanged when we make Kirby moves on the link.

Let us consider the first Kirby move, the addition of a twisted loop as in Fig. 22.6. Using Fig. 17.11, adding such a twisted loop multiplies the value of the link (the final term in Eq. 22.2) by \( e^{\pm 2\pi i c/8} \) (\( \pm \) depending on which way the loop is twisted). However, the addition of the twisted loop also changes the signature of the link \( \sigma \) by \( \pm 1 \) thus precisely canceling this factor. Thus the expression in Eq. 22.2 is certainly unchanged under the first Kirby move\(^{17} \), the Blow-Up/Blow-Down.

The manifold invariant Eq. 22.2 is known as the Witten-Reshetikhin-Turaev invariant, and was invented by Reshetikhin and Turaev [1991]. The reason it also gets named after Witten is that it gives a rigorous re-definition of the Chern-Simons manifold invariants (Eq. 5.17) discussed by Witten [1989]. This is a rather important result being that the Chern-Simons functional integral is not well defined as an integral! (See the comments in section 5.3.4)\(^{18} \).

\(^{16}\)Note that to calculate a linking matrix, we must orient all of the strands (i.e., put arrows on them). It does not matter which way these arrows point.

\(^{17}\)The killing property of Fig. 17.8 also makes Eq. 22.2 invariant under the third Kirby move, Fig. 22.5.

\(^{18}\)We mention in section 5.3.4 that the Chern-Simons partition function, among other reasons for being ill-defined, actually depends on a so-called 2-framing of the manifold. The Reshetikhin-Turaev invariant corresponds to choosing so-called canonical framing. This is discussed in depth by Atiyah [1990a]; Kirby and Melvin [1999].

for links (as discussed in section 14.3.2).
Fig. 22.10 Proof that the $\Omega$ strand satisfies the handle-slide. Here we show that any strand $a$ can freely slide over the $\Omega$ strand in the sense of Fig. 22.8. The $\Omega$ strand on the right is meant to be connected up to itself in some way in a big (potentially knotted or linked) loop which we don’t draw. In going from the first to the second line, and also in going from the second to the third line we have used the completeness relation Eq. 16.26. The equality in the second line is just sliding the vertex from the top (where $a$ and $b$ split from $c$) all the way around the $b$ strand on the right until it almost reaches the bottom $a, b, c$ vertex. Note this derivation does not even require a well defined braiding.

Note the multiplication law for connected sums of manifolds

$$Z_{WRT}(M_1 \# M_2) = D Z_{WRT}(M_1) Z_{WRT}(M_2) \quad (22.3)$$

This multiplication law is from the fact that surgery on a disjoint union of links gives a connect sum of manifolds (Eq. 22.2) and the evaluation of the disjoint union of links gives the product of the individual evaluation of the two links\(^{19}\).

Further one can extend these manifold invariants to give a topological invariant partition function of a labeled link within a manifold as in Fig. 7.1 (this was one of our general definitions of what we expected

\(^{19}\)Some references redefine $Z_{WRT}$ without the factor of $D$ out front such that $Z(M_1 \# M_2) = Z(M_1) Z(M_2)$ instead.
from a TQFT). To make this extension we simply define

\[ Z_{WRT}(\mathcal{M}; \text{labeled link}) = \frac{1}{D} \left[ e^{i2\pi \varphi/8} \right]^{\sigma} \begin{cases} \text{Evaluate link made of} \\ \Omega\text{-strands where} \\ \text{surgery on link in } S^3 \text{ gives } \mathcal{M} \\ \cup \text{labeled link} \end{cases} \]

In other words, we simply include the labeled link into the diagram to be evaluated.

Without ever saying the words “path integral” or “Chern-Simons action” we think of an anyon theory as simply a way to turn a link of labeled world lines into a number (like evaluating a knot invariant, but with rules for labeled links), and surgery on \( \Omega \) strands allows us to represent complicated manifolds.

### 22.3.1 Some examples

It is worth working through a few examples of calculating the Witten-Reshitikhin-Turaev Invariant for a few simple manifolds.

For \( \mathcal{M} = S^3 \), we don’t need to surger the manifold at all, so we don’t need any \( \Omega \) link at all. The value of the (empty) link is normalized to unity and including the prefactor in Eq. 22.2 (with signature zero) we obtain

\[ Z_{WRT}(S^3) = \frac{1}{D} \]

which matches our expectations given Eqs. 17.12 and 7.7. Note that given this relationship (See Eq. 17.6) Eq. 22.3 formula agrees with Eq. 7.15.

For \( \mathcal{M} = S^2 \times S^1 \) we need to surger a single loop in \( S^3 \) to obtain \( S^2 \times S^1 \) (See exercise 22.1). Thus we need to evaluate a single loop of \( \Omega \) string. It is an easy calculation to evaluate a loop of \( \Omega \):

\[ \ = \sum_a \frac{d_a}{D} \quad \quad \quad \ = \sum_a \frac{d_a^2}{D} = D \quad \quad (22.4) \]

Thus including the prefactor in Eq. 22.2 (the signature of the link is zero) we obtain

\[ Z_{WRT}(S^2 \times S^1) = 1 \]

which is in agreement with Eq. 7.8.

Finally let us consider the three-torus manifold \( \mathcal{M} = S^1 \times S^1 \times S^1 = T^2 \times S^1 = T^3 \). First, we note that surgery on the Borromean rings\(^{20}\) (Fig. 22.11) yields the three torus (See exercise 22.3). To evaluate the link we use the corollary of the killing property of the \( \Omega \) strand, Fig. 17.9

---

\(^{20}\)The rings are named for the crest of the royal Borromeo family of Italy, who rose to fame in the fourteenth century. However the knot (in the form of three linking triangles) was popular among Scandinavian runestones five hundred years earlier and was known as the “Walknot” or “Valknut”, or “the knot of the slain.”
to show (See exercise 22.4)
\[ Z_{WRT}(T^3) = \text{number of particle species} \quad (22.5) \]
which matches the prediction from Eq. 7.3 along with Eq. 7.5.

22.3.2 Turaev-Viro Revisited: Chain-Mail and the Turaev-Walker-Roberts Theorem

Using the ideas of surgery [Roberts 1995] produced a beautiful geometric proof of the Turaev-Walker theorem ([Turaev 1992, 1994; Walker 1991]) which relates the Turaev-Viro invariant to the Chern-Simons (Witten-Reshitikhin-Turaev) invariant of a manifold. The result is, given a modular tensor category (a modular anyon theory) we have

\[ Z_{TV}(M) = |Z_{WRT}(M)|^2 \quad (22.6) \]

We will not give the full proof here, only the general idea.

First we will require one more minor corollary. Similar to Fig. 17.9 we have the identity shown in Fig. 22.12 (See exercise 22.6).

\[ \Omega = \frac{D}{\Theta(a,b,c)} \]

Fig. 22.12 The \( \Omega \) strand fuses three lines due to the killing property. Here we have assumed an isotopy invariant theory as discussed in chapter 16, so we can draw vertices with all three lines pointing in the same direction so \( a, b, c \) fuse together to the identity. Here \( \Theta(a,b,c) = \sqrt{d_a d_b d_c} \) as in Eq. 21.3.

We now want to construct a link of \( \Omega \) strands which evaluates to the same value as the Turaev-Viro invariant discussed in chapter 21. Recall that to define the Turaev-Viro invariant, we first make a simplicial decomposition of the manifold, breaking it up into tetrahedra, we label each edge, and we sum a certain weight over all possible labelings as given in Eq. 21.1.

Given our simplicial decomposition here we will instead construct a link of \( \Omega \) strands via the following procedure: Put one loop of \( \Omega \) following the edges of each triangular face (colored gold in Fig. 22.13), and one loop of \( \Omega \) around the waist of each edge (colored purple in Fig. 22.13) in such a way that the two types of strands link with each other. Such a link is known as chain-mail\(^{21}\). We then define the so-called chain-mail invariant of the manifold \( M \) as

\[ CH(M) = D^{-N_v - N_{tet}} \left( \text{Evaluate Chain-Mail Link of } \Omega \text{ strands} \right) \quad (22.7) \]

where \( N_v \) is the number of vertices in the simplicial decomposition and \( N_{tet} \) is the number of tetrahedra\(^{22}\).  

\(^{21}\)When I have given talks on this subject I have been surprised to discover that many people don’t know that chainmail is a medieval type of armor made of linked metal loops. Of course those who had misspent youth playing Dungeons and Dragons, or reading the Lord of the Rings are very familiar with the concept and can tell you why Mithril is the best type of chainmail.

\(^{22}\)More generally the chain-mail link can be defined for any handlebody decomposition of the manifold where \( \Omega \) loops are put around 1-handles and 2-handles and \( N_v \) is then the 0-cells and \( N_{tet} \) is the 3-cells.
First, it is extremely easy to prove that the chain-mail invariant is independent of the particular simplicial decomposition (and hence is a manifold invariant as claimed). We need only show that it is unchanged under the Pachner moves (Fig. 21.4 and 21.5). This can be done entirely geometrically using only the killing property (Fig. 17.9) and the handle-slide property (Fig. 22.10) of the $\Omega$ strand (this is exercise 22.7).

Moreover, it is not hard to show that the chain-mail invariant is actually equal to the Turaev-Viro invariant. To do this we directly evaluate the chain-mail link. We start by using identity 22.12 on each $\Omega$ strand attached to each face (those drawn as gold in Fig. 22.13). This generates a factor of $D/\Theta(a, b, c)$ for each face. The remaining $\Omega$ strands (purple in Fig. 22.13) are decomposed into sums of all quantum numbers as per the definition of $\Omega$ in Fig. 17.7 each weighted by $d_a/D$. This leaves one tetrahedron of particle strings per simplex as shown on the right of Fig. 22.13. (Note that the remaining tetrahedron of strings to be evaluated is a tetrahedral diagram dual to the original tetrahedron, in agreement with the discussion below Eq. 21.4).

Putting together the factors we have obtained leaves us with the chain-mail invariant (including the prefactor in the definition) being given by

$$CH(M) = \frac{D^{N_f}}{D^{N_v+N_e+N_{tet}}} \sum_{\text{edge labels}} \prod_{\text{tetrahedra}} \tilde{G}(\text{tetrahedron}) \prod_{\text{edges}} d_{\text{edge}} \prod_{\text{triangles}} \Theta(\text{triangle})$$

with $N_v, N_e, N_f, N_{tet}$ being the number of vertices, edges, faces (triangles), and tetrahedra respectively. Finally using the well-known topological fact that in three dimensions, the Euler characteristic $N_{tet} - N_f + N_e - N_v$ is zero, the factors of $D$ are reassembled to give exactly the

![Fig. 22.13](image-url)
definition of the Turaev-Viro invariant Eq. 21.1 thus deriving
\[ CH(M) = Z_{TV}(M) \]

Finally we turn to briefly discuss the derivation of the Turaev-Walker theorem Eq. 22.6. The key to this derivation is the fact\(^{23}\) that if one uses a particularly simple decomposition of the manifold, surgery on the chain-mail link generates the connected sum of the original manifold \(M\) and its mirror image \(\overline{M}\)

\[ \text{Chain-mail link for } M \xrightarrow{\text{surger}} M \# \overline{M} \quad (22.8) \]

Evaluating the chain-mail link is therefore essentially equivalent to evaluating \(Z_{WRT}(M \# \overline{M})\)

Using the equivalence between chain-mail and the Turaev-Viro invariant we thus have (Eq. 22.3)

\[ Z_{TV}(M) \sim Z_{WRT}(M \# \overline{M}) \sim Z_{WRT}(M)Z_{WRT}(\overline{M}) \sim |Z_{WRT}(M)|^2 \]

We have written this equation with \(\sim\) rather than an equality because we have dropped factors of \(D\). To get these right we have to know more details about the particular decomposition of the manifold for which Eq. 22.8 holds so that we can keep track of the factors of \(D\) in the definition of the chain-mail invariant (Eq. 22.7). Keeping track of these factors carefully one obtains the desired Eq. 22.6.

Further Reading

Reshetikhin and Turaev [1991]; Lickorish [1993]; Witten [1989].

For more detailed discussion of Surgery, the Lickorish-Wallace theorem, and Kirby Calculus, as well as a nice discussion of manifold invariants see Prasolov and Sossinsky [1996]; Saveliev [2012] or Lickorish [1997].

Roberts/Blanchet refinement.

Mention Crane-Yetter

The following references are standards for Surgery and Kirby Calculus, although they emphasize four dimensional topology. Gompf and Stipsicz [1999]; Kirby [1989]; Akbulul [2016].

Exercises

Exercise 22.1 Surgery on a Loop

Beginning with the three-sphere \(S^3\), consider the so-called “unknot” (a simple unknotted circle \(S^1\) with no twists) embedded in this \(S^3\). Thicken the circle into a solid torus \((S^1 \times D^2)\) which has boundary \(S^1 \times S^1\). Now perform surgery on this torus by excising the solid torus from the manifold \(S^3\) and replacing it with another solid torus that has the longitude and meridian switched. I.e.,

\[ \text{Exercise 22.1 } \text{Surgery on a Loop} \]
replace $S^1 \times D^2$ with $D^2 \times S^1$. Note that both of the two solid tori have the same boundary $S^1 \times S^1$ so that the new torus can be smoothly sewed back in where the old one was removed. What is the new manifold you obtain? (This should be easy because it is in the book!)

**Exercise 22.2 Surgery on the Hopf Link** [Not hard if you think about it right!]

Consider two linked rings, known as the Hopf link (see Fig. 22.14). Consider starting with $S^3$ and embedding the Hopf link within the $S^3$ with “blackboard framing” (i.e., don’t introduce any additional twists when you embed it). Thicken both strands into solid tori and perform surgery on each of the two links exactly as we did above. Argue that the resulting manifold is $S^3$.

**Exercise 22.3 Surgery on the Borromean Rings** [Hard]

Consider the link shown in Fig. 22.11 known as the Borromean rings. Consider starting with $S^3$ and embedding the Borromean rings within the $S^3$ with “blackboard framing”. Thicken all three strands into solid tori and perform surgery on each of the three links exactly as we did in the previous two problems. Show that one gets the three torus as a result. Hint 1: Think about the group of topologically different loops through the manifold starting and ending at the same point, the so-called “fundamental group” or first homotopy group. (See section 41.3). Hint 2: If we say a path around the meridian of one of the three Borromean rings (i.e., threading though the loop) is called $a$ and the path around the meridian of the second ring is called $b$, then notice that the third ring is topologically equivalent to $aba^{-1}b^{-1}$. Hint 3: In some cases the fundamental group completely defines the manifold! (Don’t try to prove this, just accept this as true in this particular case.)

**Exercise 22.4 Evaluation of Borromean Ring $\Omega$-Link**

Use Fig. 17.9 to evaluate the $\Omega$-link of Borromean rings shown in Fig. 22.11. Use this to establish Eq. 22.5. Note that the signature of the link is zero.

**Exercise 22.5 Product of Blow Up and Blow Down**

Use the handle-slide and the killing property of $\Omega$ to prove that the diagram made of two oppositely twisted $\Omega$ loops, as shown in Fig. 22.13, gives the identity.

**Exercise 22.6 Killing Three Strands with $\Omega$**

Prove the relationship shown in Fig. 22.12.

**Exercise 22.7 Pachner Moves and the Chain-Mail Invariant**

Using killing moves (Fig. 17.9) and handle-slides (Fig. 22.10) show that the chain mail invariant Eq. 22.7 is unchanged under Pacher moves (Fig. 21.3 and 21.5). The answer is given by Roberts [1995], but it is a fun exercise. Looking up the answer spoils the fun!
A commonly discussed mechanism that derives one anyon theory from another is known as anyon condensation. The idea is modeled on the notion of conventional Bose-Einstein condensation. Under certain conditions one can imagine anyons forming a superfluid state, akin to a Bose-Einstein condensate. One can imagine making a condensate form either by continuously reducing the temperature with a fixed Hamiltonian, or by continuously changing the Hamiltonian at fixed (perhaps zero) temperature\(^1\). If one begins with a consistent anyon theory before the condensation, the system after the condensation will also be a consistent anyon theory, which we call the condensed theory\(^2\). It is believed that all continuous phase transitions that can occur between different anyon theories can be described in terms of anyon condensation\(^3\).

There is a very detailed theory of anyon condensation, worked out by Bais and Slingerland [2009] and others (see references at end of chapter). Here we will give an abbreviated discussion, along with a few explicit examples.

Let us review some aspects of Bose condensation (see Leggett [2006] or Annette [2004] for much more information about the physics of superfluids and condensates). Recall that in a Bose condensate a macroscopic number of the particles reside in one special lowest-energy single-particle eigenstate which we call the condensate wavefunction. For a uniform system (say with periodic boundary conditions) the wavefunction for this single particle eigenstate is just a constant

\[
\psi(r) = \frac{1}{\sqrt{V}}
\]  

(23.1)

with \(V\) the volume of the system. It is crucial that bosons accumulate no phase or sign when they are braided around each other or exchanged with each other. If they were to accumulate any phase or sign, this would prevent them from remaining in the eigenstate Eq. (23.1) which is everywhere real and positive. This gives us:

**Principle 1:** Bosons must experience no net phase or sign when they move around then comes back to the same configuration — i.e., when bosons exchange or braid with other particles in the condensate.

Indeed, accumulating no sign when exchanging with other identical particles is the very definition of a boson\(^4\).

With interacting bosons, one does not strictly have Bose condensation (not all of the bosons occupy the same single particle eigenstate, since

\(^1\)A phase transition that occurs at zero temperature as some parameter of the Hamiltonian is changed is often known as a “quantum phase transition”.  
\(^2\)It is sometimes possible that the condensed theory is a trivial theory — having only the vacuum particle type, and zero central charge. We should think of that as just being an uninteresting insulator. Strictly speaking this is a TQFT, just a very trivial one. It is also possible to have phase transitions from topological theories to nontopological phases of matter such as gapless phases or fermi liquids.  
\(^3\)First order, or discontinuous, phase transitions can always occur between any two phases of matter.  
\(^4\)It is possible that the condensate wavefunction has a spatial structure such as \(e^{i\phi(r)}\), which happens when there is, say, a vortex within the condensate. What is crucial is that when bosons move around within the condensate, when they get back to the same many-particle configuration the phase is the same as when they started.
Anyon Condensation

interactions kick the particles out of this eigenstate). Nonetheless, interacting bosons can condense to form superfluids which share many of the properties of Bose condensates. In particular, one still has the idea of a condensate wavefunction (or order parameter), and in order to form a condensate, no phase or sign must be accumulated when the particles exchange and braid.

To describe a condensate wavefunction (or order parameter) microscopically, one writes

$$\phi(r) = \langle \hat{\psi}(r) \rangle$$

where $\hat{\psi}(r)$ is the (second quantized) operator which annihilates a particle at position $r$. For noninteracting bosons, where many bosons are in the single particle wavefunction Eq 23.1, we obtain $|\phi|^2 = N_0/V$ where $N_0$ is the number of bosons in the single eigenstate.

The fact that this order parameter is (at least locally) number nonconserving (it destroys a particle) gives us the second important principle

**Principle 2**: Bosons can be freely absorbed by, or emitted from, the condensate.

### 23.1 Condensing Simple Current Bosons

We now would like to generalize the idea of Bose condensation to anyon theories. For simplicity begin by restricting our attention to bosons that are also simple currents and only lift this restriction in section 23.7. To remind the reader, a particle, let us call it $J$, is a simple current if $N_{Ja} = 0$ or 1 for all particle types $a$ and $c$. This condition is equivalent to the statement that $J^N = I$ for some integer $N$ where $I$ is the identity (where $J^N$ here means $N$ factors of $J$ fused together).

For a particle $J$ to condense, it must be a boson. This means that it must have trivial braiding with itself.

$$R_{J\times J}^J = 1$$

or equivalently a trivial spin factor (See Eq. 15.1)

$$\theta_J = 1$$

which is what we expect for a boson. This condition implements the above **Principle 1**: the boson must not experience a nontrivial phase as it exchanges with another particle as this would prevent a condensate wavefunction from forming.

Within the condensate, bosons may fuse with each other to form particles $J^p$ for any value of $p$. It is not hard to show that all such resulting particle types must also be bosons $\theta_{J^p} = 1$ and further, they all must braid trivially with each other (See exercise 23.1).

While one can condense bosons that are not simple currents, the rules for doing so are a bit more complicated and we will return to discuss this more complicated case in section 23.7.
We will start with an initial anyon theory which we call \( \mathcal{A} \). Within this anyon theory, we assume there is a bosonic simple current \( J \) which we intend to condense to form a new anyon theory. The final anyon theory that comes out at the end of the condensation procedure will be called \( \mathcal{U} \).

We can think of anyon condensation as proceeding in two conceptual steps as shown in Fig. 23.1. In between the initial theory \( \mathcal{A} \) and the final theory \( \mathcal{U} \) there is another theory \( \mathcal{T} \) which is not a full anyon theory, but rather a fusion algebra (or planar diagram algebra), as we will discuss further in a moment.

**Fig. 23.1** Condensing one anyon theory to another can be described as having two “steps”. The original anyon theory is labeled \( \mathcal{A} \) and the final anyon theory is labeled \( \mathcal{U} \). In between we have the intermediate theory \( \mathcal{T} \) which is not generally a full-fledged anyon theory, but rather a fusion algebra (planar diagram algebra). The first step from \( \mathcal{A} \) to \( \mathcal{T} \) involves identification and possibly splitting. The second step from \( \mathcal{T} \) to \( \mathcal{U} \) involves confinement.

### 23.2 Identification Step

The first step in the condensation process is the identification step. In this step we group the particle types from the uncondensed theory \( \mathcal{A} \) into so-called orbits.

**Definition:** The orbit of a particle type \( a \) under the action of \( J \) is the set of all particle types \( b \in \mathcal{A} \) such that \( b = J^p \times a \) for some integer \( p \). We denote the orbit as \([a]_J\), or when it not ambiguous we just write \([a]\).

One should be cautioned\(^8\) that the orbit \([a]\) is the same as orbit \([b]\) if \( b = J^q \times a \) for any \( q \).

Further, we note that if \( N \) is the smallest integer such that \( J^N = I \) then there are at most \( N \) particle types in any given orbit, although there may be fewer particles in an orbit, as we will discuss in detail in section 23.3.

The physical point here is that all of the particles types in the same orbit of the original theory \( \mathcal{A} \) are identified as being the same particle type in the \( \mathcal{T} \) theory. The physical reason for this is **Principle 2**: bosons can be freely emitted from or absorbed into the condensate. A particular particle \( a \) can absorb a boson from the condensate and become \( J \times a \) or

\(^8\)This can sometimes cause some notational confusion. It is often useful to choose a single representative of each orbit so that each orbit is uniquely denoted as a particular \([a]\) and one never writes \([b]\) if \( b = J^q \times a \).
it can absorb two bosons from the condensate to become $J^2 \times a$ and so forth. The quantum number $a$ is no longer a conserved quantum number (and therefore is not a valid particle type), but the orbit $[a]$ remains conserved and can play the role of a particle type in the condensed theory. The orbits in the condensed theory will inherit fusion rules from the fusion rules of the uncondensed theory (with some potential complications we will address in section 23.4).

### 23.2.1 Orbits of maximum size

Here we will consider the case where all of the orbits are of maximum size. I.e., all orbits have exactly $N$ particle types in them (where $N$ is the smallest positive integer so that $J^N = I$). We will return to the more complicated case where not all orbits are of maximum size in section 23.4.

We start with the original theory $A$, and each particle $a$ can be mapped to an orbit $[a]$ in the $T$ theory. The fusion rules of the $T$ theory are inherited from the fusion rules of the orginal anyon theory in a natural way which we can write in terms of the fusion multiplicity matrices as

$$N_{[a], [b]} = N_{a,b}$$

Note in particular that the identity particle $I$ of the $A$ theory maps to the orbit $[I]$ which becomes the identity particle of the $T$ theory.

**Example:** $Z_8^{(3+1/2)}$

Let us consider the anyon theory $Z_8^{(3+1/2)}$ discussed in section 20.4.2 which is equivalent to the Chern-Simons theory $SU(8)_1$. There are 8 particles which we label $p = 0, \ldots, 7$ with fusion rules

$$p \times p' = (p + p') \mod 8 \quad (23.2)$$

and $p = 0$ is the identity. The corresponding twist factors are

$$\theta_p = \exp \left[\frac{2\pi i 7}{16} p^2\right] \quad (23.3)$$

We notice that $p = 4$ has trivial twist factor $\theta_4 = 1$ and is therefore a boson. Let us call this bosonic particle $J$, and we notice that $J^2 = I$ so the maximum orbit size is 2.

In this model we have four different orbits under the action of fusing with the boson $J$, and each of these orbits is of maximum size 2. Let us write down these orbits (Recalling that $[a]$ means the orbit of $a$)

- $[0]$ which is also equal to $[4]$
- $[1]$ which is also equal to $[5]$
- $[2]$ which is also equal to $[6]$
- $[3]$ which is also equal to $[7]$
The meaning here should be obvious. Remembering that the boson \( J \), which is particle \( p = 4 \) can be absorbed or emitted for free, we then for example, must identify particles 1 and 5 into a single orbit since fusing 1 with 4 gives 5 and fusing 5 with 4 gives 1.

These four different orbits comprise the particle types of the intermediate \( T \) theory. Let us denote these four orbits as \([p] \) with \( p = 0, \ldots, 3 \). The fusion rules are inherited from the original uncondensed anyon fusion rules (Eq. 23.2) in an obvious way giving

\[
[p] \times [p'] = [(p + p') \mod 4] \tag{23.4}
\]

with \([0]\) playing the role of the identity in the \( T \) theory. To see how these fusion rules come from those of Eq. 23.2 consider, for example, \([1] \times [2] = [3] \): Here either 1 or 5 (the two particle types in the orbit \([1]\)) fused with either 2 or 6 (the two particle types in the orbit \([2]\)) will always give us 3 or 7 (the two particle types in the orbit \([3]\)).

### 23.3 Confinement Step

The particle types in the intermediate theory \( T \) form a consistent fusion algebra (and indeed a consistent planar diagrammatic algebra) but they do not generally form a consistent anyon theory, as they do not generally have a consistent braiding (or solution to the hexagon equations). The reason for this is that some of the particles in \( T \) are not valid particles of the final condensed anyon theory \( U \) and must be thrown out.

The reason some particles of \( T \) are not valid anyons in the condensed phase is that they braid nontrivially with the condensed boson. Trying to put a particle within the condensate that braids nontrivially with the condensed boson would violate **Principle 1**: when a boson in the condensate moves around, the phase must be the same when it arrives back at the same point. We thus have the rule that any particles \( a \) (or its orbit \([a]\)) allowed in the final condensed theory \( U \) must braid trivially with the condensate, meaning that

\[
R_{J \times a}^J R_{J \times a}^{J \times a} = \frac{\theta_{J \times a}}{\theta_{a}} = 1 \tag{23.5}
\]

where we have used Eq. 15.3 and the fact that \( \theta_J = 1 \). Since \( J \times a \) and \( a \) are in the same orbit \([a]\) the condition Eq. 23.5 can be rephrased by saying that an orbit (a particle type of the \( T \) theory) is allowed into the final anyon theory \( U \) if all of the particles in the orbit have the same spin factor \( \theta \). Such particles that are allowed in \( U \) we say are *deconfined*, meaning that they can travel freely within the condensate. The particle types from \( T \) that braid nontrivially with the condensate are not allowed within the condensate and we say they are *confined*.

Although the confined particles of the \( T \) theory are not part of the final condensed anyon theory, they still have physical meaning. The full \( T \) theory can be physically realized as a 1+1 dimensional theory.
living on the edge of a droplet of the $U$ anyon theory living inside a larger region of the $A$ uncondensed theory. The reason for this is that if we restrict motion of particles to a one-dimensional edge, there is no possibility of braiding one particle around each other and there is thus no problem with any of the particles of the full $T$ theory — both the confined and deconfined particles can live there. The $T$ theory, since it describes a 1+1 dimensional edge is not a braided anyon theory, but is rather a fusion algebra (or a planar diagram algebra).

If we try to drag one of the confined particles of the $T$ theory into the condensed $U$ droplet, its nontrivial braiding with the condensate creates a “branch-cut” in the condensate along its path into the condensate and destroys the condensate along this path. This costs an energy proportional to the distance the particle has been dragged into the $U$ region. Thus there is a force pushing these confined particle back to the edge of the droplet. The particles are confined to the edge.

$Z_8^{(3+1/2)}$ Again

Let us return to our example of $Z_8^{(3+1/2)}$ and determine which of our orbits (particle types of $T$) are confined or deconfined. Recall the rule that an orbit is deconfined if all of the constituent particles in the orbit have the same twist factor $\theta$. From Eq. 23.3 we have

$$\begin{align*}
\theta_0 &= \theta_4 = 1 \\
\theta_1 &\neq \theta_5 \\
\theta_2 &= \theta_6 = -i \\
\theta_3 &\neq \theta_7
\end{align*}$$

Thus, the only two particle types allowed in the final condensed $U$ theory are the orbits $[0]$ (which is the identity) and $[2]$ with corresponding spin factors $\theta_{[0]} = 1$ and $\theta_{[2]} = -i$. The only nontrivial fusion we obtain from Eq. 23.4 is $[2] \times [2] = [0]$.

We thus recognize this condensed anyon theory as the (left-handed) semion theory! Further we establish that if we condense a semion droplet within a $Z_8^{(3+1/2)}$ background there will be two additional particle types $([1]$ and $[3])$ that remain confined at the edge of the droplet.

23.4 Splitting: Orbits not of maximum size

Above in section 23.2.1 we assumed all of the orbits were of maximum size. That is, if $N$ is the smallest positive integer so that $J^N = I$, then all orbits are of size $N$.

If we have a situation where some orbits are not of maximum size, then we have a new physical phenomenon, known as splitting. This phenomenon is a reflection of the fact that assigning each orbit $[a]$ from the uncondensed theory $A$ to be a particle type of the intermediate
theory $\mathcal{T}$ will not give an acceptable fusion algebra. Let us see how this happens.

Let us suppose we have some particle $a$ such that $J^p \times a = a$ with $0 < p < N$ (In fact, $p$ must divide $N$). We start by recalling

$$a \times \bar{a} = I + \ldots$$

On the other hand, we can also write

$$a \times \bar{a} = (J^p \times a) \times \bar{a} = J^p \times (a \times \bar{a}) = J^p \times (I + \ldots) = J^p + \ldots$$

We thus conclude that we must have

$$a \times \bar{a} = I + J^p + \ldots$$

Now we claim that this feature will result in the orbits in $\mathcal{A}$ not producing an acceptable fusion algebra in $\mathcal{T}$. As in the previous example, let us divide all of the particles in $\mathcal{A}$ into their orbits under the action of $J$, which we write as $[a]$. The fusion equation Eq. 23.6 then would imply the fusion for the orbits

$$[a] \times [\bar{a}] = [I] + [J^p] + \ldots$$

where here we have used that $[J^p]$ is in the $J$ orbit of the identity (i.e., $[J^p] = [I]$). Eq. 23.7 now presents an inconsistency as one of our rules of fusion algebras (Eq. 8.6) is that $N_{a[a]} = 1$, i.e., the identity field should occur only once on the right hand side. We conclude that this is not acceptable as a fusion algebra for $\mathcal{T}$.

To resolve this problem the orbit $[a]$ must split into multiple particle types in $\mathcal{T}$ which we will write as $[a]_i$ with $i = 1, 2, \ldots, q_a$ for some number $q_a$.

Most generally we can write the mapping between the original $\mathcal{A}$ and $\mathcal{T}$ as

$$a \rightarrow \sum_{i=1}^{q_a} n_a^i [a]_i$$

where now $[a]_i$ are particle types of the $\mathcal{T}$ theory and the $n$’s are integers.

If the orbit of $[a]$ is maximal, then $[a]$ does not need to split, meaning $(n_a^1 = 1$ and $q_a = 1$, and we don’t need to write a subscript on $[a]$). However, if the orbit is not maximal, then $[a]$ must split into multiple different particles $[a]_1, [a]_2, \ldots$ such that the twist factors all agree

$$\theta_{[a]_i} = \theta_a \ .$$

As in the simple case with no splitting, the fusion rules of the $\mathcal{T}$ theory must be consistent with those of the uncondensed $\mathcal{A}$ theory. In particular

$$a \times b = \sum_c N_{a[b]}^c c$$

On the other hand, we can also write

$$a \times \bar{a} = (J^p \times a) \times \bar{a} = J^p \times (a \times \bar{a}) = J^p \times (I + \ldots) = J^p + \ldots$$

We thus conclude that we must have

$$a \times \bar{a} = I + J^p + \ldots$$

Now we claim that this feature will result in the orbits in $\mathcal{A}$ not producing an acceptable fusion algebra in $\mathcal{T}$. As in the previous example, let us divide all of the particles in $\mathcal{A}$ into their orbits under the action of $J$, which we write as $[a]$. The fusion equation Eq. 23.6 then would imply the fusion for the orbits

$$[a] \times [\bar{a}] = [I] + [J^p] + \ldots$$

where here we have used that $[J^p]$ is in the $J$ orbit of the identity (i.e., $[J^p] = [I]$). Eq. 23.7 now presents an inconsistency as one of our rules of fusion algebras (Eq. 8.6) is that $N_{a[a]} = 1$, i.e., the identity field should occur only once on the right hand side. We conclude that this is not acceptable as a fusion algebra for $\mathcal{T}$.

To resolve this problem the orbit $[a]$ must split into multiple particle types in $\mathcal{T}$ which we will write as $[a]_i$ with $i = 1, 2, \ldots, q_a$ for some number $q_a$.

Most generally we can write the mapping between the original $\mathcal{A}$ and $\mathcal{T}$ as

$$a \rightarrow \sum_{i=1}^{q_a} n_a^i [a]_i$$

where now $[a]_i$ are particle types of the $\mathcal{T}$ theory and the $n$’s are integers.

If the orbit of $[a]$ is maximal, then $[a]$ does not need to split, meaning $(n_a^1 = 1$ and $q_a = 1$, and we don’t need to write a subscript on $[a]$). However, if the orbit is not maximal, then $[a]$ must split into multiple different particles $[a]_1, [a]_2, \ldots$ such that the twist factors all agree

$$\theta_{[a]_i} = \theta_a \ .$$

As in the simple case with no splitting, the fusion rules of the $\mathcal{T}$ theory must be consistent with those of the uncondensed $\mathcal{A}$ theory. In particular

$$a \times b = \sum_c N_{a[b]}^c c$$
in the $\mathcal{A}$ theory implies
\[
\left( \sum_{i=1}^{q_a} n_i^a [a]_i \right) \times \left( \sum_{i=1}^{q_b} n_i^b [b]_i \right) = \sum_c N_{a,b}^c \left( \sum_{i=1}^{q_c} n_i^c [c]_i \right)
\]
within the $\mathcal{T}$ theory. This consistency implies the relationship
\[
d_a = \sum_{i=1}^{q_a} n_i^a d_{[a]_i}
\]
(23.10)

between the quantum dimensions in the $\mathcal{A}$ theory (left) and the quantum dimensions in the $\mathcal{T}$ theory (right). Once the particles have split, it is then possible to have a consistent set of fusion rules in the $\mathcal{T}$ theory. Once these fusion rules have been established, one can determine which fields are confined in order to determine the final condensed anyon theory $\mathcal{U}$.

While the phenomenon of identification (section 23.2) has a fairly obvious physical interpretation, it is often not as obvious how to interpret the phenomenon of splitting — except to say that it is required for consistency. However, a physical understanding is given by realizing the presence of a condensate can cause certain physical quantities to be locally conserved where they are indefinite in the uncondensed phase. It is the presence of these new locally conserved quantities which allow us to form $[a]_i$ where $i$ can take $q_a$ different values — corresponding to the $q_a$ different values that the conserved quantity may take. This picture of emergent conserved quantities is elucidated by Burnell et al. [2011, 2012].

Example: $SU(2)_4$

Let us consider the example of the Chern-Simons theory $SU(2)_4$. We list the fields, their quantum dimensions, and their fusion rules in table 23.1.

<table>
<thead>
<tr>
<th>particle</th>
<th>$d$</th>
<th>$\theta$</th>
<th>$\times$</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1</td>
<td>1 [\sqrt{3}]</td>
<td>$e^{2\pi i/8}$</td>
<td>1</td>
<td>0 + 2</td>
<td>1 + 3</td>
<td>2 + 4</td>
</tr>
<tr>
<td>1</td>
<td>$\sqrt{3}$</td>
<td>$e^{2\pi i/3}$</td>
<td>1 + 3</td>
<td>1 + 3</td>
<td>0 + 2 + 4</td>
<td>1 + 3</td>
<td>2</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>$e^{2\pi i/8}$</td>
<td>2 + 4</td>
<td>1 + 3</td>
<td>0 + 2</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>3</td>
<td>1</td>
<td>4</td>
<td>3</td>
<td>2</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

Table 23.1 Data for $SU(2)_4$. Left: quantum dimensions and twist factors for the different particles. Note that 0 is the identity. Right: Nontrivial fusion rules. Note that the fusion rules are given by Eq. (19.10) and the quantum dimensions are given by Eq. (19.2) given $d = \sqrt{3}$. You can check the consistency of $d_a$ with the fusion rules by using Eq. (8.10) (i.e., $d_a$ should be the largest eigenvalue of the fusion matrix $[N_{a,b}^c]$). Particle 4 is a simple current boson which we will attempt to condense.

We notice that particle 4 is a simple current with orbit of length 2...
(4 × 4 = 0). Let us now list the orbits in this theory

\[ [0] = [4] \quad \text{Maximum size orbit (also the identity)} \\
[1] = [3] \quad \text{Maximum size orbit} \\
[2] \quad \text{Not a maximum size orbit. This orbit must split} \]

These orbits are just read off from the bottom line of the table 23.1 under fusion with the field 4, we have 0 mapping to 4 and vice versa; we have 1 mapping to 3 and vice versa, but 2 just maps to itself.

The fusion rules for the orbits, which we read off of table 23.1, are

\[
\begin{align*}
[2] \times [2] &= [0] + [2] + [0]
\end{align*}
\]

where [0] is the identity orbit. It is the last line here which demonstrates explicitly the problem noted in Eq. 23.7 — we should not have the identity twice on the right hand side. To fix this problem we split the particle [2] into two pieces [2]_1 and [2]_2.

\[
\begin{align*}
[1] \times [1] &= [0] + [2]_1 + [2]_2 \\
[1] \times ([2]_1 + [2]_2) &= [1] + [1] \\
([2]_1 + [2]_2) \times ([2]_1 + [2]_2) &= [0] + ([2]_1 + [2]_2) + [0]
\end{align*}
\]

While this is not quite the full fusion rules of the \( T \) theory, one can nonetheless extract a unique set of fusion rules for the \( T \) theory consistent with Eq. 23.11-23.13, which are shown in table 23.2.

<table>
<thead>
<tr>
<th>particle</th>
<th>( d )</th>
<th>( \theta )</th>
<th>( \times )</th>
<th>[1]</th>
<th>[2]_1</th>
<th>[2]_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>[0]</td>
<td>1</td>
<td>1</td>
<td>[1]</td>
<td>[0] + [2]_1 + [2]_2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>[1]</td>
<td>( \sqrt{3} )</td>
<td>( e^{2\pi i/3} )</td>
<td>[1]</td>
<td>[1]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>[2]_1</td>
<td>1</td>
<td>( e^{2\pi i/3} )</td>
<td>[2]_1</td>
<td>[1]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>[2]_2</td>
<td>1</td>
<td>( e^{2\pi i/3} )</td>
<td>[0]</td>
<td>[2]_1</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 23.2 Data for the intermediate \( T \) theory obtained from condensing the 4 particle in \( SU(2)_4 \). This is the fusion theory describing the edge of a condensed droplet.

Several things are worth noting on this table. First, note that \( d_{[2]_1} + d_{[2]_2} \) is the same as \( d_2 \) from the original \( SU(2)_4 \) as is required by Eq. 23.10. Secondly note that the twist factors are unchanged (even if a particle

\[1\text{Eq. 23.10 is already written as a proper fusion rule. Eq. 23.11 implies }[1] \times [2]_1 = [1] \text{ and } [1] \times [2] = [1]. \quad \text{Next we note that the left hand side of Eq. 23.13 can be rewritten as } ([2]_1 \times [2]_1) + ([2]_2 \times [2]_2) + 2 \times [2]_1 \times [2]_2 \text{ which, comparing to Eq. 23.13 immediately implies that } [2]_1 \times [2]_2 = [0]. \quad \text{To pin down the remaining fusion rule we use associativity } [2]_2 = ([2]_1 \times [2]_2) \times [2]_2 = [2]_1 \times [2]_2 \times [2]_2 \text{ which implies the only consistent set of fusion rules to be } [2]_1 \times [2]_2 = [2]_1 \text{ and } [2]_1 \times [2]_2 = [2]_2. \]
Anyon Condensation

type splits) as stated in Eq. 23.9.

We can obtain the final $\mathcal{U}$ anyon theory, with a proper braiding, by throwing out the particles that are confined. Looking back at table 23.1 we see that the orbit $[1]$ is made up of particles 1 and 3 which have different spin factors. This implies that particle $[1]$ must be confined (it braids nontrivially with the condensate, see Eq. 23.5). However, the orbit $[2]$ is made of a single particle type and therefore is deconfined (even if it splits). Thus our final anyon theory after confinement is given by table 23.3. We recognize the resulting anyon theory as $SU(3)_1$ or equivalently $\mathbb{Z}_3^1$. See section 20.4.1.

<table>
<thead>
<tr>
<th>particle</th>
<th>$d$</th>
<th>$\theta$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>i</td>
<td>1</td>
</tr>
<tr>
<td>$[2]_1$</td>
<td>i</td>
<td>$e^{2\pi i/3}$</td>
</tr>
<tr>
<td>$[2]_2$</td>
<td>i</td>
<td>$e^{2\pi i/3}$</td>
</tr>
</tbody>
</table>

Table 23.3 Data for the final $\mathcal{U}$ anyon theory obtained from condensing the 4 particle from $SU(2)_4$. We recognize this theory as $SU(3)_1$ or equivalently $\mathbb{Z}_3^1$.

23.5 Other Features of Condensation

A few other features of condensation are worth mentioning. First, if we start with a modular anyon theory, $\mathcal{A}$, then the condensed theory $\mathcal{U}$ is also modular. Further for modular theories, the central charge (modulo 8) remains unchanged

$$c_{\mathcal{A}} = c_{\mathcal{U}} \mod 8$$  \hspace{1cm} (23.14)

Secondly, there is a beautiful relationship between the total quantum dimensions of the uncondensed theory $\mathcal{A}$, the fusion algebra $\mathcal{T}$ and the final theory $\mathcal{U}$. Recalling that total quantum dimension is defined by

$$D = \sqrt{\sum_a d_a^2}$$

We then have

$$\frac{D_{\mathcal{A}}}{D_{\mathcal{T}}} = \frac{D_{\mathcal{T}}}{D_{\mathcal{U}}}$$  \hspace{1cm} (23.15)

Let us check these relations for the $SU(2)_4$ condensation example. From tables 23.1, 23.2, and 23.3 we obtain

$$D_{\mathcal{A}=SU(2)_4} = \sqrt{12} \quad D_{\mathcal{T}} = \sqrt{6} \quad D_{\mathcal{U}=SU(3)_1} = \sqrt{3}$$

in agreement with Eq. 23.15. Also (From Eq. 17.16) we can calculate that

$$c_{SU(2)_4} = c_{SU(3)_1} = 2 \hspace{1cm} \text{mod 8}$$

in agreement with Eq. 23.14.
23.6 Cosets

In Chern-Simons theory, one of the most common ways to construct new TQFTs is the idea of a coset theory first discussed in the context of conformal field theory (See Di Francesco et al. [1997]; Moore and Seiberg [1989]; Goddard et al. [1985]). Given Lie groups $G$ and $H$ such that $H$ is a subgroup of $G$, we may consider theories $G_k$ (Chern-Simons theory $G$ at level $k$) and correspondingly $H_k'$. There is then a well defined way to make a so-called coset theory which we write as $G_k/H_k'$. One rough physical interpretation of this construction is that we are gauging the subgroup $H$, essentially making these degrees of freedom redundant. One can have more complicated cosets where we embed $H$ into a product of Lie groups $G 	imes G'$, to construct coset theories like $G_k \times G_{k'}/H_{k''}$.

While cosets of this type can seem quite complicated, they actually have an extremely simple interpretation in terms of boson condensation. To construct $G_k/H_k'$ we first construct $G_k \times H_{k'}$ where the overbar means we should switch the chirality of the theory$^{12}$ Then if we condense all possible simple current bosons in the product theory $G_k \times H_{k'}$, we obtain the coset theory $G_k/H_{k'}$. (This is much simpler than the conventional coset construction!) This technique can be generalized in an obvious way. For example, to construct $G_k \times G_{k'}/H_{k''}$ we first construct $G_k \times G_{k'} \times H_{k''}$ and then condense all of the simple current bosons.

Note that given Eq. 23.14 and Eq. 17.27 the central charge of a coset is equal to the sum of central charges in the numerator minus the sum of central charges in the denominator.

**Example: $SU(2)_2/U(1)_2$**

To construct the coset $SU(2)_2/U(1)_2$ we want to first construct $SU(2)_2 \times U(1)_2$ and then condense all simple current bosons. Let us recall the data for $SU(2)_2$ (See section 18.3.1) and $U(1)_2$ (See section 20.4.2) which are shown in Table 23.4.

<table>
<thead>
<tr>
<th>particle</th>
<th>d</th>
<th>$\theta$</th>
<th>$\times$</th>
<th>$\sigma$</th>
<th>$\psi$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$I$</td>
<td>1</td>
<td>$\frac{1}{16}$</td>
<td>$e^{2\pi i/16}$</td>
<td>$\sigma$</td>
<td>$I$</td>
</tr>
<tr>
<td>$\sigma$</td>
<td>$\sqrt{2}$</td>
<td>$-1$</td>
<td>$-$</td>
<td>$\psi$</td>
<td>$\sigma$ $I$ $+\psi$</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>particle</th>
<th>d</th>
<th>$\theta$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>$e^{2\pi i/8}$</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>$-1$</td>
</tr>
<tr>
<td>3</td>
<td>1</td>
<td>$e^{2\pi i/8}$</td>
</tr>
</tbody>
</table>

Table 23.4 Data for $SU(2)_2$ (top) and $U(1)_2$ (bottom). The overline indicates we take the mirror image theory, meaning all of the twist factors are complex conjugated compared to the definition given in section 20.4.2.

$^{12}$In Chern-Simons theory, $G_{-k} = \overline{G_k}$. Switching chirality in the diagrammatic algebra can be achieved by evaluating the mirror image of the diagram.
The product theory $SU(2)_2 \times U(1)_2$ has particles types of the form $(a, b)$ where $a$ is from the $SU(2)_2$ theory and $b$ is from the $U(1)_2$. The twist factor of such a product particle is $\theta_{(a,b)} = \theta_a \theta_b$.

One can see from the table that the product particle $(\psi, 2)$ is a boson simple current (twist factor $\theta = 1$), so we can condense it. (In fact, not including the identity particle $(I, 0)$, this is the only boson simple current). There are 12 particles in the product theory which divide into 6 orbits which all are of maximum size (so there is no splitting). These orbits (under the action of fusion with the $(\psi, 2)$ boson) are

\[
[I, 0] = [\psi, 2] \quad [\sigma, 0] = [\sigma, 2] \quad [\psi, 0] = [I, 2] \\
[I, 1] = [\psi, 3] \quad [\sigma, 1] = [\sigma, 3] \quad [\psi, 1] = [I, 3]
\]

These are the six particle types of the $T$ theory. Finally, examining the twist factors we can see that only the three orbits $[I, 0]$ and $[\sigma, 1]$ and $[\psi, 0]$ are deconfined. The final condensed anyon theory $U = SU(2)_2/U(1)_2$ is given in table 23.5. We recognize this result as being simply the Ising TQFT (See section 18.3.1).

<table>
<thead>
<tr>
<th>particle</th>
<th>$d$</th>
<th>$\theta$</th>
<th>$\times$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$[I, 0]$</td>
<td>1</td>
<td>1</td>
<td>$[\sigma, 0]$</td>
</tr>
<tr>
<td>$[\sigma, 1]$</td>
<td>$\sqrt{2}$</td>
<td>$e^{2\pi i/16}$</td>
<td>$[\sigma, 0]$</td>
</tr>
<tr>
<td>$[\psi, 0]$</td>
<td>1</td>
<td>$-1$</td>
<td>$[\sigma, 0]$</td>
</tr>
</tbody>
</table>

Table 23.5 The final anyon theory data for the coset $SU(2)_2/U(1)_2$ is just the Ising TQFT.

## 23.7 More General Condensations

One can also consider condensations of objects that are not simple currents. There is a fairly simple rule for constructing all possible condensations of a modular anyon theory which has been worked out by Neupert et al. [2016].

Given a modular anyon theory (a modular tensor category or TQFT) with $N$ particle types. Each condensation is described by a matrix $M$ with nonnegative integer entries with the following properties

\[
[M, S] = [M, T] = 0 \quad (23.16)
\]

with $M_{I, I} = 1$ where $S$ and $T$ are the modular $S$ and $T$ matrices respectively (See section 17.3). The $M$ matrix must be of the form

\[
M = nn^T \quad (23.17)
\]

where $n$ is a rectangular matrix with nonnegative integer entries and $T$ means transpose. The $n$ matrix is the same coefficients as in Eq. 23.8
except we only include resulting particles that are deconfined (i.e., we count the particles in the $U$ theory and not all the particles in the $T$ theory).

From the $n$ matrix we construct the $S$ and $T$ matrices of the condensed ($U$) theory as

$$S_n = n S_{\text{condensed}} \quad T_n = n T_{\text{condensed}}$$

(23.18)

We must then confirm that $S_{\text{condensed}}$ is valid meaning that it is symmetric, unitary, and generates non-negative fusion coefficients with the Verlinde formula (Eq. 17.13).

Note that $M$ being a permutation matrix is a valid solution and this simply permutes the particles of the original theory and should not be considered a condensation.

### 23.8 Condensation and Boundary Modes

[MOVE THIS SECTION? OR remove reference to topological phases]

Many topological phases of matter have the property that they must have gapless modes along their boundary when you have a finite region of the matter (say, a droplet or a disk). We will explore this in detail for quantum Hall states in section *** below. Here however, we introduce some general principles.

As mentioned in section 17.3.3, if the central charge of the TQFT is nonzero, then the edge of the system necessarily carries heat and therefore must be gapless (i.e., has low energy thermal excitations). However, even when the central charge is zero and the thermal conductance is zero, there can be cases where the edges of a system must be gapless. It turns out that the condition for this to be true is intimately related to condensation! Without giving detailed proof we will state the result (See Refs. Burnell [2018]; Kong [2014]; Levin [2013])

The edge of a topologically ordered system can only be gapped if one can condense a set of bosons from this system and obtain a topologically trivially ordered system (a system with no anyons).

(See footnote [2] from this chapter). The converse of this statement is not generally true, as a gapped system can always be made gapless by altering the Hamiltonian a bit near the edge to allow gapless excitations.

While we won’t give a any proof of this\(^\text{13}\), but instead just give a cartoon of what it means. It is perhaps easier to think about a general boundary between a TQFT $C$ and another one $D$. If we think of the Hamiltonian transitioning slowly as a function of position between the region of $C$ and $D$ our above arguments are that a smooth transition occurs only if $C$ can be obtained by a condensation transition from $D$ or vice versa. It is exactly this type of smooth transition that we need in order to avoid a gapless mode between the two regions (at a cartoon level one can think about some abstract order parameter vector changing

\(^\text{13}\)Indeed, while there is no rigorous proof of this criterion, although the arguments given in the literature are fairly strong! See Kong [2014]; Levin [2013] for example.
direction in some abstract space as we move from one region to another). This criterion (that $\mathcal{D}$ should come from $\mathcal{C}$ by condensation, or vice versa) suffices to justify the above condition for when we can have a gapless edge to the vacuum if we just choose, say $\mathcal{D}$ to be the vacuum (the trivial TQFT). However, it does not actually give the right rule for determining if one can have gapless modes on the boundary between regions of $\mathcal{C}$ and of $\mathcal{D}$.

In fact the correct, more general rule [Bais et al. 2009] is given by:

The boundary between two topologically ordered systems $\mathcal{C}$ and $\mathcal{D}$ can only be gapped if the system $\mathcal{C} \times \bar{\mathcal{D}}$ can have a gapped edge to the trivial theory, i.e., if a set of bosons from $\mathcal{C} \times \bar{\mathcal{D}}$ can be condensed to give a topologically trivially ordered system.

If $\mathcal{C}$ is obtained from $\mathcal{D}$ by a boson condensation, or vice versa, then this criterion is certainly satisfied. However, there are cases where $\mathcal{C}$ is not obtained from $\mathcal{D}$ by condensation (and neither $\mathcal{D}$ from $\mathcal{C}$) and yet this criterion can be satisfied. The reason for this difference is that this criterion allows another physical process whereby a particle near the boundary of $\mathcal{C}$ binds with a particle near the boundary of $\mathcal{D}$ to form an object that condenses along the boundary.

Further Reading

Bais and Slingerland [2009] is the original discussion of anyon condensation. Full disclosure, I was the referee for this paper. I spent a lot of time reading it in great detail and I ended up deciding it was pretty brilliant.

A nice review of the physics of anyon condensation is given by Burnell [2018].

Eliëns et al. [2014] gives methods of extracting detailed data, such as $F$-matrices of a condensed theory from the data for the uncondensed theories.

Neupert et al. [2016] is also a good reference on the mathematics of condensation.

Much of the structure of anyon condensation (at least for condensation of simple currents) was worked out originally in the context of so-called “fixed point resolutions” of conformal field theories. See for example, Schellekens [1999].

A more mathematical discussion of condensation is given by Kong [2014].

Chapter 40 shows how to use the computer program Kac to work out the results of certain condensations.

Exercises

Exercise 23.1 Fusion of Bosonic Simple Currents
Given an anyon theory with a bosonic simple current \( J \), such that \( R^J_J^2 = 1 \) and \( J^N = I \), show that all of the particle types \( J^p \) with \( 0 < p < N \) are also bosons, and further that the braiding any two of these particle types is trivial \( R^{J^p, J^p} = 1 \). Hint: This is fairly trivial given some results of chapter 20.

Exercise 23.2 Condensation to the Vacuum
The toric code is one of the simplest TQFTs, which we will study in great depth in later chapters. The particle types and fusion relations are given in Table 26.1, and the \( S \)- and \( T \)-matrices are given in Eq. 26.5.

Show that there are two possible bosons that can condense, and that the resulting \( \mathcal{U} \) theory in either case is the trivial TQFT (no particles but the vacuum).

Exercise 23.3 Splitting
Consider \( \text{SU}(2)_2 \times \text{SU}(2)_2 \). There is a single simple current boson that can be condensed. Find the \( \mathcal{T} \) theory (there is a splitting!) and the final \( \mathcal{U} \) theory after condensation.

Exercise 23.4 Cosets
(a) Calculate the properties of the coset \( \text{SU}(16)_1 / \text{SU}(2)_2 \).
(b) Calculate the properties of the coset \( \text{SU}(2)_1 \times \text{SU}(2)_1 / \text{SU}(2)_2 \).

Exercise 23.5 General Condensation Method
The \( S \)-matrix for \( \text{SU}(2)_4 \) is given by

\[
S_{ij} = \frac{1}{\sqrt{3}} \sin \left( \frac{(i+1)(j+1)\pi}{6} \right)
\]

where \( i, j \in 0, \ldots, 4 \). (If you are adventurous you can confirm this formula by using Table 23.1 and Eq. 17.20.)

Using the method of section 23.7 confirm that the matrix \( n \) given by

\[
\begin{pmatrix}
1 & 0 & 0 & 0 & 1 \\
0 & 0 & 1 & 0 & 0 \\
0 & 0 & 1 & 0 & 0
\end{pmatrix}
\]

satisfies Eqs. 23.10 and 23.16.

Show that the \( S \)- and \( T \)-matrices of \( \text{SU}(3)_1 \) or equivalently \( \mathbb{Z}_3^1 \) (See section 20.4.1) satisfy the requirement for the condensed phase given by Eq. 23.18.
Part V

Toric Code Basics
Introducing Quantum Error Correction

Before we look at the toric code, it is worth introducing some ideas of quantum error correction. While initially the ideas of error correction may seem somewhat different from what we have been discussing, we will see that (at least some quantum error correcting codes) are extremely closely related to the topological ideas we have been discussing in prior chapters. Some of this material may be well known to most readers, but we reiterate it for completeness and to orient the discussion.

24.1 Classical Versus Quantum Information

24.1.1 Memories

Classical Memory

The unit of classical information is a bit\(^2\) — a classical two state system which can take the values 0 or 1. A memory with \(N\) bits can be in any one of \(2^N\) states — each state corresponding to a particular bit-string, such as 011100111.

Quantum Memory

The unit of quantum information is the quantum bit or qubit which is a quantum two state system — i.e. a two-dimensional complex Hilbert space spanned by vectors which we usually call \(|0\rangle\) and \(|1\rangle\). A qubit can be in any state

\[ |\psi\rangle = \alpha|0\rangle + \beta|1\rangle \]

with arbitrary complex prefactor \(\alpha, \beta\) (where we normalize wavefunctions so \(|\alpha|^2 + |\beta|^2 = 1\)).

A quantum memory with \(N\) qubits is a vector within a \(2^N\) dimensional complex Hilbert space. So for example, with 2 qubits the general state of a system is specified by four complex parameters

\[ |\psi\rangle = \alpha|00\rangle + \beta|01\rangle + \gamma|10\rangle + \delta|11\rangle \]  

(24.1)

with the normalization condition \(|\alpha|^2 + |\beta|^2 + |\gamma|^2 + |\delta|^2 = 1\). So to specify the state of a quantum memory with 2 bits, you have to specify four complex parameters, rather than, in the classical case just stating which of the four states the system is in.
24.2 **Errors**

An error is some process which accidentally changes the state of the memory away from the intended state. Often we take as an error model the case where only one bit or one qubit is affected at a time (a “minimal” error) although more complicated errors can occur in practice.

24.2.1 **Classical Error Correction**

There is a simple way to protect the information stored in a classical memory from errors. Instead of storing a single bit 0 or 1, we will store multiple copies of the bit. For example in Table 24.1 we use three “physical” bits to store one “logical” bit of information.

<table>
<thead>
<tr>
<th>logical bit</th>
<th>physical bits</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>000</td>
</tr>
<tr>
<td>1</td>
<td>111</td>
</tr>
</tbody>
</table>

Table 24.1 Three bit repetition code. Stores a single logical bit of information using three physical bits. The code space is the set 000 and 111.

Our memory should always either be in the state 000 or 111 — we call these two possibilities the code space. If we detect the system being in any other state of the three bits (i.e., not in the code space) we know an error has occurred. If an error does occur on one of the physical bits (i.e., if one of the bits is accidentally flipped) we can easily find it, because it would leave our memory with not all of the physical bits being the same. For example, if our system starts as 000, an error introduced on the second bit would leave it in the form 010. Then, by just using a majority-rule correction system, it is easy to figure out what happened and flip the mistaken bit back. So our error correction protocol would be to continuously compare all three bits, if they don’t all match, flip the one which would bring them back to matching. Assuming errors are rare enough (and only occur on one bit at a time\(^3\)) this scheme is an effective way to prevent errors. For added protection one can use more redundant physical bits, such as five physical bits or seven physical bits for a single logical bit. Such larger codes could withstand several bit-flip errors at a time and would still allow successful correction. For example, the five bit code could withstand two bit-flip errors at a time and correction via majority rule would still be successful.

One might think the same sort of approach would work in the quantum world: make several copies of the qubit you want to protect, and then compare them to see if one has changed. Unfortunately, there are two big problems with this. The first is the so-called no-cloning theorem — it is not possible to make a perfect clone of a qubit. The second problem is that measuring a state inevitably changes it.

\(^3\)If two bit flips happen at the same time, then an uncorrectable logical error occurs. It is thus imperative that we check the state of our physical bits very frequently so that we catch errors and correct them before multiple errors can occur.
24.3 Quantum No Cloning Theorem

Because the quantum no-cloning theorem is such an integral part of the discussion surrounding quantum error correction (and because the proof is easy) it is worth going through the proof.

The result (usually credited to Wootters and Zurek [1982] and Dieks [1982]) is such a straightforward result of quantum mechanics that some people have argued whether it deserves to be called a theorem. Nonetheless, the statement of the “theorem” is as follows:

**Theorem**: Given a qubit in an arbitrary unknown state $|\phi_1\rangle$ and another qubit in a known initial state $|\phi_2\rangle$, there does not exist any unitary operator $U$ (i.e., any quantum mechanical evolution) such that

$$U(|\phi_1\rangle \otimes |\phi_2\rangle) = e^{i\chi(\alpha)} |\phi_1\rangle \otimes |\phi_1\rangle$$

for all possible input $|\phi_1\rangle$.

The point of the theorem is that there is no way to copy $|\phi_1\rangle$ into the auxiliary qubit $|\phi_2\rangle$. The reason we are looking for a unitary operator $U$ is that all time evolutions in quantum mechanics correspond to unitary operators. The phase $e^{i\chi}$ allows some arbitrary phase to occur during the cloning process which we can allow to be a function of the cloned state $|\phi_1\rangle$.

**Proof of Theorem**: Suppose such a unitary operator as specified in Eq. (24.2) did exist. This means we can properly copy two orthogonal states $|0\rangle$ and $|1\rangle$, meaning

$$U(|0\rangle \otimes |\phi_2\rangle) = e^{i\chi(0)} |0\rangle \otimes |0\rangle$$
$$U(|1\rangle \otimes |\phi_2\rangle) = e^{i\chi(1)} |1\rangle \otimes |1\rangle$$

Quantum mechanical operators must be linear so we can try applying this operator to the linear superposition $\alpha|0\rangle + \beta|1\rangle$ and we must get

$$U(\{ \alpha|0\rangle + \beta|1\rangle \} \otimes |\phi_2\rangle) = \alpha e^{i\chi(0)}|0\rangle \otimes |0\rangle + \beta e^{i\chi(1)}|1\rangle \otimes |1\rangle.$$ 

But this is not what a putative cloning device must give. Instead a clone of the bit should have given the outcome

$$e^{i\chi(\alpha|0\rangle + \beta|1\rangle)} \left[ \alpha|0\rangle + \beta|1\rangle \right] \otimes \left[ \alpha|0\rangle + \beta|1\rangle \right]$$

which is not generally the same result. Thus no cloning device is consistent with the linearity inherent in quantum mechanical evolution. □

---

4One can object that one can make a measurement that has the effect of projecting, rather than a unitary operation. However, there a philosophy known sometimes as “The Church of the Larger Hilbert Space” which says that we should simply treat our measurement apparatus as part of the quantum mechanical system, in which case all time evolutions become unitary again. The reason it is called a “church” is because it is almost a religious view of how one should think about quantum measurements.
24.4 Quantum Error Correction

Perhaps the most surprising thing about quantum error correction is that it is possible at all! This was discovered in 1995 by Shor, and shortly thereafter by Steane.

In the next chapter we will introduce the toric code (Kitaev), which is a quantum error correcting code closely related to anyons. However, here, we will briefly introduce a very simple error correcting code introduced in the original work by Shor to try to explain how such codes typically work.

24.4.1 Bit Flip Correcting Code

Let us consider the following simplified problem. Suppose we know that the only error that can ever occur on our physical system is the application of a Pauli \( \sigma_z \) operator, i.e., a so-called bit-flip error. We can protect our qubit from such an error in the following way:

Given a qubit

\[ |\psi_1\rangle = \alpha |0\rangle + \beta |1\rangle \]

let us encode this physical qubit into three logical qubits

\[ |\psi_4\rangle = \alpha |000\rangle + \beta |111\rangle \]

where by \(|0\rangle\) we mean \( |0\rangle \otimes |0\rangle \otimes |0\rangle \) and similarly for \(|111\rangle\). The code space is the Hilbert space spanned by wavefunctions of the form Eq. 24.4.

We must not measure any of the three physical bits since this will collapse the entire wavefunction to either \(|000\rangle\) or \(|111\rangle\). However, we can measure the product of two bits, such as

\[ \hat{O}_{12} = \sigma_1^x \sigma_2^x \quad \hat{O}_{23} = \sigma_2^x \sigma_3^x \]

The wavefunction \(|\psi_4\rangle\) is in a +1 eigenstate of the operator \( \hat{O}_{12} \) so we can measure this operator without collapsing the wavefunction. The purpose of this operator is to check that bits 1 and 2 are the same. Similarly \( \hat{O}_{23} \) checks that bits 2 and 3 are the same. These operators are known as stabilizer operator (these operators leave the code space unchanged) and their eigenvalues are known as the syndrome since they are meant to diagnose whether the wavefunction has developed any "sickness", i.e., whether an error has occurred.

---

6 The discussion that follows (the remainder of this chapter) is included for completeness, but is not crucial on a first reading.

7 This does not violate the no-cloning theorem, since we are entangling two qubits with the initial qubit, rather than cloning the initial qubit. This procedure can be achieved with a quantum circuit as shown in Fig. 24.1.

8 When we use Pauli matrices we are thinking of \(|0\rangle\) as being spin up and \(|1\rangle\) as being spin down. So \(\sigma_x|0\rangle = |0\rangle\) and \(\sigma_x|1\rangle = -|1\rangle\).

9 We can also define \(\hat{O}_{13}\), but this is redundant information since

\[ \hat{O}_{13} = \hat{O}_{12}\hat{O}_{23} \]

Peter Shor is probably the single person most responsible for creating the field of quantum computing. While there were a few early pioneers in the 1980’s, such as Feynman, Yuri Manin (see footnote 19 of chapter 2), and David Deutsch, there was not a lot of interest in the field for two key reasons: (a) it was not clear you could do anything useful with a quantum computer, and (b) it was not clear you could actually build a quantum computer. Shor conquered both of these questions. In 1994, Shor invented the quantum algorithm for efficiently factoring large numbers. Being that most cryptography schemes rely on large numbers not being efficiently factorizable this was obviously (a) useful, particularly to governments and spy agencies who were then willing to invest large amounts of money into pursuing this type of science. The following year, Shor invented error correcting codes, which then strongly suggested that (b) quantum computers could, at least in principle, be built.
Quantum Error Correction

### Table 24.2

<table>
<thead>
<tr>
<th>Error created by</th>
<th>Stabilizers with $-1$ eigenvalue</th>
<th>Fix by applying</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\sigma_x^1$</td>
<td>$\hat{O}_{12}$</td>
<td>$\sigma_x^1$</td>
</tr>
<tr>
<td>$\sigma_x^2$</td>
<td>$\hat{O}<em>{12}$, $\hat{O}</em>{23}$</td>
<td>$\sigma_x^2$</td>
</tr>
<tr>
<td>$\sigma_x^3$</td>
<td>$\hat{O}_{23}$</td>
<td>$\sigma_x^3$</td>
</tr>
</tbody>
</table>

If an error is created by the operator in the first column, the stabilizer(s) in the middle column will be measured to have a $-1$ eigenvalue, indicating an error. The error is corrected with the operator in the right column.

If we find that both operators $\hat{O}_{12}, \hat{O}_{23}$ are in the $+1$ eigenstate, then we conclude that all three bits are the same, so the wavefunction is properly in the code space (i.e., is of the form of Eq. 24.4). However, if one (or both) of these operators are measured in the $-1$ eigenstate, then we know that an error has occurred. Assuming no more than one $\sigma_x$ error has occurred (i.e., we start with a wavefunction of the form of Eq. 24.4 and only one physical bit is flipped over) we can easily identify the problematic bit. For example, if $\sigma_x$ has been applied to bit 1, then we would find $\hat{O}_{12}$ in the $-1$ eigenstate but $\hat{O}_{23}$ remains in the $+1$ eigenstate. We can then correct the problematic bit by flipping it over again with $\sigma_x$. The error detection and correction rules are given in Table 24.2.

Note that these stabilizer operators do not collapse superpositions of the form of Eq. 24.4 since they do not actually measure the value of any of the bits, they only check to see if two bits are the same as each other.

One might think that Eq. 24.4 (along with the error correction rules of Table 24.2) would constitute a quantum error correcting code. Unfortunately, this is not the case. While we have constructed a code which can correct errors created by $\sigma_x$ (i.e., bit flip errors) one can also have sign errors created by $\sigma_z$ (Errors created by $\sigma_y$ can be thought of as the product of $\sigma_z$ and $\sigma_x$). Applying a $\sigma_z$ operator to any of the three bits in Eq. 24.4 results in

$$|\psi_3\rangle = \alpha|000\rangle - \beta|111\rangle$$

and this cannot be detected by our stabilizers $\hat{O}_{12}$ and $\hat{O}_{23}$.

### 24.4.2 Nine Qubit Shor Code

Shor [1995] found that it is indeed possible to protect a qubit from both $\sigma_x$ and $\sigma_z$ errors by using 9 physical qubits. Again we consider a general qubit of the form

$$|\psi\rangle = \alpha|0\rangle + \beta|1\rangle$$
Introducing Quantum Error Correction

Shor encoded the qubit into nine physical bits in the following way

\[
|\psi_9\rangle = \frac{\alpha}{\sqrt{2}} \left[(|000\rangle + |111\rangle) \otimes (|000\rangle + |111\rangle) \otimes (|000\rangle + |111\rangle)\right] + \frac{\beta}{\sqrt{2}} \left[(|000\rangle - |111\rangle) \otimes (|000\rangle - |111\rangle) \otimes (|000\rangle - |111\rangle)\right]
\]

The code space is the Hilbert space spanned by wavefunctions of the form of Eq. 24.6. The stabilizers of this code are then as follows:

\[
\hat{O}_{12} = \sigma_1^z \sigma_2^z \quad \hat{O}_{23} = \sigma_2^z \sigma_3^z \\
\hat{O}_{45} = \sigma_1^z \sigma_2^z \quad \hat{O}_{56} = \sigma_2^z \sigma_3^z \\
\hat{O}_{78} = \sigma_1^z \sigma_2^z \quad \hat{O}_{89} = \sigma_2^z \sigma_3^z
\]

\[
\hat{O}_{1-6} = (\sigma_1^x \sigma_2^x \sigma_3^x)(\sigma_4^x \sigma_5^x \sigma_6^x) \\
\hat{O}_{4-9} = (\sigma_1^x \sigma_2^x \sigma_3^x)(\sigma_7^x \sigma_8^x \sigma_9^x)
\]

The more interesting stabilizers are given in Eqs. 24.10 and 24.11. These stabilizers are meant to detect, and allow correction, of sign errors, i.e., errors produced by application of \(\sigma_z\) operators. While these stabilizers look somewhat different, in fact these are again quite similar to the simple stabilizers we considered in Eq. 24.5 but in a different basis. Let us define

\[
|+\rangle = \frac{1}{\sqrt{2}} (|000\rangle + |111\rangle) \\
|\rangle = \frac{1}{\sqrt{2}} (|000\rangle - |111\rangle)
\]

With this notation, our error correcting code Eq. 24.6 appears as

\[
|\psi_9\rangle = \alpha |++\rangle + \beta |--\rangle
\]

which we recognize as being the same form as Eq. 24.3. Further let us define \(X\) to be the operator that flips over one of these effective qubits and \(Z\) to be the operator that measures the qubit, as follows

\[
X|\pm\rangle = |\mp\rangle \\
Z|\pm\rangle = \pm|\pm\rangle
\]
Note now that

\[
\sigma_j^z |\pm\rangle = \frac{1}{\sqrt{2}} (|000\rangle \pm |111\rangle) \quad (24.12)
\]

for \( j = 1, 2 \) or 3. So that \( \sigma_j^z \) with \( j = 1, 2 \) or 3 plays the role of the \( X \) operator (the bit-flip operator) on the \( |\pm\rangle \) qubit made from the first three qubits (so let us call this \( X_{123} \)). Note that the effect of all three \( \sigma_j^z \) on this wavefunction is the same.

Similarly we have

\[
(\sigma_1^x \sigma_2^x \sigma_3^x) |\pm\rangle = (\sigma_1^x \sigma_2^x \sigma_3^x) \frac{1}{\sqrt{2}} (|000\rangle \pm |111\rangle) \quad (24.13)
\]

So that \( (\sigma_1^x \sigma_2^x \sigma_3^x) \) plays the role of \( Z \) operating on the \( |\pm\rangle \) qubit made from the first three spins (let us call this \( Z_{123} \)).

With this notation, the stabilizers in Eqs. 24.10 and 24.11 can then be rewritten as operators on three of these effective qubits

\[
\hat{O}_{1-6} = Z_{123}Z_{456} \quad \hat{O}_{4-9} = Z_{456}Z_{789}
\]

entirely analogous to the stabilizers Eqs. 24.5. These stabilizers can detect when \( X_{123} \) or \( X_{456} \) or \( X_{789} \) are applied to the system. But these \( X \) operators are equivalent to the \( \sigma_z \) operators on the original qubits. Thus these two stabilizers can detect \( \sigma_z \) errors, and tell us what operator to apply in order to repair the errors.

Thus we are able to write out Table 24.3 which gives the full error detection and correction rules for the code Eq. 24.6. A few things are worth noting about this table. First, consider the last three rows, which address sign errors created by \( \sigma_z \) operators. Note that, for example, an error created by \( \sigma_1^z \) can be corrected by either \( \sigma_1^z \) or \( \sigma_2^z \) or \( \sigma_3^z \). Second, note that the table addresses the possibility of errors being created by \( \sigma_x \) or \( \sigma_z \). If an error is created by \( \sigma_y \) it is simply considered as the product of \( \sigma_y \) and \( \sigma_x \), and can be corrected accordingly. All Hermitian operations that can be performed on a single qubit are some linear combination of \( \sigma_x, \sigma_y \) and \( \sigma_z \), so this code allows correction of any error that can occur on a single qubit.
Introducing Quantum Error Correction

Table 24.3 Error Detection and Correction Rules for Eq. 24.4. If an error is created by the operator in the first column, the stabilizer(s) in the middle column will be measured to have a $-1$ eigenvalue, indicating an error. The error is corrected with the operator in the right column.
Introducing the Toric Code

We will describe the toric code approach to error correction which is conceptually one of the simplest error correction scheme, as well as being very possibly the best error correction scheme by many measures\(^1\). We will see that the toric code is essentially a topological quantum field theory, which is why we are studying it in this book! As with so many great ideas in this field, the toric code was invented by Kitaev [1997]\(^2\).

25.1 Toric Code Hilbert Space

We imagine an \(N_x\) by \(N_y\) square lattice with a qubit (or spin-\(\frac{1}{2}\)) on each edge, as shown in Fig. 25.1, where the boundaries of the lattice are made periodic hence forming a torus (hence the name “toric”). The total number of spins is \(N = 2N_xN_y\) and correspondingly the dimension of the Hilbert space is \(2^N\).

\[\Rightarrow\]

Fig. 25.1 The Hilbert space of the toric code: An \(N_x\) by \(N_y\) square lattice with qubits (or spin-\(\frac{1}{2}\)), represented as dots, on each edge. The lattice is wrapped up to make it periodic in both directions. i.e., the orange dashed lines at the top and the bottom are identified, and the orange dotted lines at the left and right are identified. Once the system is made periodic this is topologically a torus There are 72 spins in this picture so the Hilbert space has dimension \(2^{72}\).

\(^1\)The statement that it is the “best” is based on the fact that the so-called surface codes (which is essentially the toric code) has the highest known error threshold. The idea of a threshold is the following: If you can make measurements on each bit at a rate of one per second, and similarly you can perform operations on your qubit at a rate of one per second, how many errors per second can you successfully correct? If you can correct errors sufficiently faster than they are created you can prevent any errors from occurring and your quantum information will live forever. Codes are compared to each other to see which one can withstand the greatest error rate, and very often the toric code wins. (See \*\*\*\*\*\*\*\*\*\*). One should be cautioned that the rules of the competition can be modified to favor one or another type of code. (Different rules corresponding to different types of potential hardware). For example, one might declare that performing a measurement on a qubit is faster than flipping qubits. One might declare that single bit operations are faster than two-qubit operations, and so forth.

\(^2\)Alexei Kitaev is one of the great geniuses of modern physics (MacArthur Genius Fellow, Breakthrough Prize Winner, etc., etc.). He is perhaps best known for inventing several of the most important algorithms of quantum computing (See for example the discussion of Kitaev-Solovay in section \[11.1.1\]), but in fact these are not even a small fraction of his highly influential works which span a wide range of modern fields. He visibly gets excited and smiles when he realizes something new about physics, and (being that this seems to happen frequently) he also seems to be one of the happiest people around.
We will think of the qubits along the edges as spin-\(\frac{1}{2}\) particles. We choose to work with a basis of up and down spins for our Hilbert space (\(\sigma_z\)-eigenstates). I.e, we have \(2^N\) basis states with all possible combinations of some spins pointing up and some spins pointing down. A convenient notation for this basis is given by coloring the edges containing down spins blue but leaving uncolored (i.e, black) the edges with up spins as shown in Fig. 25.2.

![Fig. 25.2 A particular basis state of the Hilbert space, working in the up-down basis (\(\sigma_z\)-eigenstates). Here we denote down spins by thick (blue) lines. Up spins are denoted by not uncolored edges (black).](image)

### 25.2 Vertex and Plaquette Operators

Let us now define some simple operators on this Hilbert space which we will use to build the toric code.

**Vertex Operators**

The first operator we define is known as a vertex operator. Given a vertex \(\alpha\) which consists of four incident edges \(i \in \text{vertex } \alpha\), we define the vertex operator as

\[
V_{\alpha} = \prod_{i \in \text{vertex } \alpha} \sigma_z^{(i)}
\]  

(25.1)

I.e., this is just the product of \(\sigma_z\) applied to the four spins incident on the vertex. This operator simply counts the parity of the number of down spins (number of colored edges) incident on the vertex. It returns +1 if there are an even number of incident down spins at that vertex and returns −1 if there are an odd number. This is depicted graphically in Fig. 25.3.

Note that since the eigenvalues of \(V_{\alpha}\) are +1 or −1, we have

\[
V_{\alpha}^2 = 1
\]

(25.2)

which can also be seen by just squaring \(V_{\alpha}\) in the defining Eq. (25.1) and using the fact that \(\sigma_z^2 = 1\). On a lattice of \(N_x\) by \(N_y\) sites, there are a total of \(N = N_x N_y\) different vertex operators.
25.2 Vertex and Plaquette Operators

Vertex and Plaquette Operators

Fig. 25.3 The vertex operator $V_\alpha$ returns +1 if there are an even number of incident down spins (blue lines) at vertex $\alpha$ and returns −1 if there are an odd number.

Plaquette Operators

We now define a second operator known as the plaquette operator. Given a plaquette $\beta$ which contains four edges in a square (edge $i \in$ plaquette $\beta$) we define

$$P_\beta = \prod_{i \in \text{plaquette } \beta} \sigma_x^{(i)}$$

(25.3)

which flips the state of the spins on all of the edges of the plaquette as depicted in Fig. 25.4. On an $N_x \times N_y$ lattice, there are a total of $N_xN_y$ plaquette operators.

Fig. 25.4 The plaquette operator (bright pink) flips the state of the spin on the four edges of a plaquette. Blue edges become black and black edges become blue. The small black arrow points to a particular vertex to emphasize that the number of blue edges incident on this vertex has the same parity both before and after the plaquette flip (implying that the plaquette and vertex operators commute as mentioned in Eq. 25.7 below).

Since flipping a plaquette brings one back to the original configuration we have

$$P_\beta^2 = 1$$

(25.4)

which can also be seen by squaring $P_\beta$ in the defining Eq. 25.3. Given Eq. 25.4 the only two possible eigenvalues of $P_\beta$ are $\pm 1$. The eigenvectors corresponding to each of these eigenvalues can also be determined. In the basis we are using, the spin-up/spin-down basis corresponding to
Introducing the Toric Code

uncolored/colored edges, the $P_\beta$ operator is off-diagonal — it flips spins around a plaquette. Thus the eigenvectors can be written as

\begin{align*}
\text{eigenvector with } +1 \text{ eigenvalue} &= \frac{1}{\sqrt{2}} \left( \begin{array}{c}
\text{unflipped plaquette} \\
\text{flipped plaquette}
\end{array} \right) \\
\text{eigenvector with } -1 \text{ eigenvalue} &= \frac{1}{\sqrt{2}} \left( \begin{array}{c}
\text{unflipped plaquette} \\
-\text{flipped plaquette}
\end{array} \right)
\end{align*}

(25.5) (25.6)

Two examples of such eigenvectors with +1 eigenvalue are shown in Fig. 25.5.

|\Psi\rangle = \frac{1}{\sqrt{2}} \left( \begin{array}{c}
\begin{array}{c}
\text{blue lines}
\end{array} \\
\begin{array}{c}
\text{blue lines}
\end{array}
\end{array} \right) + \frac{1}{\sqrt{2}} \left( \begin{array}{c}
\begin{array}{c}
\text{blue lines}
\end{array} \\
\begin{array}{c}
\text{blue lines}
\end{array}
\end{array} \right)

\begin{align*}
\text{Caution: In this section we define vertex operators (Eq. 25.1) with } \\
\sigma_z \text{ operators and plaquette operators with } \sigma_x \text{ operators (Eq. 25.3). In Kitaev's original work the convention is the opposite: vertex operators are defined as a product of } \sigma_x \text{ operators and plaquette operators are a product of } \sigma_z. \text{ Kitaev's convention is used more commonly in the quantum information community, while our convention is perhaps more typical in the condensed matter community. As we will see in section 29.7 below, Kitaev's convention also makes the toric code look more like a lattice gauge theory. We can transform between the two conventions in either of two ways. Method 1: we can rotate all of our spins so as to exchange } x \text{ and } z. \text{ Method 2: we can make a duality transform on the lattice so that plaquettes become vertices and vertices become plaquettes. This duality approach will be discussed in section *** below.}
\end{align*}

25.2.1 Operators Commute

I claim that all of the plaquette operators and all of the vertex operators commute with each other. It is obvious that

$$[V_\alpha, V_{\alpha'}] = 0$$

since $V_\alpha$’s are only made of $\sigma_z$ operators and all of these commute with each other (either two $\sigma_z$ operators act on different edges, in which case
they commute, or they act on the same edge in which case they are the same operator so they commute). Similarly

$$[P_\beta, P_{\beta'}] = 0$$

since $P_\beta$’s are made only of $\sigma_x$ operators and all of these commute with each other.

The nontrivial statement is that

$$[V_\alpha, P_\beta] = 0 \quad (25.7)$$

for all $\alpha$ and $\beta$. The obvious case is when $V_\alpha$ and $P_\beta$ do not share any edges, then the two operators obviously commute. When they do share edges, geometrically they must share exactly two edges, in which case the commutation between each shared $\sigma_x^{(i)}$ and $\sigma_z^{(i)}$ accumulates a minus sign, and since there are exactly two shared edges the net sign accumulated is $(-1)^2 = +1$ meaning that the two operators commute. An example of this commutation is shown in Fig. 25.4 where a small black arrow points to a particular vertex. Note that the vertex operator has the same eigenvalue both before and after the application of the plaquette operator (eigenvalue = $-1$ since there is an odd number of incoming blue lines). This shows explicitly that the vertex operator at this vertex commutes with the bright pink plaquette operator.

### 25.2.2 Is This a Complete Set of Operators? (Not quite!)

We have $N_x N_y$ vertex operators and $N_x N_y$ plaquette operators — all of these operators commute, and each of these operators has 2 eigenvalues. This appears to match the fact that there are $2N_x N_y$ spins in the system each of which can point up or down, thus apparently giving the same number of degrees of freedom. So is our set of $V_\alpha$ and $P_\beta$ operators a complete set of operators on this Hilbert space? I.e., if we describe the eigenstate of each of these operators do we determine a unique state of the Hilbert space?

It turns out that the $V_\alpha$ and $P_\beta$ operators do not quite form a complete set of operators on the Hilbert space. The reason they fail to form a complete set is that there are two constraints on these operators

$$\prod_\alpha V_\alpha = 1 \quad (25.8)$$
$$\prod_\beta P_\beta = 1 \quad (25.9)$$

To see that Eq. 25.8 is true, note that each edge occurs in exactly two operators $V_\alpha$ (since each edge is attached to exactly two vertices). Thus when we multiply all the $V_\alpha$’s together, each $\sigma_x^{(i)}$ occurs exactly twice, and $(\sigma_x^{(i)})^2 = 1$. Thus the product of all the $V_\alpha$’s is the identity. The argument is precisely the same for multiplying together all of the $P_\beta$’s.
Introducing the Toric Code

Thus we can freely specify the eigenvalues ($\pm 1$) of $(N_x N_y - 1)$ operators $V_\alpha$, but the value of the one remaining $V_\alpha$ is then fixed by the values chosen for the other $(N_x N_y - 1)$ of them. Similarly with the $P_\beta$'s, we can only specify $(N_x N_y - 1)$ eigenvalues ($\pm 1$), and then the last eigenvalue is fixed by the value chosen by the other $(N_x N_y - 1)$. So specifying the eigenvalues of these commuting operators specifies only $2(N_x N_y - 1)$ binary choices. Since we started with $2N_x N_y$ spins which can be in two states (up or down) we must still have two binary choices (two $\pm 1$ degrees of freedom) remaining that we have not fixed. These two degrees of freedom (two binary choices) left unspecified are going to be two error protected qubits (our logical qubits) in the toric code scheme for building a quantum error correcting code.

25.3 Building the Code Space

We are now in a position to build our quantum error correcting code. In particular, we want to define our code space — the space of possible allowed states of our system that we use for encoding quantum information. Analogous to the simple classical codes discussed in section 24.2.1 we must have some error checking protocol that continually checks that the system remains in the code space.

We will now state two simple rules that define our code space. We must (as often as possible) check to see that the two rules remain satisfied. If we find that they are not satisfied we know a physical error has occurred, the system has left the code space, and we must then go about trying to correct it.

**Rule 1:** Specify that $V_\alpha = 1$ for every vertex.

This condition guarantees that there are an even number of blue lines (down spins) incident on every vertex. It is easy to see that this can be interpreted as a constraint that all configurations must form closed loops of the blue lines. There can be no ends of lines, and no branching of lines. An example of such a loop configuration is shown in Fig. 25.6.

Thus our error checking protocol examines every vertex frequently and if it ever finds that $V_\alpha = -1$ then we know we are no longer in the code space, i.e., an error has occurred that we must try to repair.

**Rule 2:** Specify that $P_\beta = 1$ for every plaquette.

As mentioned near Fig. 25.5 this assures that every plaquette is in an equal superposition of flipped and unflipped states with a plus sign between the two pieces. Note in particular that, because the $P_\beta$ and $V_\alpha$ operators commute, the action of flipping a plaquette will not ruin the fact that Rule 1 is satisfied (that is, that we are in a loop configuration).

The quantities $V_\alpha$ and $P_\beta$ are known as the stabilizers of the code — the code space is unchanged under the application of these operators (See also the discussion of stabilizers and syndromes in section 24.4.1).
If these operators are measured to have a $-1$ eigenvalue then we know an error has occurred.

We now have the following prescription for constructing a wavefunction that satisfies both Rule 1 and Rule 2, i.e., a wavefunction in the code space: First start in any state of which satisfies Rule 1, i.e., some loop configuration (this is a state with well defined spins up and down in the $\sigma_z$ basis). We call this configuration the reference configuration. Then to this reference configuration wavefunction, we add all other wavefunctions that can be obtained by flipping plaquettes. We thus have

$$|\psi\rangle = \sum_{\text{all loop configs that can be obtained by flipping plaquettes from a reference loop config}} N^{-1/2} |\text{loop config}\rangle \quad (25.10)$$

where $N$ is a normalization constant that counts the total number of terms in the sum. By adding up all such flipped configurations, we assure that every plaquette is in the correct superposition of flipped and unflipped and we satisfy Rule 2 (Recall from Fig. 25.5 that adding flipped and unflipped gives $P_\beta = +1$).

We now make a crucial observation: flipping plaquettes never changes the parity (eveness or oddness) of the number of loops that go around a nontrivial cycle of the torus. To see this, try drawing a line around a cycle of the torus, as shown with the dashed red line in Fig. 25.7. If one flips a plaquette (bright pink in the figure), it does not change the parity of the number of blue edges that cross through the dashed red line. As a result of this observation, we realize that the sum in Eq. 25.10 does not include all possible loop configurations, but rather contains all loop configurations with fixed parity of loops going around the cycles (matching the parity of loops around the cycles in the reference configuration).

Thus there are four independent wavefunctions of the form of Eq. 25.10 which differ in whether the reference configuration has an even or an odd number of blue edges going around each cycle. All of these states satisfy the constraint rules that all $V_\alpha = 1$ and all $P_\beta = 1$. We will call these states

$$|\psi_{eo}\rangle, \quad |\psi_{oe}\rangle, \quad |\psi_{ee}\rangle, \quad |\psi_{oo}\rangle \quad (25.11)$$

where $e$ and $o$ stand for an even or an odd number of blue lines going around a given cycle. So for example, we have

$$|\psi_{eo}\rangle = \sum_{\text{all loop configs that have an even number of blue strings around vertical cycle and odd number of blue strings going around horizontal cycle}} N^{-1/2} |\text{loop config}\rangle$$

Or graphically, we have Fig. 25.8.
Introducing the Toric Code

Fig. 25.7 A loop configuration (blue edges) on a torus (dotted orange lines on opposite sides are identified as in Fig. 25.2). Drawing a line (dashed red) around one of the cycles of torus, one can see that flipping a plaquette, such as the one marked in bright pink, does not change the parity of the number of blue edges cutting through the dashed red line. Further, it does not matter where we place the dashed red line. So for example, we can change the $y$-coordinate of the horizontal dashed red line, and the number of blue edges it cuts through is always odd. Similarly if we change the $x$-coordinate of the vertical dashed red line, the number of blue edges it cuts through is always even.

$$|\psi_{ee}\rangle = |\psi\rangle + |\psi\rangle + |\psi\rangle + \ldots$$

$$|\psi_{eo}\rangle = |\psi\rangle + |\psi\rangle + |\psi\rangle + \ldots$$

Fig. 25.8 Top Line: Graphical depiction of $|\psi_{ee}\rangle$ which is a sum of all wavefunctions having an even number of blue strings running around each cycle. Bottom line: Graphical depiction of $|\psi_{eo}\rangle$ which has an even number of strings around one cycle (the meridian, short direction, or vertical in the planar diagram) and an odd number around the other (the longitude, long direction, or horizontal in the planar diagram).

The code space is spanned by these four wavefunctions. Equivalently, the most general wavefunction we can write within the code-space (i.e., that that satisfies the two above rules, that all $V_{\alpha} = 1$ and all $P_{\beta} = 1$) is of the form

$$|\psi\rangle = A_{ee}|\psi_{ee}\rangle + A_{eo}|\psi_{eo}\rangle + A_{oe}|\psi_{oe}\rangle + A_{oo}|\psi_{oo}\rangle \quad (25.12)$$

for arbitrary complex coefficients $A_{ee}, A_{eo}, A_{oe}, A_{oo}$ subject to the normalization condition $|A_{ee}|^2 + |A_{eo}|^2 + |A_{oe}|^2 + |A_{oo}|^2 = 1$. It is these coefficients which store the two so-called “logical” qubits of quantum information that we are trying to protect with this coding scheme (compare Eq. 24.1). The underlying spins on the lattice that make up the code are the “physical” qubits. Wavefunctions of the form of Eq. 25.12 are thus the code-space.

Note that in order to turn the $|\psi_{ee}\rangle$ wavefunction into the $|\psi_{eo}\rangle$ wave-
function we would need to insert a single blue loop around the horizontal (longitude) cycle, which involves flipping an entire row of spins at once. If one were to try to flip only some of these spins, we would have an incomplete loop, or an endpoint of a blue line, violating the rule that \( V_\alpha = 1 \) for all vertex sites, therefore not being in the code-space. It is this fact that allows us to test for errors (by looking for such endpoints) and correct them efficiently, as we shall see next.

25.4 Errors and Error Correction

Let us now turn to study possible errors in more detail. What does a physical error look like in this system? Imagine an evil demon arrives\(^4\) and, unbeknownst to us, applies an operator to one of the physical spins in the system to try to create an error. We will start by considering the cases where this operator is \( \sigma_x \) (section 25.4.1) or \( \sigma_z \) (section 25.4.2). We will consider combinations of these operators in section 25.4.3 and argue that we do not have to consider other possibilities in section ??.

25.4.1 \( \sigma_x \) Errors

Let us first consider the case where the error operator is \( \sigma_x \). I.e., starting in one of the code space wavefunctions, \( \sigma_x \) is applied on edge \( i \) to create an error. This operator commutes with all the plaquette operators \( P_\beta \) but anticommutes with the vertex operators \( V_\alpha \) which intersect that edge. This means, if we start in the code space (all \( V_\alpha = +1 \), all \( P_\beta = +1 \)), and apply this error operator \( \sigma_x^{(i)} \), we then end up in a situation where the two vertices attached to the edge \( i \) are now in the wrong eigenstate \( V_\alpha = -1 \). One way to see this is to realize that before \( \sigma_x \) is applied, all of the vertices have an even number of blue edges (down spins) coming into them. By flipping over one edge with \( \sigma_x \) we obtain two vertices with an odd number of blue edges coming into them (one at each end of this edge). A more formal way to see this is to realize that in the original state \( |\psi\rangle \) we have

\[
V_\alpha |\psi\rangle = |\psi\rangle
\]

meaning we start in the +1 eigenstate. We then apply the error operator \( \sigma_x^{(i)} \) to both sides. Assuming \( i \) is one of the edges incident on the vertex \( \alpha \), we have

\[
\sigma_x^{(i)} |\psi\rangle = \sigma_x^{(i)} V_\alpha |\psi\rangle = -V_\alpha \sigma_x^{(i)} |\psi\rangle
\]

or

\[
V_\alpha [\sigma_x^{(i)} |\psi\rangle] = -[\sigma_x^{(i)} |\psi\rangle]
\]

showing we end up in the \(-1\) eigenstate of the vertex operator.

To show these errors graphically we will no longer draw the up and down spins (the blue edges) but instead we just draw the \( \sigma_x \) operator as a dark red line, and the vertices which are in the \(-1\) eigenstate as an orange \( \times \) as shown in Fig. 25.9.

\[\text{Fig. 25.9 Starting in the code space, a } \sigma_x \text{ operator applied to the red edge (i) creates two vertices in the } V_\alpha = -1 \text{ eigenstate marked with the orange } \times \text{'s.}\]

\(^4\) It is not entirely necessary to anthropomorphize the error creating process.
Introducing the Toric Code

Fig. 25.10 Left: When two $\sigma_x$ errors (the red edges) are made on edges that share a vertex (the small black arrow), the shared vertex is hit with $\sigma_x$ twice, and thus returns to the $V_\alpha = +1$ state. Only the two vertices at the end of the “string” are in the $V_\alpha = -1$ state. Middle: A longer string of errors. Note that we can only measure the endpoints of the string, not where the errors were made, so we cannot tell if the error string goes down two steps then two steps to the right, or if goes two steps to the right then down two steps. Right: If we detect the errors as in the middle panel and we try to correct it by dragging the errors back together, but we choose the incorrect path for the string, we end up making a closed (red) loop of $\sigma_x$ operators — which acts as the identity on the code space, so we still successfully correct the error!

So it is clear what our error correction protocol must do. It must frequently measure the state of the $V_\alpha$ operators. If it finds all $V_\alpha$ operators in the +1 eigenstate then the system is still in the code space. On the other hand, if it finds a pair of adjacent vertices in the in the $V_\alpha = -1$ state, we know that a $\sigma_x$ operator has been applied on the intervening edge by some error creating process. Once we have identified the error it is easy to correct it by applying $\sigma_x$ on the same edge, thus returning the system to its original state and to the code space.

Now suppose that the error creating demon is very fast and manages to make several such $\sigma_x$ errors very quickly. If these errors are well separated from each other, we will easily find multiple pairs of vertices in the $V = -1$ state, with the two members of each pair separated from each other by one edge distance. These can similarly be identified by our correction scheme and repaired, returning us to the code space again.

However, it could be the case that two errors are created on two edges that share a single vertex, as shown on the left of Fig. 25.10. The vertex that is shared (marked by a small black arrow in the figure) gets hit by $\sigma_x$ twice. The first time it is hit by $\sigma_x$ the eigenvalue of $V_\alpha$ flips from its initial state +1 to -1 but then when it is hit the second time, this flips it back to the +1 state. Thus after the two $\sigma_x$ operators have been applied on the two red edges on the left of Fig. 25.10 only the two vertices, marked with an orange $\times$, at the end of the red “string” are in the $V_\alpha = -1$ state and are then detectable as errors.

Nonetheless, the error correction scheme is still fairly straightforward. One checks the state of all the vertices and when $V_\alpha = -1$ is found, one tries to find the closest other vertex with $V_\alpha = -1$ to pair it with. Then one applies a string of $\sigma_x$ operators to correct these errors. One can think of this as dragging the errors back together and annihilating them with each other again.

It is important to realize that we cannot see the error operators themselves (which we have drawn as a red string in Fig. 25.10) by making measurements on the system — we can only detect the endpoints of string, the vertices, marked with an orange $\times$, where $V_\alpha = -1$. For
example, in the middle panel of Fig. 25.10 we cannot tell if the error string goes down two steps and then to the right, or if it goes to the right two steps then down. We only know where the endpoints of the string are.

Suppose now we detect the two errors shown in the middle panel of Fig. 25.10. We may try to correct these errors by guessing where the red string is, and applying $\sigma_x$ along this path to bring the endpoints back together and reannihilate them. However, it is possible that we guess incorrectly as shown in the right panel of Fig. 25.10. In this case we will have ended up producing a (red) closed loop of $\sigma_x$ operators applied to the original state. Fortunately, a product of $\sigma_x$ operators around a closed loop is precisely equal to the product of the plaquette operators $P_\beta$ enclosed in the loop

$$\prod_{i \text{ around loop}} \sigma^{(i)}_x = \prod_{\beta \text{ enclosed in loop}} P_\beta$$

(25.13)

Since the code space is defined such that all of the plaquettes operators are in the +1 eigenstate, this red loop of $\sigma_x$ thus acts as the identity on the code space, and we still successfully correct the error!

If it is not already obvious, the reason for the equality Eq. ?? can be seen in the right of Fig. 25.10. Within the enclosed region, each edge is acted on by two plaquette operators (marked in bright pink) each applying $\sigma_x$ to the edge, and since $\sigma_x^2 = 1$ this means there is no net effect on the internal edges, and one is left with only $\sigma_x$ applied to the (red) loop bounding the region.

On the other hand, if a loop of errors occurs which extends around a cycle, as shown in Fig. 25.11 (think of this as dragging the error, the orange ×, all the way around the cycle and re-annihilating it again) then, although we return to the code space (there are no $V_\alpha = -1$ vertices remaining) we have changed the parity of the number of down spins around a cycle thus scrambling the quantum information and make an error in the logical bits. In fact what we get in this case is a transform that switches the even and odd sectors around one cycle:

$$A_{cc}|\psi_{cc}\rangle + A_{co}|\psi_{co}\rangle + A_{ac}|\psi_{ac}\rangle + A_{oo}|\psi_{oo}\rangle \longrightarrow$$

$$A_{ac}|\psi_{cc}\rangle + A_{oa}|\psi_{oa}\rangle + A_{cc}|\psi_{oe}\rangle + A_{oa}|\psi_{oo}\rangle$$

(25.14)

If the red error loop had gone around the other cycle, the second index of $A_{ij}$ would have flipped rather than the first index.

The powerful idea of the toric code is that by having a very large torus, it requires a very large number of physical errors (spin flips) to make this loop go all the way around the cycle and actually scramble the quantum information (the logical qubits). If we are continuously checking for $V_\alpha = -1$ physical errors we can presumably correct these errors before a logical error can arise.

**Fig. 25.11** If an error string (red) goes all the way around a cycle, it changes the parity of the number of blue loops running around the cycle. Although this wavefunction is in the code space, it has been scrambled compared to the original wavefunction with no error string. I.e., a logical error has occurred.
25.4.2 $\sigma_z$ Errors

We can also consider what happens if the error operator applied to the system is a $\sigma_z$ operator. Much of the argument in this case is similar to the $\sigma_x$ case.

The $\sigma_z$ operator on an edge anticommutes with the two adjacent plaquettes $P_\beta$ which each include that edge. Applying $\sigma_z$ to a system in the code space takes the system out of the code space and results in the two adjacent plaquettes having eigenvalue $P_\beta = -1$ as shown in Fig. 25.12.

To be more explicit about this, recall that in the code space each plaquette is in a superposition (a positive sum) of flipped and unflipped plaquettes (See Fig. 25.5). In Fig. 25.13 we can see how applying a $\sigma_z$ operator to one edge results in the difference of the flipped and unflipped plaquettes, and thus $P_\beta = -1$.

Analogous to the above discussion of $\sigma_x$ errors, our $\sigma_z$ error correction protocol should frequently check for pairs of neighboring plaquettes where $P_\beta = -1$ and if these are found the protocol should correct the error by applying $\sigma_z$ to the intervening edge.

We next consider the case where several $\sigma_z$ errors are created before being corrected. Again if the errors are spatially well separated they can be identified and corrected without regards to each other. Whenever we find pairs of neighboring plaquettes in the $P_\beta = -1$ state, we correct them each individually. However, the situation is more complicated if two such errors occur on different edges of the same plaquette.

Starting in the code space, if two $\sigma_z$ errors are applied on two different edges of the same plaquette (marked in dark green in the left of Fig. 25.14) then the $P_\beta$ eigenvalue of that plaquette (marked with an arrow) is flipped twice. As shown in the left of Fig. 25.14 this results in two plaquettes in the $P_\beta = -1$ state being separated by a plaquette (marked with the arrow) in the $+1$ state. In the figure we have marked the $P_\beta = -1$ plaquettes with bright green $\times$. These $\times$’s are connected together with a bright green line which cuts through the dark green edges. Sometimes we say that the bright green line is a path on the dual lattice (meaning it goes from center of plaquette to center of plaquette).

The error correction procedure uses multiple $\sigma_z$ operators to bring these defects back together and reannihilate them.

If still more $\sigma_z$ errors are created, they can form a string, as shown in the middle of Fig. 25.14. The $\sigma_z$ errors here are applied on dark green edges. The bright green path on the dual lattice cuts through all of these...
25.4 Errors and Error Correction

Fig. 25.14 Left: When two $\sigma_z$ operators (dark green) are applied to two different edges of the same plaquette (the one with the arrow) then this plaquette ends up in the $P_\beta = +1$ state. The two plaquettes with $P_\beta = -1$ are marked with the bright green $\times$ and are connected together by the bright green error string. Middle: A string of several $\sigma_z$ errors. The bright green line cuts through each of the dark green edges where $\sigma_z$ is applied. The bright green line lives on the dual lattice, meaning it goes from plaquette center to plaquette center. Right: A closed loop of $\sigma_z$ errors. This is equal to the product of all of the enclosed $V_\alpha$ operators. In the code space, this product is equal to $+1$.

dark green edges. As in the $\sigma_x$ case, one is not able to actually detect the bright green string (or the dark green edges), but can only see the endpoints of the string as plaquettes where $P = -1$.

Analogous to the $\sigma_x$ case, if from errors, or from an attempt to correct errors, the $\sigma_z$ error string forms a closed loop as illustrated as the bright green line in the right of Fig. 25.14 this loop of $\sigma_z$ operators is equal to the product of the enclosed $V_\alpha$ operators.

$$\prod_{i \text{ around dual loop}} \sigma_z^{(i)} = \prod_{\alpha \text{ enclosed in dual loop}} V_\alpha \quad (25.15)$$

Since within the code space, $V_\alpha = 1$, a closed loop returns the system to its original state. This means that reannihilating the defects (the bright green $\times$’s) successfully correct the errors independent of the path that we use to bring the defects back together.

Another way of seeing Eq. 25.15 is to think in terms of the blue loops of down spins discussed above. The $\sigma_z$ operators register $-1$ each time they intersect a blue loop. On the other hand the blue loops must be closed so the number of intersections between a blue loop and the loop of the blue $\sigma^z$ error string in the figure must be even (since a blue loop going into the region enclosed by the bright green string must also come out), thus forcing the product of the blue $\sigma_z$ operators to have a value of 1.

On the other hand, if the loop of $\sigma_z$ operators goes all the way around the cycle, it then scrambles the logical qubits. In particular, one can see that if there is a string of $\sigma_z$ going all the way around one cycle as shown in Fig. 25.14 this operator then counts the parity of the number of blue edges going around the opposite cycle, as shown in the figure. Thus, applying the string of $\sigma_z$ operators around the (horizontal) cycle makes the transformation

$$A_{ce} |\psi_{ce}\rangle + A_{eo} |\psi_{eo}\rangle + A_{oc} |\psi_{oe}\rangle + A_{oo} |\psi_{oo}\rangle \rightarrow \quad (25.16)$$

If the string of $\sigma_z$ were to go around the opposite (vertical) cycle, the
Introducing the Toric Code

**Fig. 25.16** Left: Both $\sigma_x$ and $\sigma_z$ are applied on the same edge. This results in two neighboring vertex defects (orange $\times$’s) with $V_\alpha = -1$ and two neighboring plaquette defects (bright green $\times$’s) with $P_\beta = -1$. Middle: Additional $\sigma_x$ operators are applied along the red string to move the vertex defects apart, and an additional $\sigma_z$ operator is applied on the dark green edge to move the plaquette defects apart. Right: A closed (red) loop of $\sigma_x$ operators is equal to the product of all of the enclosed $P_\beta$ operators. As compared to Fig. 25.10, here there is an enclosed plaquette defect (with the bright green $\times$) so the value of the loop is $-1$.

minus sign would be applied instead to $A_{eo}$ and $A_{oo}$.

### 25.4.3 Combinations of $\sigma_x$ and $\sigma_z$

We have discussed errors created by $\sigma_x$ and $\sigma_z$, but we should also consider what happens when both types of errors are created. If we have an error correction protocol that removes $\sigma_x$ errors and another protocol that removes $\sigma_z$ errors, and as long as these two procedures don’t interfere with each other, we should be able to remove combinations of the two.

To give a detailed example let us suppose, starting in the code space, both a $\sigma_x$ operator and a $\sigma_z$ operators are applied to a single edge as shown in the left of Fig. 25.16. We obtain from these error operators both the orange $\times$’s at the two adjacent vertices (two vertices with $V_\alpha = -1$, from the application of $\sigma_x$) and the bright green $\times$’s on the two adjacent plaquettes (two plaquettes with $P_\beta = -1$ from the application of $\sigma_z$). Assuming our error checking procedure finds all of these errors, it can correct the errors by applying both a $\sigma_x$ and a $\sigma_z$ to this edge again.

However, we might want to be a bit more cautious if we are to consider both $\sigma_x$ errors (vertices with $V_\alpha = -1$) and $\sigma_z$ errors (plaquettes with $P_\beta = -1$) at the same time. For example, in the middle of Fig. 25.16 we imagine $\sigma_z$ is applied to the two edges marked in dark green, thus making $P_\beta = -1$ for the two plaquettes marked with bright green $\times$’s, and $\sigma_x$ is applied to the six edges marked in red so that $V_\alpha = -1$ at the two orange $\times$’s. On the right of the figure we imagine more $\sigma_z$ operators applied so that the orange $\times$’s come together and reannihilate forming a closed red loop. As in the right of Fig. 25.10 the closed red loop is equal to the product of $P_\beta$ for the enclosed plaquettes (Eq. 25.13). However, in this case one of the enclosed plaquettes (the one marked with the green $\times$) is in the $P_\beta = -1$ eigenstate. Thus forming the closed red loop now gives an overall sign of $-1$. Similarly if we had moved the bright green $\times$ in a loop around one of the orange $\times$’s and reannihilated it, as shown in Fig. 25.17 we would also have generated a $-1$.

We have thus uncovered an important principle: If you move a defect
around in a loop and reannihilate it, you make a measurement of what is contained in that loop!

From the context of our code, should we be worried about having accumulated an overall sign? No! As mentioned in chapter 11 note 8 typically in quantum computing we are not interested in the overall phase of a result, which is hard — if not impossible — to control anyway. The logical qubits are protected up to an overall phase prefactor only.

25.4.4 Other Errors

We have now considered errors created by the application of $\sigma_x$ operators and $\sigma_z$ operators. Further we argued that we are not concerned about the overall phase of the system, so application of a constant prefactor is not something we worry about. In fact, products and sums of all combinations of $\sigma_x$, $\sigma_z$ and constants can generate all possible error operators that could ever be applied to our system! Thus our error correction scheme can in principle correct all possible errors!

25.5 The Toric Code on Different Lattices, and Different Topologies

Before ending this chapter it is worth making a short detour to consider the possibility of building the toric code with a lattice of a different geometry. In building the toric code we could have used a triangular lattice, a honeycomb$^6$, or even an “irregular lattice”$^6$. Whatever the geometry of the lattice (regular or irregular) the vertex term must assure an even number of blue edges into each vertex (no matter how many edges join at each vertex), and the plaquette term must flip all of the edges of a plaquette (no matter how many edges the plaquette has). An example of an irregular lattice is shown in Fig. 25.18.

6 Which should not be called a lattice, although people insist on incorrectly calling it a lattice anyway! See Simon [2013], for example, for proper definition of a lattice!

![Fig. 25.18 Part of an irregular lattice. A vertex operator with three legs is marked in green. This operator still gives $+1$ or $-1$ if an even or odd number of down spins are incident on the vertex. A plaquette operator with five sides is marked in bright pink. The plaquette operators flip all of the spins along its edges.](image)

For any lattice geometry (regular or irregular) the vertex and plaque-
Introducing the Toric Code

Tetrakise Operators are a commuting set. Thus the code space can be described as (Eq. 25.10) a sum of all loop configurations that can be reached from a reference configuration by flipping plaquettes. What is crucial to the toric code is that the topology of the system is a torus, so that we still have four orthogonal states in the code space characterized by the number of (blue) loops around the two cycles as shown in Fig. 25.8.

One we allow different lattice geometries, we may also ask the question of what happens if our system has different topologies, i.e., is not a torus, but is some general more general object.

To understand the dependence on the topology of the surface let us consider an arbitrary two-dimensional orientable closed surface\(^7\). In two dimensions, all such surfaces can be fully described topologically by their genus \(g\), or number of handles: a sphere, \(g = 0\), has zero handles; a torus, \(g = 1\), has one handle; a two-handled torus, \(g = 2\), has two handles, and so forth. We will make use of the famous Euler characteristic: for any polyhedral decomposition of an orientable two manifold, we have the beautiful formula\(^8\)

\[
2 - 2g = \text{(Number of Vertices)} - \text{(Number of Edges)} + \text{(Number of Faces)}
\]

An example of this identity is shown in Fig. 25.19. In section 25.2.2 we counted the total number of degrees of freedom in the system and we found that, after fixing all of the vertex and plaquette degrees of freedom, on a torus, we were left with two qubits remaining, and these became the protected logical qubits of our toric code. Let us now try to figure out, for a more general surface, how many degrees of freedom we cannot specify once we have specified all the vertex and plaquette eigenvalues. Since there is one spin on each edge we can rewrite Euler’s equation as

\[
2 - 2g = \text{(Number of Vertices)} - \text{(Number of Edges)} + \text{(Number of Faces)}
\]

\[
\left[ \text{Number of Vertex Ops} + \text{Number of Plaquette Ops} - 2 \right] + 2g = \text{Number of Spins} \tag{25.17}
\]

We can read this equation as follows. The right hand side is the total number of binary (±1 or ↑, ↓) choices we can make (number of 2-state degrees of freedom). The brackets counts the number of these choices we specify by fixing all of the vertex and plaquette eigenvalues: we have one choice for each vertex operator and one for each plaquette operator, but then we subtract two binary choices because of the constraints Eqs. 25.8 and 25.9 (Once all but one vertex is specified, the last vertex is fixed and is not an additional degree of freedom, and similar for plaquettes). The remaining term, \(2g\), is the number of binary choices (number of qubits) that we cannot fix by specifying the vertex and plaquette operator eigenvalues. Thus, in the case of a spherical surface, there are no qubits remaining unspecified; in the case of a torus \((g = 1)\) there are two qubits left (as we calculated above); in the case of a two handled torus, \(g = 2\), there are four qubits, and so forth.

We might have suspected that the number of qubits we could make

\(^7\) In fact the toric code can be defined on a non-orientable surface such as a Klein bottle. However, since the rest of this book focuses on orientable manifolds, we will do the same here for simplicity.

\(^8\) Euler noticed this in 1758 for the case of convex polyhedra \((g = 0)\). However, his proof of the statement was incorrect. It was correctly proven by Legendre in 1794.
out of a toric code on a $g$-handled torus would be related to the number of different distinguishable loops on the surface. Indeed this turns out to be correct! For a $g = 1$ genus (regular) torus, there are two independent cycles (the longitude and the meridian). Similarly on a $g$-handled torus there are $2g$ independent cycles. Each qubit represents the evenness or oddness of the number of (blue) loops going around each cycle.

Shor’s original nine-bit error correcting code (Section ??) can also be viewed as a toric code — but on a surface with singular points (See exercise 25.3).

25.6 $\mathbb{Z}_N$ Toric Code (Briefly)

One of the simplest generalizations of the toric code is known as the $\mathbb{Z}_N$ toric code. (While this is not too complicated a generalization, it is also not entirely crucial to the development of ideas here and can be skipped on a first reading\textsuperscript{10}). In this model instead of having a two state system (spin-up/spin-down) on each edge, we will put an $N$ state system\textsuperscript{11} on each edge which we will write as $s \in 0, \ldots, (N - 1)$, instead of saying a spin is up or down (or the edge is blue or black). It is worth noting that it will sometimes be more natural to think of the exponential $e^{2\pi i s/N}$ which naturally has only $N$ possible states.

We can use a regular or irregular lattice for this model. However, now for $N > 2$ we must put arrows on each edge to define a direction. The direction chosen does not particularly matter. An irregular lattice with arrows is shown in Fig. 25.20 Each edge $i$ is labeled with a quantum number $s_i \in 0, \ldots, N$. (The $N = 2$ case corresponds to the regular toric code with $s = 0$ corresponding to a black edge [spin up] and $s = 1$ corresponding to a blue edge [spin down]). We will think of this quantum number as being some sort of fictitious $\mathbb{Z}_N$ valued current running along the edge in the direction of the arrow.

We now define an operator $Q_\alpha$ that measures the total $\mathbb{Z}_N$ current sink or source at a vertex $\alpha$ to be\textsuperscript{12}

$$Q_\alpha = \left[ \sum_{i \in \text{edges arriving at } \alpha} s_i - \sum_{i \in \text{edges leaving } \alpha} s_i \right] \mod(N) \quad (25.18)$$

where an edge touching a vertex is denoted as "arriving" at a vertex if its arrow is pointed at the vertex and is denoted as "leaving" if its arrow is pointed away from the vertex. This operator has $N$ possible eigenvalues given by the possible charges $0, \ldots, (N - 1)$. The sum of charges over the entire system must be zero

$$\sum_\alpha Q_\alpha \mod(N) = 0$$

since this expression counts each edge as being a leaving vertex once and counts each edge as being an arriving vertex once. In a form more

\textsuperscript{9}The number of independent cycles (or number of holes) in a manifold is known as the first Betti number. For a $g$-handled torus, this number is $2g$. More precisely the first Betti number is equal to the number of closed circular cuts required to reduce the manifold to a simply connected region. Another way to define the first Betti number is as the number of generators in the so-called first homology group, for those who know homology theory.

\textsuperscript{10}Our presentation will be a bit faster here with the assumption that the reader is now familiar with the conventional toric code.

\textsuperscript{11}A two state system is known as a qubit. A three state system is sometimes known as a qutrit. A general $d$ state system is known as a $d$ qudit. One should not call an $n$ state system "qunit" though since the word "nits" is often used for an object having $e$ states, with $\ln e = 1$. (Also "nits" are lice eggs. Yuck.)

\textsuperscript{12}The term $\mod(N)$ means take the remainder after dividing by $N$. Because we are working $\mod(N)$ it is not possible to say whether a vertex is a current sink or a current source since a value $Q_\alpha = m$ is the same as $Q_\alpha = m + pN$ for any integer $p$, positive or negative.
Introducing the Toric Code

analogous to the constraint Eq. (25.8) we could instead write

$$\prod_{\alpha} e^{2\pi i Q_{\alpha} / N} = 1$$

We can also define another operator which changes the value of the variables on the edges of a plaquette $\beta$

$$A_\beta = \begin{cases} s_i \to (s_i + 1) \text{mod}(N) & \text{arrow pointing clockwise around } \beta \\ s_i \to (s_i - 1) \text{mod}(N) & \text{arrow pointing counterclockwise around } \beta \end{cases}$$

(25.19)

Since $(A_\beta)^N$ is the identity, there are $N$ possible eigenvalues of this operator which are given by $e^{2\pi ik/N}$ with $k = 0, \ldots, (N - 1)$.

Each edge with a fixed arrow direction bounds two plaquettes such that the arrow points clockwise with respect to one plaquette and counterclockwise with respect to the other plaquette as shown for example in Fig. 25.21. Applying $A_\beta$ to the plaquettes on both sides of an edge thus leaves the edge separating the two unchanged. This immediately implies that applying $A_\beta$ to all plaquettes in the system is the identity:

$$\prod_\beta A_\beta = 1$$

analogous to Eq. (25.9)

A useful commutation relation (analogous to $\sigma_x \sigma_z = -\sigma_z \sigma_x$ for the conventional toric code) is (See exercise 25.1).

$$e^{2\pi i s_j / N} A_\beta = e^{\pm 2\pi i / N} A_\beta e^{2\pi i s_j / N}$$

(25.20)

where $s_j$ is the operator that measures the quantum number on an edge $j$ on the boundary of plaquette $\beta$. The $\pm$ on the right is positive if the arrow on the edge is going clockwise around $\beta$ and is minus if the arrow on the edge is going counterclockwise.

25.6.1 Code Space

Analogous to the discussion of section 25.2.1, it is easy to establish that the operators $A_\beta$ and $Q_{\alpha}$ all commute with each other (See exercise 25.1).

Our code space for this system will be the space of states such that we have the vertex condition $Q_{\alpha} = 0$ (or to look more similar to the regular toric code, all $e^{2\pi i Q_{\alpha} / N} = 1$) for all vertices and the plaquette condition $A_\beta = 1$ for all plaquettes.

Let us first examine the condition $Q_{\alpha} = 0$ for all vertices. This means that there is no net current flow into or out of any vertex analogous to the loop condition for the conventional toric code (See near Fig. 25.8). This is very much like Kirchhoff’s first law in circuit analysis except that the current is $\mathbb{Z}_N$ valued, i.e., its value is periodic modulo $N$. In Fig. 25.22 we show two configurations of edge variables on a torus geometry that satisfy this condition for a $\mathbb{Z}_3$ toric code.
Note that if we start with a situation where all \( Q_\alpha = 0 \), flipping a plaquette by applying \( A_\beta \) (shown in Fig. 25.22) does not change the condition that \( Q_\alpha = 0 \) (This is equivalent to our statement that \( Q_\alpha \) and \( A_\beta \) commute). The second code space condition, \( A_\beta = 1 \), is analogous to the plaquette condition for the usual toric code. The code space wavefunction can then be written in the usual form

\[
|\psi\rangle = \sum_{\text{all configs that can be reached by applying } A_\beta \text{ starting with a reference config that satisfies } Q_\alpha = 0} N^{-1/2} |\text{config}\rangle 
\] (25.21)

We should be a bit cautious if we define our code space as the \( A_\beta = 1 \) state. As discussed in the toric code, we would want to have an error correcting protocol that continually checks to make sure that the system has not left the code space. Hence we might want to make continuous measurements of \( A_\beta \). Unfortunately, \( A_\beta \) is not a Hermitian operator, so it is not directly measurable. However, we can certainly measure \( \frac{1}{2}(A_\beta + A_\beta^\dagger) \) and \( \frac{1}{2i}(A_\beta - A_\beta^\dagger) \) which both have the same eigenvectors as \( A_\beta \) (since \( A_\beta^\dagger = A_\beta^{-1} \)) and this will work just as well.

As with the regular \( (\mathbb{Z}_2) \) toric code, there are multiple orthogonal wavefunctions that one can obtain depending on the reference configuration used. For each noncontractable cycle in the system, we can define a \( \mathbb{Z}_N \) valued current flow going around that cycle. For example, in the torus shown in Fig. 25.22 there are 2 units (mod 3) of current going vertically, and 3 units (mod 3) going horizontally. Note that flipping a plaquette by applying \( A_\beta \) (also shown in the Figure) does not change
Introducing the Toric Code

**Fig. 25.23** Introduction of an error string that creates two vertex defects in an example of the $\mathbb{Z}_3$ toric code. The error string (red) has an arrow pointing from the $Q_\alpha = -1$ defect to the $Q_\alpha = +1$ defect. Each edge along the path is incremented one unit if its arrow is aligned with the red string arrow and is decremented one unit if its arrow is opposed to the red string arrow. Note that on the left, an element of the code space is obtained by applying $A_\beta$ to all possible plaquettes in all possible ways and summing the result. If we start in the code space and apply the red error string, note that the error string is not measurable, only the defects at the end points are.

the value of the current going around either noncontractable cycle. Thus since a genus $g$ surface has $2g$ noncontractable cycles, we expect that the $\mathbb{Z}_N$ toric code should be able to store $2g N$-state systems of quantum information in its quantum memory. I.e, the Hilbert space of this quantum memory has dimension $N^{2g}$.

### 25.6.2 Errors

We can now consider what sort of error processes can occur. The first type is a vertex error (analogous to the $\sigma_x$ error of the conventional toric code). Here we can consider, for example, incrementing the quantum number (the “current" label) along one edge (mod $N$), to create a defective vertex at either end of that edge. One of these vertices will then have $Q_\alpha = +1$ and the other will have $Q_\alpha = -1$.

More generally we can consider moving such defects away from each other analogous to Fig. 25.10. An example of this is illustrated in Fig. 25.23. This generates an error string (shown red in the figure) connecting the two defects. The error string (red) has an arrow on it, and will increment the quantum number of any edge along its path if the arrow of the edge is aligned with that of the string, and will decrement the quantum number of the edge if the arrow of the edge is antialigned with the arrow of the string. At the head of the string is a vertex defect with $Q_\alpha = +1$ and at the tail is a defect with $Q_\alpha = -1$. Defects with $Q_\alpha = \pm m$ can analogously be made by incrementing or decrementing edges by $m$ steps. Multiple strings may be introduced and the defects may be brought together, adding together modulo $N$. Defects are always created in $\pm$ pairs, so that if we start in the code space, the sum of $Q_\alpha$ over the entire system must remain zero (mod $N$).

If we create a defect pair and move the $Q_\alpha = +1$ defect in a clockwise circle then reannihilate it with the $Q_\alpha = -1$ defect, this leaves behind a closed loop of red-string, analogous to the right of Fig. 25.10. However,
25.6 \( Z_N \) Toric Code (Briefly) 347

A\( \beta \) = \( e^{2\pi i/N} \)

A\( \beta \) = \( e^{-2\pi i/N} \)

Fig. 25.24 Application of the operator \( X_j = e^{2\pi i s_j/N} \) on an edge (shown as dark green lines) multiplies the eigenvalue of \( A_\beta \) on the plaquette to the left of the arrow by \( e^{2\pi i/N} \) and multiplies the eigenvalue of the plaquette on the right by \( e^{-2\pi i/N} \). Multiple applications of these operators can move the defective plaquettes away from each other, as shown in the Figure — only the bright green \( \times \)'s are defects. In the figure, the two vertical dark green bonds are \( X_j \) operators and the horizontal dark green bond is a \( X_j^{-1} \) operator.

it is easy to see that such a clockwise oriented closed loop is equal to the product of \( A_\beta \) operators for the enclosed plaquettes.

\[
\text{Moving } (Q_\alpha = +1) \text{ defect in clockwise loop } = \prod_{\beta \text{ enclosed}} A_\beta \quad (25.22)
\]

analogous to Eq. 25.13 Moving in a counterclockwise loop will correspondingly give a product of \( A_\beta^{-1} \). Analogous to the conventional toric code, since in the code space the \( A_\beta \)'s are all in the +1 eigenstate, this means that starting in the code space, creating two defects dragging, moving them around and reannihilating returns the system to the original state in the code space. However, dragging a defect around a noncontractable cycle (around a handle of the torus for example), and then reannihilating puts the system in a different state of the code space and creates a logical error for this quantum memory.

We now turn to consider plaquette defects. Here we imagine starting in the code space and applying the operator \( X_j = e^{2\pi i s_j/N} \) on edge \( j \) (where \( s_j \) is the operator that measures the quantum number on the edge). Due to the commutation relation Eq. 25.20 the action of the operator \( X_j \) changes the eigenvalue of \( A_\beta \) on the neighboring two plaquettes multiplying the eigenvalue on the neighboring plaquette to the left of the arrow on the edge \( j \) by \( e^{2\pi i/N} \) and multiplying the other neighbor by \( e^{-2\pi i/N} \). Correspondingly using the operator \( X_j^{-1} \) will multiply the eigenvalue of \( A_\beta \) by the inverse phase. Starting in the code space (where all \( A_\beta \) are in the +1 eigenstate) application of \( X_j \) or \( X_j^{-1} \) will create a pair of neighboring defective plaquettes. Applying multiple such operators can move the defective plaquettes away from each other as shown in Fig. 25.24 (analogous to Fig. 25.14 for the conventional \( N = 2 \) toric code).

If, starting in the code space, a \( A_\beta = e^{2\pi i/N} \) defect is created and
then moved in a clockwise loop and then reannihilated with its partner, the system is returned to the code space. In the process a net phase is accumulated given by

\[ \text{Moving } (V_\beta = e^{2\pi i/N}) \text{ defect in clockwise loop } = \prod_{\alpha \text{ enclosed}} e^{2\pi iQ_\alpha/N} \quad (25.23) \]

To see this result, it is easiest to imagine a situation where all the arrows on the (dark green in Fig. 25.24) edges point towards the center of the loop. Moving the \( A_\beta = e^{2\pi i/N} \) defect in a clockwise loop then corresponds to applying \( X_j \) operators on all the edges, thus these operators measure the sum of all the edges entering the enclosed region which is thus the sum of all the \( Q_\alpha \) inside the region (See Eq. 25.18). (One can then check that reversing the orientation of the arrow on an edge does not change this result.) As in the case of the conventional toric code, if a plaquette defect is moved around a noncontractable cycle, a logical error is imparted to the code space.

**Further Reading**

Look at Kitaev? (Laumann?)

---

**Exercises**

**Exercise 25.1 Commutation of operators in the \( \mathbb{Z}_N \) toric code**

(a) Show that all of the operators \( A_\beta \) (Eq. 25.19) and \( Q_\alpha \) (Eq. 25.18) commute.

(b) Prove Eq. 25.20

**Exercise 25.2 Code Space Degeneracy of the \( \mathbb{Z}_N \) Toric Code**

Generalize the Euler characteristic argument of section 25.5 to confirm that the dimension of the code space of the \( \mathbb{Z}_N \) toric code is \( N^{2g} \) where \( g \) is the genus of the surface.

**Exercise 25.3 Shor’s Code is a Toric Code!**

Consider 9 edges assembled into three spheres in the following way. First make a single sphere out of two vertices (put one at the north pole and one at the south pole for simplicity), three edges connecting the two vertices, and three plaquettes filling out the sphere. Now make three spheres in this way. Connect these spheres together at the vertices but do not add any new plaquettes or edges. We now have three vertices, each connected to six edges (three on one sphere and three on another sphere). This makes a manifold with three singular points as shown in Fig. 25.25. Consider now the toric code applied to this system. There is one noncontractable loop in the figure which plays the role of a qubit. There are nine plaquette operators and three vertex operators, making a total of 12 stabilizers. Show that this system is precisely the Shor error correcting code introduced in section ??.

---

*Fig. 25.25 Three spheres (shown elongated into bananas) connected together at their north and south poles. Each sphere consists of three edges, two vertices, and three plaquettes.*
The Toric Code as a Phase of Matter and a TQFT

We have introduced the toric code as a way to store quantum information. However, it is also possible to construct the toric code such that it is a quantum phase of matter, i.e., the ground state of Hamiltonian. As such the toric code becomes an example of topologically ordered quantum matter — a physical system that is described at low temperature and long wavelength by a topological quantum field theory.

To recast the toric code as a phase of matter, we simply write a Hamiltonian which is a sum of all of our vertex and plaquette operators described in section 25.2

\[ H_{\text{toric code}} = -\frac{\Delta_v}{2} \sum_{\text{vertices } \alpha} V_\alpha - \frac{\Delta_p}{2} \sum_{\text{plaquettes } \beta} P_\beta \]  

(26.1)

where here \( \Delta_v > 0 \) and \( \Delta_p > 0 \) have dimensions of energy and set the energy scale of the problem. The operators \( V_\alpha \) and \( P_\beta \) all have eigenvalues \( \pm 1 \), and since the operators all commute with each other (see section 25.2.1) the lowest energy configuration (i.e., the ground state space) is obtained by simply setting all of the \( V_\alpha = 1 \) and \( P_\beta = 1 \).

In other words, the ground state space is exactly the code space!

If the system is on a torus geometry, there will be a four-fold degenerate ground state corresponding to the four orthogonal wavefunctions in the code space (Eqn. 25.11). As discussed in section 25.5 if the system is on a genus \( g \) surface, we would instead have a \( 2^{2g} \)-fold degenerate ground state (corresponding to the \( 2g \) qubits in the code space). The dependence on topology strongly suggests to us that our physical system is described by a topological quantum field theory!\(^1\)

26.1 Excitations

If there are vertices with \( V_\alpha = -1 \) or plaquettes where \( P_\beta = -1 \), the system is not in the ground state space. These occurrences, which we called errors previously, in the language of topologically ordered matter should now be considered to be particle (or quasiparticle\(^5\)) excitations.

Let us list all of the types of particles we can find

1. We always have a vacuum or identity particle (which can also be thought of as the absence of a particle) which we call \( I \).
2. We can have a vertex where \( V_\alpha = -1 \) instead of \( V_\alpha = +1 \). The
3. Often for simplicity one sets \( \Delta_v = \Delta_p = \Delta \). To simplify even more, set the energy scale of the problem to unity with \( \Delta_v = \Delta_p = 1 \).
4. Since \( V_\alpha \) and \( P_\beta \) have eigenvalues \( \pm 1 \), we have included a factor of \( 1/2 \) out front so that the difference between energies of these two eigenstates is \( \Delta_v \) or \( \Delta_p \) respectively.
5. The term “quasiparticle” is used for objects that act as relatively weakly interacting individual particles in low energy theories but are emergent from some other more complicated collective degrees of freedom. The distinction between particle and quasiparticle is not used consistently. We often speak of protons, pions, electrons, etc, as particles rather than quasiparticles, but they too are presumably just the low energy excitations of a more complicated theory.

1In fact the toric code is the paradigmatic example!

4From only this ground state degeneracy, we can conclude that the TQFT describing this system is abelian. See exercise 26.1

5The term “quasiparticle” is used for objects that act as relatively weakly interacting individual particles in low energy theories but are emergent from some other more complicated collective degrees of freedom. The distinction between particle and quasiparticle is not used consistently. We often speak of protons, pions, electrons, etc, as particles rather than quasiparticles, but they too are presumably just the low energy excitations of a more complicated theory.
The energy of each vertex defect is $\Delta_v$.

(3) We can have a plaquette where $P_\beta = -1$ instead of $P_\beta = +1$. The energy of each plaquette defect is $\Delta_p$.

The vertex and plaquette defects are often called electric and magnetic particles respectively ($e$ and $m$). The intuition for the names here is that the vertex defect is some sort of “charge” at the vertex (hence “electric”) and the plaquette defect is some sort of flux through the plaquette (hence “magnetic”).

**Caution:** Sometimes the the vertex defect is instead called magnetic and the plaquette defect is called electric! We will see in section 29.7 why this other labeling is sensible as well.

Fortunately, which particle is labeled $e$ and which one $m$ is at this point just a convention. The TQFT which will describe the system is actually the same independent of which one we label $e$ and which one we label $m$. See the discussion in chapter **. For the time being we will follow the convention that the vertex defect is electric and the plaquette defect is magnetic. Sometimes, however, it is easier to just call the defects vertex and plaquette defects to avoid confusion, and we will try to do this as much as possible!

Since vertex defects are produced in pairs, and can be brought back together and annihilated in pairs, we know we must have

$$\text{vertex defect} \times \text{vertex defect} = I$$

Similarly since plaquette defects $m$ are produced in pairs, and can be brought back together and annihilated in pairs we must also have

$$\text{plaquette defect} \times \text{plaquette defect} = I$$

In the more common notation we would write

$$e \times e = I$$

$$m \times m = I .$$

We might then wonder what happens if we bring together a vertex and a plaquette defect. They certainly do not annihilate! Thus

(4) We define another particle type, called $f$, which is the fusion of the magnetic and the electric particles (fusion of vertex and plaquette defect)

$$f = e \times m$$

The energy of such a particle is $(\Delta_v + \Delta_p)$.

Sometimes (particularly in the gauge theory literature) a particle that is a combination of an electric and magnetic particle is called a **dyon**.

We have the fusion relation

$$f \times f = I$$
which we can see by associativity and commutativity of fusion
\[ f \times f = (e \times m) \times (e \times m) = (e \times e) \times (m \times m) = I \times I = I \]

These three particle types \( e, m, f \) and the vacuum \( I \), are the only particle types there in this theory: First, they form a closed set under the fusion rules. Secondly, as we discussed in section ?? any “error” in this system (or excitation out of the ground state) can be described in terms of combinations of applications of \( \sigma_x \) and \( \sigma_z \), hence all excitations are generated by only the \( e \) and \( m \) particles. The full fusion relations are given in Table. 26.1. There are no non-abelian fusions here (i.e., each fusion gives a unique outcome) so we conclude we have an abelian theory.

Note that there are exactly four particle types (including the identity), and there are exactly four ground states on the torus, in agreement with the general principle that the number of particle types should match the number of ground states on the torus (See Eq. ?? for example).

### 26.2 Statistical Properties of Vertex and Plaquette Defects

Let us first consider the vertex defect particles (we have been calling this particle \( e \), but for more clarity we will stick with the term “vertex defect”). These particles are both created and moved around by applying \( \sigma_x \) operators. All of the \( \sigma_x \) operators commute with each other, so the end result is the same independent in what order we create, move, and annihilate the vertex defect particles with these \( \sigma_x \) operators.

There are now several “experiments” we can do to test the statistics of these particles. For example, we can create a pair of vertex defects, move one around in a circle and reannihilate, then compare this to what happens if we put another vertex defect inside the loop before the experiment. Since all of the creation and moving of particles is achieved by applying \( \sigma_x \), and all \( \sigma_x \) operators commute with each other, we see that the presence of another vertex defect inside the loop does not alter the phase of moving the defect around in a circle. Entirely analogously braiding a plaquette defect particle around another plaquette defect particle accumulates no phase: the defect is both created and moved by the \( \sigma_z \) operator and all of these \( \sigma_z \) operators commute with each other.

The experiments just described tells us that the vertex and plaquette defects (\( e \) and \( m \)) are either bosons or fermions. It is fairly obvious that braiding bosons around bosons never accumulates a phase. This is less obvious for fermions: while exchanging two fermions accumulates a \(-1\) sign, taking a fermion in a loop all the way around another fermion accumulates two minus signs (hence a \(+1\) sign, or no phase) since it is equivalent to two exchanges. Thus we have not yet determined the full statistical properties of these particles. To do so we will need to examine the twist factor \( \theta \) for these particles (See chapter ??).

To determine the twist factor of a particle, we will make a curl in a
Suppose we create a pair of vertex defect particles and move one around in a circle then reannihilate the pair (as in the right panel of Fig. 25.10). This process is created by a string of $\sigma_x$ operators. Recall that, if there are no vertex defects inside the loop this process does not accumulate a phase because the string of $\sigma_x$ operators around the loop is equivalent to the product of the $P_B$ plaquette operators enclosed (Eq. 25.13) — and in the ground state (the code space), the $P_B$ operators are in the $+1$ state. However, if there is one vertex defect particle inside the loop (as in Figs. 26.3 or right panel of Fig. 25.10), this means that one of the $P_B$ operators is actually in the $-1$ state. In this case the phase of taking the vertex defect particle around the loop is actually $-1$. So there is a phase of $-1$ for taking a vertex defect around a plaquette defect.

Another way to understand the braiding statistics is as follows: The operator that takes a vertex defect particle in a loop (the red loop in Fig. 26.3) is made of $\sigma_x$’s. The operator that creates a pair of plaquette defects and pulls them apart (The bright green line in Fig. 26.3) is made of $\sigma_z$’s (the dark green lines). These two operators intersect on a single edge (the one which has both a red line and a dark green line Fig. 26.3).
The $\sigma_x$ and $\sigma_z$ on this edge anticommute. Thus if we consider the processes of taking a vertex defect particle in a circle and then creating the plaquette defects and pulling them apart, and then compare it to the process where the plaquette defects are pulled apart first and then the vertex defect particle is taken in a circle which surrounds one of the two plaquette defect particles (as in Fig. 26.3), these two processes must differ by $-1$ due to the anticommutation of the two operators.

We can check that one accumulates exactly the same phase if we take an plaquette defect in a circle around a vertex particle. Taking a plaquette defect around in a loop is a process created by a string of $\sigma_x$ operators as shown in Fig. ?? (see also right of Fig. 25.14). If there are no vertex defects enclosed in the loop, this process does not accumulate a phase because the string of $\sigma_x$ operators around the loop is equivalent to the product of the $V_\alpha$ vertex operators enclosed (Eq. 25.15) — and in the ground state (the code space), the $V_\alpha$ operators are in the +1 state. However, if there is one vertex defect inside the loop, this means that one of the $V_\alpha$ operators is actually in the $-1$ state. In this case the phase of taking the plaquette defect around the loop is actually $-1$. So there is a phase of $-1$ for taking a plaquette defect around a vertex defect, showing nontrivial anyonic braiding with each other.

We have thus shown that taking an electric particle around a magnetic particle\(^6\) accumulates a phase of $-1$. In the language of chapter 13 we have just shown that

$$R_{ef}^m R_{me}^f = -1.$$  \hspace{1cm} (26.4)

Due to this $-1$ phase from taking $e$ around $m$, or $m$ around $e$, sometimes
354  The Toric Code as a Phase of Matter and a TQFT

\[ f = e \times m = - \]

\[ f = e \times m = - \]

\[ f = e \times m = - \]

\[ f = e \times m = - \]

**Fig. 26.4** The \( f \) particle is a fermion since its twist factor is \( \theta_f = -1 \). We can derive this by using \( f = e \times m \) and using the fact that \( e \) braiding around \( m \) gives a \(-1\) sign.

7Semions are particles that accumulate \( \pm i \) under exchange (see for example, section 18.1.2) so wrapping one semion all the way around another accumulates a phase of \(-1\). I personally do not like to use the term “relative semion” since the toric code is mostly unrelated to the semion theory.

**26.2.2 Properties of \( f \), the fermion**

Since \( f \) is made up of an \( m \) bound to an \( e \), we can derive the properties of \( f \) from our knowledge of the properties of \( m \) and \( e \). Braiding \( e \) around \( f \) is equivalent to taking \( e \) around both \( e \) and \( m \). Since \( e \) around \( e \) accumulates no phase, and \( e \) around \( m \) accumulates a phase of \(-1\), we conclude that taking \( e \) around \( f \) accumulates a phase of \(-1\). Similarly, taking \( m \) around \( f \) also accumulates a phase of \(-1\). Taking \( f \) around \( f \) is then equivalent to taking \( e \) and \( m \) both around \( f \), and since each of these processes accumulates a phase of \(-1\), the braiding of \( f \) around \( f \) accumulates a phase of \(+1\). As discussed in the case of \( e \) and \( m \) this tells us that \( f \) is either a boson or a fermion. To determine which one, we must again determine the twist factor \( \theta_f \). Perhaps the simplest way to determine this is via the diagrams in Fig. 26.4 which show that \( \theta_f = -1 \), hence \( f \) is actually a fermion (hence the notation, \( f \) for fermion). The astute reader will realize that this manipulation is actually a special case of the ribbon identity Eq. 15.3 (compare Eq. 26.4).

**26.3 S and T matrices**

We can summarize our findings about this anyon theory by stating the modular \( S_{ij} \) matrix, which lists the braiding result obtained by taking particle \( i \) around particle \( j \) as shown in Fig. 7.13 and the \( T \) matrix (Eq. 17.14) which simply lists the twist factors along the diagonal.

Listing the particles in the order \( I, e, m, f \) we can write \( S \) and \( T \) as in

\[
S = \frac{1}{D} \begin{pmatrix}
1 & 1 & 1 & 1 \\
1 & 1 & -1 & -1 \\
1 & -1 & 1 & -1 \\
1 & -1 & -1 & 1
\end{pmatrix}
\quad T = \begin{pmatrix}
1 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 \\
0 & 0 & 1 & 0 \\
0 & 0 & 0 & -1
\end{pmatrix}
\quad \text{(26.5)}
\]

where unitarity fixes the total quantum dimension \( D = 2 \) in agreement also with the definition Eq. 17.11 that \( D^2 = \sum_i d_i^2 \). We can also check that the central charge (via Eq. 17.16) must be zero mod 8.
26.4 Charge-Flux Model

We can describe the statistics of the particles in the toric code using a charge-flux model somewhat analogous to Chern-Simons theory\(^8\). Here let us define

- **electric particle** = \( e \) = particle bound to 1 unit of electric charge
- **magnetic particle** = \( m \) = particle bound to \( \pi \) units of magnetic flux
- **fermion** = \( f \) = particle bound to 1 unit of electric charge and \( \pi \) units of magnetic flux

It is easy to see that this charge and flux will correctly give the +1 and -1 phases accumulated from braiding particles.

26.5 \( \mathbb{Z}_N \) Toric Code (Briefly)

To generalize the toric code Hamiltonian Eq. 26.1 to the case of the \( \mathbb{Z}_N \) toric code discussed in section 25.6, we need to find operators analogous to \( V_{\alpha} \) and \( P_{\beta} \) in which will favor \( Q_{\alpha} = 0 \) on the vertices and \( A_{\beta} = 1 \) on the plaquettes as the ground state. We will choose to work with the following Hamiltonian\(^9\)

\[
H_{\mathbb{Z}_N \text{ toric code}} = -\Delta_v \sum_{\text{vertices} \alpha} \delta Q_{\alpha,0} - \Delta_p \sum_{\text{plaquettes} \beta} \hat{P}_{\beta} \quad (26.6)
\]

where \( \delta \) is a Kronecker delta function and

\[
\hat{P}_{\beta} = \frac{1}{N} \sum_{p=0}^{N-1} (A_{\beta})^p . \quad (26.7)
\]

The operator \( \hat{P}_{\beta} \) gives unity on a plaquette where the eigenvalue of \( A_{\beta} \) is 1, and gives zero otherwise (To see this recall that \( (A_{\beta})^N = 1 \) so the eigenvalues of \( A_{\beta} \) can only be \( e^{2\pi i k/N} \) for some integer \( k \)). Similarly the Kronecker delta function gives unity on a vertex where \( Q_{\alpha} = 0 \) and gives zero otherwise.

The \( N = 2 \) case of this Hamiltonian is not quite identical to Eq. 26.1 for the conventional toric code. Whereas \( V_{\alpha} \) and \( P_{\beta} \) in Eq. 26.1 have eigenvalues ±1, the terms \( \delta Q_{\alpha,0} \) and \( \hat{P}_{\beta} \) here have eigenvalues 0 and 1. However, we do have (for the \( N = 2 \) case) that

\[
V_{\alpha} = \frac{1}{2} (\delta Q_{\alpha,0} - 1) \quad P_{\beta} = \frac{1}{2} (\hat{P}_{\beta} - 1)
\]

and note that Eq. 26.1 already has the factors of \( \frac{1}{2} \) out front of each term compared to that of 26.6. Thus we see that Eq. 26.1 and Eq. 26.6 differ only by the addition of an unimportant constant.

The ground state of the Hamiltonian Eq. 26.6 is any state where all

\[^{8}\text{Precisely the toric code can be described as a Chern-Simons theory with two Chern-Simons fields } a_\mu^I \text{ with } I = 1, 2 \text{ and an action which we write as } S = \frac{1}{4\pi} \int d^3 x K_{1,\gamma} a_\mu^1 a_\mu^2 a_\mu^\gamma \text{ and a purely off-diagonal coupling matrix } \]

\[
K = \begin{pmatrix} 0 & 2 \\ 2 & 0 \end{pmatrix}
\]

Such Chern-Simons theories with multiple fields will be discussed in more detail in chapter **.

\[^{9}\text{This form is compatible with the more general form used in chapter 26 below.}\]
\( Q_\alpha = 0 \) and all \( A_\beta = 1 \). I.e., the code space of the \( \mathbb{Z}_N \) toric code. The defect energies are \( \Delta_\alpha \) for a vertex defect and \( \Delta_\beta \) for a plaquette defect.

If we move vertex defects together, their value of \( Q_\alpha \) add modulo \( N \), where \( Q_\alpha = 0 \) corresponds to an unexcited vertex and all other values of \( Q_\alpha \) are excitations (i.e., we have \( \mathbb{Z}_N \) fusion rules). Similarly if we move plaquette defects together, their eigenvalues of \( A_\beta = e^{2\pi ik/N} \) with \( k \) an integer multiply with \( A_\beta = 1 \) being an unexcited plaquette and all other values being excited (again this is \( \mathbb{Z}_N \) fusion rules). We can also construct quasiparticles that correspond to combinations of excited vertices and excited plaquettes. The most general particle type can then be notated as \([n, k]\) with \( n, k \in 0, \ldots, (N - 1) \) by which we mean

\[
[n, k] \rightarrow \left\{ \begin{array}{ll}
Q_\alpha &= n \\
A_\beta &= e^{2\pi ik/N}
\end{array} \right.
\]

thus there are a total of \( N^2 \) particle species. If a particle has both nontrivial vertex quantum number and nontrivial plaquette quantum number it is sometimes called a dyon. The fusion rules for these particles are then given by

\[
[n_1, k_1] \times [n_2, k_2] = \left[ (n_1 + n_2) \text{mod}(N), (k_1 + k_2) \text{mod}(N) \right]
= [n_1 + n_2, k_1 + k_2]
\]

where in the second line addition is assumed to be modulo \( N \). Note that these fusion rules imply we have an abelian anyon theory.

Following similar arguments to section 26.2 it is easy to show that all of the vertex defects and all of the plaquette defects are bosons — and further, all vertex defect braid trivially with all vertex defects and all plaquette defects braid trivially with all plaquette defects. However, given Eqs. 26.22 and 26.23 it is clear that the vertex defects braid nontrivially with the plaquette defects, which also gives nontrivial braiding properties to the dyons. In particular these two equations showed that taking a \( Q_\alpha = +1 \) charge clockwise around a \( A_\beta = e^{2\pi i/N} \) plaquette give a phase of \( e^{2\pi i/N} \) which we can rewrite as \( R \)-matrices

\[
R_{[0,1],[1,1]}^{[0,1],[0,1]} R_{[1,1]}^{[1,0],[0,1]} = e^{-2\pi i/N}
\]

and more generally we can similarly derive the phase for taking an \([n_1, m_1]\) particle counterclockwise all the way around an \([n_2, m_2]\) particle

\[
R_{[n_1+n_2+k_1+k_2],[n_1+k_1]}^{[n_2+k_2],[n_1,k_1]} R_{[n_1+n_2+k_1+k_2]}^{[n_1+k_1+n_2+k_2]} = e^{-2\pi i(n_1k_2+n_2k_1)/N}
\]

where addition is assumed to be modulo \( N \). Further, using the ribbon identity Eq. 15.3 we obtain

\[
\theta_{[n,k]} = e^{-2\pi i nk/N}
\]

Using either Eq. 26.9 directly to evaluate the relevant diagram, or using

---

10. The exponent on the right has a minus sign because the \( R \) matrix is defined as the phase associated with a counterclockwise exchange.
Eq. [17.20] we can obtain the $S$-matrix for the $\mathbb{Z}_N$ toric code

$$S_{[n'k'],[n,k]} = \frac{1}{N} e^{2\pi i (nk' + n'k)/N} \quad (26.11)$$

We can write a charge-flux model description for each of these particle types by modeling the $[n,k]$ particle as $n$ units of electric charge bound to $2\pi k/N$ units of magnetic flux.

**Exercises**

**Exercise 26.1 Show Toric Code is Abelian from Ground State Degeneracies Only**
Given that the ground state degeneracy for the toric code on an $g$-handled torus is $2^{2g}$, show that this must be an abelian TQFT with $2^g$ particle types.
(Hint: See exercise 8.2 and (gs degen on a torus is # of particle types))

**Exercise 26.2 Ground State Degeneracy of Toric Code**
Starting with the fusion rules of the toric code, calculate the ground state degeneracy of the toric code on $g$-handled torus using the methods of section 8.4.

**Exercise 26.3 Twist of the Plaquette Defect**
In section 26.2 we explicitly calculate that the twist factor for a vertex defect particle is trivial. Use an analogous technique to show the twist factor for the plaquette defect particle is also trivial.

**Exercise 26.4 Checking the Verlinde Formula for the Toric Code**
Confirm the Verlinde formula Eq. [17.13] for the toric code.

**Exercise 26.5 Twist factor for excitations of the $\mathbb{Z}_N$ toric code**
Confirm Eq. [26.10]
Robustness of Topologically Ordered Matter

Topologically ordered matter, matter that can be described as a TQFT, has a remarkable property known as topological robustness — the topological properties of the phase of matter are unchanged if the Hamiltonian is changed a small bit (so long as the gap to making excitations remains). While this property is very general, we will illustrate it using the toric code, and generalization to other types of topologically ordered matter are then fairly obvious.

27.1 Perturbed Hamiltonian

Let us start with the toric code Hamiltonian, and add some perturbing term.

\[ H = H_{\text{toric code}} + \delta H \]  

(27.1)

where \( H_{\text{toric code}} \) is the toric code Hamiltonian defined in Eq. 26.1. In Eq. 27.1 \( \delta H \) is some arbitrary small perturbation Hamiltonian which is the sum of arbitrary local terms\(^1\). The claim is that for small enough \( \delta H \) (but not necessarily infinitesimally small!), the topological properties of the phase of matter — such as the 4-fold degenerate ground state, the types of excitations, and their braiding statistics — will remain entirely unchanged in the large system size limit\(^2\). We will focus first on the ground state degeneracy, but the arguments for the other topological properties will follow similarly.

27.1.1 Robustness of Ground State Degeneracy

Rough Argument

The general idea is as follows: When \( \delta H \) is turned on, it can be treated in perturbation theory order by order. If \( \delta H \) is small, then higher orders in perturbation theory are successively less important. Now at some \( M^{th} \) order in perturbation theory we can consider \( M \) applications of \( \delta H \) to the unperturbed system. Each application of \( \delta H \), which is some arbitrary local operator, can create, annihilate, or move some quasiparticle excitations. The key point is that in order to break the ground state degeneracy in any way, which is equivalent to causing a logical error in the toric code when used as a quantum memory, a quasiparticle must be taken all the way around a cycle of the torus. For a very large torus this

\(^1\) An operator is local if it acts on a finite number of spins which are all near each other — meaning within some fixed finite distance as we take the thermodynamic limit in the problem. A term which would not be local would be a product of all of the spins in the system, or a product of all the spins in a string all the way around a cycle of the torus.

\(^2\) This statement should become exponentially more accurate as the size of the system is made larger and larger.
Robustness of Topologically Ordered Matter

can only happen at extremely high order in perturbation theory, and hence any splittings are only exponentially small.

In More Detail

Let us now make this argument more rigorous. It is easiest to choose a particularly simple form for $\delta H$ to work with. However, once we understand the physical principle, we will realize that the actual form we choose doesn’t matter and the results hold much more generally. For simplicity, let us choose $\delta H$ to be the sum of $\sigma_x$ over all spins

$$\delta H = J \sum_1 \sigma_x^{(i)}$$  (27.2)

where $J$ is a parameter with dimensions of energy which is assumed to be small compared to the excitation gap $\Delta$ of the toric code.\(^3\)

We treat $\delta H$ in a Brillouin-Wigner perturbation expansion which “integrates out”\(^4\) excitations and leaves us with an effective Hamiltonian within the previously degenerate ground state space. (See appendix \(^{27.3}\) for detailed derivation of this perturbation expansion). The new effective Hamiltonian within this previously degenerate ground state space is given by

$$H^\text{eff}_{pn} = E_0 + \langle p | \delta H \frac{1}{1 - G \delta H} | n \rangle$$  (27.3)

$$= E_0 + \langle p | \delta H | n \rangle + \langle p | \delta H G \delta H | n \rangle + \langle p | \delta H G \delta H G \delta H | n \rangle + \ldots$$

where $| p \rangle$ and $| n \rangle$ are states in the ground state space of the unperturbed model, and $E_0$ is the unperturbed ground state energy. Here $G$ is the Green’s function

$$G(E) = \sum_{n \in \text{excited}} \frac{| n \rangle \langle n |}{E - E_n^0}$$

and the sum is over (non-ground state) eigenstates $| n \rangle$ of the unperturbed model whose energies are $E_n^0$. Note that one must plug in the eigenenergies $E$ of $H^\text{eff}$ into $G(E)$ to find self-consistent solutions\(^5\).

It is crucial to note that each factor of $G$ has an energy denominator on the order of the gap $\Delta$ or larger, and each factor of $\delta H$ has an energy numerator on the order of the coupling constant $J$. Thus the expansion shown in Eq. (27.3) is actually an expansion in the ratio $J/\Delta$.

Our claim is that, to very high order in this expansion, the effective Hamiltonian within this ground state space will remain proportional to the identity matrix. I.e., the ground state space does not split at all, as we previously claimed. To see why this is, let us consider the lowest few orders of the perturbation expansion. The first term is $\langle p | \delta H | n \rangle$. Here $| n \rangle$ and $| p \rangle$ are states in the ground state of the unperturbed Hamiltonian. Each term in $\delta H$ applies a $\sigma_x$, thus creating two vertex defect excitations. Since $\delta H | n \rangle$ is an excited state and $| p \rangle$ is an unexcited state, there is zero overlap between the two.

The next order term of the perturbation expansion is $\langle p | \delta H G \delta H | n \rangle$.
such that $\delta H$ is applied twice. Two applications of $\delta H$ can create two pairs of quasiparticles in different places; or the first $\delta H$ could create one pair and the second $\delta H$ could be applied to a neighboring spin to pull the two quasiparticles further apart (as in the left of Fig. 25.10). In either case the matrix element will still be zero since there would be no overlap between the excited state $\delta HG\delta H|n\rangle$ and the unexcited state $|p\rangle$. However, there is a third process we could consider, where the first $\delta H$ acts on edge $i$ to create a pair of quasiparticles and then the second $\delta H$, acting on the same edge $i$, annihilates the quasiparticles. In this case we return again to the ground state that we started in and we have a nonzero overlap. (The value of the matrix element is $J/(E - E_{\text{excited}}) \sim J/\Delta$.) Crucially, it does not matter which of the four ground states $|n\rangle$ we started in, we are always returned to the same ground state when the quasiparticles are annihilated again. So the contribution to $H^{\text{eff}}_{pn}$ is just $\delta_{pn}J/(E - E_{\text{excited}})$, meaning this does not cause any splitting of the ground state degeneracy.

The argument continues similarly at higher orders of perturbation theory. Each application of $\delta H$ creates, annihilates, or moves quasiparticles (Considering the form of $\delta H$ in Eq. 27.2, all quasiparticles are vertex defects in this simplified argument). In order to get a nonzero matrix element, the successive applications of the $\delta H$ operators must return the system to the ground state space at the last step — i.e., any quasiparticles that are created must then be reannihilated. If all the quasiparticles are reannihilated, the value of the matrix element is simply $J$ for each application of $\delta H$ with the Green’s functions contributing energy denominators corresponding to the energies of the (necessarily) excited intermediate states. When all of the quasiparticles are reannihilated at the last step, the system returns to exactly the same ground state it started from unless a quasiparticle has gone all the way around the cycle of the torus, a case we will address in a moment.

If we assume that no quasiparticles have gone all the way around the cycle of the torus then $H^{\text{eff}}$ must be proportional to the identity. Each ground state returns to the same ground state, and the matrix elements are independent of which ground state you started in. In this case the ground state degeneracy does not split. However, if a quasiparticle does go around a cycle of the torus (as in Fig. 25.11) then the final ground state is different from the initial ground state, meaning that there is an off-diagonal contribution to $H^{\text{eff}}$ (compare Eq. 25.14), and $H^{\text{eff}}$ is no longer proportional to the identity matrix. Diagonalization of $H^{\text{eff}}$ then gives a splitting of the previously degenerate ground state.

Crucially, for a torus whose (smallest) cycle length is $L$, it requires at least $L$ applications of $\delta H$ to move a quasiparticle around the cycle. This means that only at $L^{th}$ order in perturbation theory will we find off-diagonal terms in the effective Hamiltonian $H^{\text{eff}}$. Since each $\delta H$ is of size $J$ and each order in perturbation theory comes with a Green’s function $G$ with energy denominator on order of the gap $\Delta$, then the size of these off-diagonal terms is given by $\sim (J/\Delta)^L$ which is exponentially small in the size of the system, assuming $J$ is smaller than order $\Delta$. Thus
the ground state degeneracy only splits an exponentially small amount, with the splitting going to zero as the size of the system gets large.

The argument is quite similar for other possible forms of $\delta H$. For example, let us consider the perturbation

$$\delta H = J \sum_i \sigma_z^{(i)} \quad (27.4)$$

Very similarly here each application of $\delta H$ creates, annihilates, or moves a quasiparticle; only this time the quasiparticles are plaquette defects. Again, as long as no quasiparticles are moved around a cycle of the torus, the resulting contributions to $H_{\text{eff}}$ are proportional to the identity. However, if a plaquette quasiparticle is moved around a cycle (as in Fig. 25.15) then there will be a contribution to $H_{\text{eff}}$ which is different for the different ground state sectors (compare Eq. 25.16) thus breaking the ground state degeneracy. As with the argument for the vertex defects, the splitting of the ground state degeneracy only occurs at order $L$ in perturbation theory (with $L$ the length of the cycle of the torus) and is thus exponentially small in the size of the system.

It is clear that this general argument is not specific to the particular form of $\delta H$ we have chosen. In order to have such an exponentially protected ground state degeneracy, $\delta H$ need only be local, i.e., it can be any sum of terms, each of which operates on a finite set of spins near each other. In particular, we need to have a situation where the number of applications of $\delta H$ which is required to cause a quasiparticle to go around the cycle of the torus should scale with the linear size of the system.

It is important to realize that in the presence of the perturbation Hamiltonian, the ground states have virtual excitations mixed into them. (Sometimes one says that the ground state is “dressed” with virtual excitations). If one were to measure the plaquette and vertex operators, one would find some number of virtual excitations are in the $-1$ state rather than the $+1$ state. However, if $\delta H$ is small, the density of these virtual excitations is low, and such excitations occur in local annihilatable pairs which, in the language of error correcting codes, are easily correctable. In the language of this Hamiltonian, these do not change the topological properties of the phase of matter.

27.1.2 Quasiparticles

One can further ask what happens to the nature of the quasiparticles when a perturbation is applied to the Hamiltonian. In absence of the

---

6 For example, at first order in the perturbation series there is a contribution $G\delta H|\psi\rangle$ to the wavefunction where $|\psi\rangle$ is in the code-space. For $\delta H$ of the form of Eq. 27.2 or 27.4 the operator $\delta H$ creates exactly two defects when acting on the code space. See Eq. 27.9 and particularly exercise 27.1.

7 One might worry about the convergence of the perturbation theory. (Rigorous perturbation arguments using a different technique have been given by Bravyi et al. [2010].) Obviously if the perturbation is large enough that some excited states become lower energy than the ground state, then the argument no longer makes sense. Indeed, it is generally assumed that the perturbative expansion should hold until the point where the gap closes. However, one might also worry that the perturbative expansion may fail to converge since, while each successive order in the expansion is reduced by $(J/\Delta)$, there are also many many more terms to consider in each higher order of perturbation theory. At least for small enough $J/\Delta$ (but not requiring infinitessimally small $J/\Delta$) this issue does not cause trouble as discussed in exercise 27.1.
perturbation, the quasiparticles have well-defined topological properties—in particular they have nontrivial braiding statistics (which we worked out for the toric code in chapter 26). We claim that, so long as we are considering physics at long length scales, these topological properties will remain unchanged under small perturbations.

Rough Idea

The general strategy is again to perform a perturbation expansion. We will find that in the presence of the perturbation, the quasiparticles will develop a surrounding cloud of virtual excitations\(^8\) (this is often called a “dressing”). The quasiparticles are then no longer point particles, but rather develop a length scale which becomes longer and longer as the perturbation is increased. However, the braiding properties of the quasiparticles will remain unchanged, so long as the particles remain far enough apart that their dressing clouds do not intersect.

In More Detail

For simplicity let us focus on the perturbing Hamiltonian Eq. 27.2 which only makes vertex defects. If the perturbing Hamiltonian is small, we will mix into the ground state a low density of virtual excitations, most of which will simply be two vertex defects separated by a single edge (See exercise 27.1).

The perturbation we have chosen (Eq. 27.2) commutes with the plaquette operators \(P_\beta\), so that the plaquette excitations can still be described as simply a plaquette where \(P_\beta = -1\). I.e., given any eigenstate of the system (with or without the perturbation Eq. 27.2) one may obtain another eigenstate by applying \(\sigma_z\) to an edge which then flips the eigenvalue of \(P_\beta\) on the neighboring plaquettes. The lowest energy configuration (as in the unperturbed toric code) is when all of the plaquettes have eigenstate \(P_\beta = +1\) and we consider \(P_\beta = -1\) to be a plaquette excitation.

Let us now start in the perturbed ground state and add a pair of plaquette excitations. We can create these particles and move them around by applying \(\sigma_z\) to edges as in Fig. 25.14. As we move the plaquette excitations through the system, they will cross through some of the virtually excited red edges—i.e., in only some terms of the superposition that make up the wavefunction does the plaquette defect cross through red edges. These terms of the wavefunction incur a minus sign due to the plaquette defect particle motion. We can think of these signs as being the result of the plaquette defect swimming through the soup of pairs of virtual vertex defect particles.

Now suppose we have added a pair of vertex quasiparticles that have been pulled far apart from each other\(^9\). If we measure every vertex, we would obtain a picture like Fig. 27.1. Here the isolated quasiparticles that are not part of a local pair can still be clearly identified. They are connected together by a long red string. We should remember, as discussed in section 25.4.1 that the red string is not actually measurable

\[^{8}\text{As mentioned at the end of the last section, the ground state becomes “dressed” with virtual excitations too!}\]

\[^{9}\text{When we add a perturbing Hamiltonian like Eq. 27.2 we can give the quasiparticles dynamics, allowing them to move from their initial positions. While there is nothing in principle wrong with this, it does make it harder to study the quasiparticles if they keep moving. To avoid this problem we can imagine a “pinning potential” to trap the quasiparticles and prevent them from moving from whatever position we put them at (For example, we can lower the prefactor of } V_\alpha \text{ in the Hamiltonian Eq. 25.1 only at one vertex which makes it energetically favorable for the vertex defects to sit at that position).}\]
and its actual position is unknown (or we can say the position of the red string fluctuates) — only the endpoints are measurable. Note that the position of the endpoints may also fluctuate due to virtual excitation pairs that may appear at the end position to extend or shorten the long red string. This makes the quasiparticle at the end of the string look like a cloud of finite size rather than a point particle.

Now let us repeat the experiments of creating and moving plaquette defects, but having first introduced the two additional quasiparticles separated by a long distance as in Fig. 27.1. Now as we move the plaquette defect, the wavefunction again picks up phases associated with swimming through the thick soup of virtual excitations. However, now, if it crosses over the long red line as well, this will then create a $-1$ braiding phase. Thus, the phase of braiding the plaquette defect around the vertex defect quasiparticle is independent of the fact that there are lots of pairs of virtual vertex defect particles.

We should be cautious, however, if we pass the plaquette defect close to the end of the long red string — i.e., close to the position of the (not-virtual) vertex defect quasiparticle. Because of the fluctuations of the position of the quasiparticle, it may not be possible to determine if the plaquette defect has gone around the vertex defect or not.

We could just as well have chosen the perturbation Eq. (27.4) which would make virtual plaquette defect excitations, fluctuates the position of the plaquette defect particle, but leave the vertex defect excitations unchanged. The argument would be almost exactly the same. Again we find a braiding phase of $-1$.

Treating the case where the perturbation creates both vertex and plaquette excitations is a bit harder to do rigorously, and some different techniques become useful which we will not discuss here. However, generally the same principles apply: the quasiparticles develop some finite size (which grows larger for larger perturbation), but as long as the quasiparticles are braided at distances much larger than this size, the braiding phases are unchanged by the perturbation.

27.2 Topologically Ordered Matter

27.2.1 Importance of Rigidity

The fact that the quasiparticle statistics and ground state degeneracy are unchanged when the Hamiltonian is modified a small bit should not be surprising to us given the ideas of rigidity we have run into in section 9.3 and 13.3: It is not possible to perturb the properties of a TQFT a small amount and maintain a consistent TQFT. As we perturb the Hamiltonian with any local operators, to have any change in the long length scale properties of the TQFT that results, the system needs to go through a phase transition. In other words, the topological properties are a property of the entire phase of matter.
27.2.2 The Notion of Topological Order

The type of protection from small perturbations that we have just discovered, particularly in the discussion of section 27.1.1, is the basis for a very useful definition of topological order. Let us assume we have multiple degenerate ground states on a surface with nonzero genus (i.e., a torus, two-handled torus, etc) which we call $|\psi_i\rangle$ with $i = 1, \ldots, M$ with $M > 1$ the ground state degeneracy. We often define topological order to be the property that

\[
\langle \psi_i | \hat{O} | \psi_j \rangle = C_\hat{O} \delta_{ij} \tag{27.5}
\]

where $\hat{O}$ is any local operator and where $C_\hat{O}$ is a constant depending on the particular local operator $\hat{O}$ we are considering. Finite size corrections to this result, if any, must be exponentially small in the size of the system. In other words, the multiple ground states look just like each other locally, but are mutually orthogonal. Note that since any perturbation we add to the Hamiltonian is assumed local, this condition guarantees that the ground state degeneracy remains robust under perturbations.

This definition of topological order is effectively the same as saying that the system is described by a TQFT. While a TQFT must have the property Eq. 27.5 it is not immediately obvious that having this property is enough to conclude that we have a TQFT. However, there are no systems known with this property that are not essentially TQFTs.

27.2.3 Defining a Topological Phase of Matter

The classic definition of a phase of matter is a region of thermodynamic parameter space where certain (or even most) physical properties of a system are robustly unchanged. For example, water is one phase of matter (robustly characterized as being liquid over a large range of temperature and pressure, whereas another phase of matter of the same system is ice, which is robustly characterized as being solid\(^\text{12}\).

For our consideration of topological quantum systems, we would like to ask how we should best characterize different possible phases of our systems. One obvious possibility is to characterize each topologically ordered phase of matter in terms of its TQFT in the long length scale limit. Recall that the perturbed toric code only has the the same topological properties as the unperturbed toric code (same ground state degeneracy, same quasiparticle statistics) if we look at long length scales (large system, excitations far apart). Indeed, characterization in terms of the long length scale TQFT is one good way to define a topological phase of matter.

However, there is also a more abstract (and topological!) way to define a phase of matter, which can be invoked much more generally — even in cases where we do not have a nontrivial TQFT. Here we use as inspiration the perturbation theory argument in the prior section and we note that this argument only fails when the gap to make excitation

12 The example of water/ice is actually not a particularly good example for our purposes. All of the physics discussed in this book is zero temperature — phase transitions can occur as a function of some tuning parameter in the Hamiltonian. Such zero temperature phase transitions are known as quantum phase transitions. A nice example of a quantum phase transition is the zero temperature phase transition between solid Helium and liquid Helium as a function of pressure.
becomes zero. Thus we make the following definition:

**Definition of a Topological Phase of Matter:** Two (zero temperature) gapped states of matter are in the same topological phase of matter if and only if you can continuously deform the Hamiltonian to get from one state to the other without closing the excitation gap.

This sort of definition can obviously be used much more generally to distinguish different phases of matter. Further this definition fits with our intuition about topology:

Two objects are topologically equivalent if and only if you can continuously deform one to the other.

### 27.3 Appendix: Brillouin-Wigner Perturbation Theory

Here we will discuss a slight variant of Brillouin-Wigner perturbation theory designed for the case of interest where we have a degenerate ground state and a gap to excitations $\Delta$.

Here we assume a Hamiltonian $H = H_0 + \delta H$. The orthonormal eigenstates of $H_0$ are $|n\rangle$ with energies $E_n$ and the eigenstates of $H$ we call $|\tilde{n}\rangle$ with energies $E_{\tilde{n}}$. The degenerate ground state space of $H_0$ we call $S$ (of some dimension $d$) and all other eigenstates of $H_0$ are in the space $\bar{S}$. The energy of the ground state space of $H_0$ is $E_0$.

Our strategy will be to “integrate out” the excited states leaving ourselves with an effective Hamiltonian within the ground state space $S$.

Projectors onto the unperturbed ground state space and unperturbed excited state space are

$$P = \sum_{n \in S} |n\rangle \langle n|, \quad Q = 1 - P = \sum_{n \in \bar{S}} |n\rangle \langle n|$$

As we turn on the perturbation, the $d$ orthogonal states in the space $S$ will generally split in energy (although for our TQFT we expect this splitting to be exponentially small if $\delta H$ is local). Our approach remains valid for perturbation strengths smaller than the point where the splitting is on the order of the gap $\Delta$.

Let us start with a $d$-dimensional basis of perturbed wavefunctions $|\tilde{m}\rangle$ with energies $E_{\tilde{m}}$ in the perturbed ground state space $S$. Let us project these states into the original ground state space

$$P|\tilde{m}^u\rangle = |\psi_{\tilde{m}}\rangle = \sum_{n \in S} C_{n\tilde{m}} |n\rangle$$

We have marked the state $|\tilde{m}\rangle$ with a superscript $u$ because we will work with an unnormalized state on the left so that $|\psi_{\tilde{m}}\rangle$ is normalized to unity $\langle \psi_{\tilde{m}} | \psi_{\tilde{m}} \rangle = \sum_{n \in S} |C_{n\tilde{m}}|^2 = 1$. 

We write the Schrödinger equation for the perturbed state as \((H_0 + \delta H)|\tilde{m}^u\rangle = E_{\tilde{m}}|\tilde{m}^u\rangle\) as

\[
(E_{\tilde{m}} - H_0)|\tilde{m}^u\rangle = \delta H|\tilde{m}^u\rangle
\]  
(27.8)

Applying \(Q\) to the left and dividing through by \(E_{\tilde{m}} - H_0\) we obtain

\[
Q|\tilde{m}^u\rangle = G\delta H|\tilde{m}^u\rangle
\]

where \(G\) is the Green’s function

\[
G = \sum_{n \in S} \frac{|n\rangle \langle n|}{E_{\tilde{m}} - E_n}
\]

Note that this is nonsingular as long as the energy \(E_{\tilde{m}}\) of the states that were in the ground state space do not come close to the energies of the unperturbed excited states \(E_n\).

We then write

\[
|\tilde{m}^u\rangle = P|\tilde{m}^u\rangle + Q|\tilde{m}^u\rangle = |\psi_{\tilde{m}}\rangle + G\delta H|\tilde{m}^u\rangle
\]

which we can rewrite as

\[
(1 - G\delta H)|\tilde{m}^u\rangle = |\psi_{\tilde{m}}\rangle
\]

or

\[
|\tilde{m}^u\rangle = \frac{1}{1 - G\delta H}|\psi_{\tilde{m}}\rangle
\]  
(27.9)

Note in particular that only the first term in the series is within the space \(S\), and all other terms are entirely outside of this space.

To find the energy of the state \(|\tilde{m}^u\rangle\) we apply an arbitrary state \(|p\rangle\) within the space \(S\) to the left of Eq. 27.8 to obtain

\[
\langle E_{\tilde{m}} - E_0|p|\tilde{m}^u\rangle = \langle p|\delta H|\tilde{m}^u\rangle = \langle p|\delta H \frac{1}{1 - G\delta H}|\psi_{\tilde{m}}\rangle
\]

\[
(E_{\tilde{m}} - E_0)C_{p\tilde{m}} = \sum_{n \in S} \langle p|\delta H \frac{1}{1 - G\delta H}|n\rangle C_{n\tilde{m}}
\]

We recognize this as an eigenvalue/eigenvector problem within the space \(S\)

\[
(E_{\tilde{m}} - E_0)C_{p\tilde{m}} = \sum_{n} H_{p\tilde{m}}^\text{eff} C_{n\tilde{m}}
\]

where \(H_{p\tilde{m}}^\text{eff}\) is the effective Hermitian Hamiltonian within the low energy subspace

\[
H_{p\tilde{m}}^\text{eff} = \langle p|\delta H \frac{1}{1 - G\delta H}|n\rangle
\]

Note that this is a bit more tricky than just a simple eigenvalue problem.
because $G$ is implicitly dependent on $E_{\tilde{m}}$. This self-consistency has a few implications, including the fact that the eigenvectors $C'_{\hat{m} \tilde{m}}$ or equivalently $|\psi_{\tilde{m}}\rangle$ are not orthogonal. It is the eigenvector $|\hat{m}\rangle$ of the full Hamiltonian which are orthogonal, and $|\psi_{\tilde{m}}\rangle$ are normalized projections of $|\tilde{m}\rangle$.

**Further Reading**

Mention Wen being the creator of the definition of topological order

---

**Exercises**

**Exercise 27.1 Convergence of Perturbation Expansion**

Considering a large toric code system with $N$ edges, with the Hamiltonian Eq. 27.1 and a simple perturbation Eq. 27.2.

(a) If all the vertices are simultaneously measured, show that, to second order in perturbation theory, the probability of finding no defects is

$$P_{\text{box}} = \frac{1}{1 + (N/4)[J/\Delta]^2}$$

with $\Delta = \Delta_v$ being the energy of a vertex defect. Show that the probability of exactly one edge being flipped (i.e., finding two vertex defects, separated by one edge) is $1 - P_{\text{box}}$. Show that to this order in perturbation theory the probability that a given edge is flipped is

$$p_{\text{edge,ex}} = \frac{1}{4} (J/\Delta)^2$$

(b) Assume that the probability of a particular edge being flipped (meaning that $\sigma_x$ has acted on that edge) is small. In this case we can ignore the possibility of a defect being created and then moved by one step as in the left of Fig. 25.10. I.e., we are assuming that all applications of $\sigma_x$ are on isolated edges far from other defects. We can then think of the system as being $N$ independent uncoupled edges each of which is a two level systems. Show that the probability of a given edge being flipped is exactly

$$p_{\text{edge,ex}} = \frac{2 + (J/\Delta)^2 - 2\sqrt{1 + (J/\Delta)^2}}{2 + 2(J/\Delta)^2 - 2\sqrt{1 + (J/\Delta)^2}} \approx \frac{1}{4} (J/\Delta)^2 + \ldots$$

Hint: think about a two level system!

This approximation corresponds to resumming a subset of the terms in the perturbation expansion (out to infinite order) corresponding to isolated pairs of vertex defects. At least for small $J/\Delta$ the neglect of other terms is justified by the low density of defects.
Abstracting the Toric Code: Introduction to Tube Algebra

In section 25.5 we pointed out that we can build the toric code on any lattice (see also Fig. 28.1). Indeed, in many respects it is easiest to dispense with the lattice altogether. This simplifies a lot of the thinking and allows us to generalize the model to describe more complex TQFTs which we will do in chapter *** - ***.

The basic idea here is that the toric code can be viewed as simply a gas of fluctuating loops or strings — without need to tie it to the underlying lattice. The rules that describe this loop gas are planar diagrammatic rules — the same kind of diagrammatic rules we have been using all along! If we want to put the model back on a lattice at the end of the day, we can do this, but in fact many of the manipulations are in fact simpler without the burden of the lattice.

Before making the transition to the continuum, it is useful to first work with a trivalent lattice where only three edges meet at a vertex. For example, we could take a honeycomb as shown in the top Fig. 28.1. This eliminates situations where four lines intersect at a corner as shown in the bottom of Fig. 28.1. (We need not work with a regular lattice, see section 25.5, but it is convenient for simplicity.) The rule that all vertices have an even number of blue edges coming into them (Rule 1 from section ??) now means that the allowed configurations in the toric code ground state are nonintersecting loops as compared to the intersecting loops that are allowed, for example, on a square lattice.

### 28.1 Toric Code as a Loop Gas

We start by abstracting the toric code to simply a gas of fluctuating non-intersecting loops — no longer paying attention to a lattice. An example of a loop gas configuration is shown in Fig. 28.2.

We can write the toric code wavefunction (See Eq. 25.10) in the form of

$$|\psi\rangle \sim \sum_{\text{all loop configs that can be obtained from a reference loop config via allowed moves}} |\text{loop config}\rangle$$

(28.1)

where the allowed moves are inspired by the moves we can make on a lattice. We list these allowed moves in Fig. 28.3.

---

**Fig. 28.1** The toric code loop gas can be constructed on any lattice. It is useful to choose a lattice where only three edges meet at a vertex, such as the honeycomb shown on the top. This eliminates the possibility of four lines intersecting at a single corner as shown with the square lattice as shown on the bottom.

**Fig. 28.2** A loop gas in two dimensions. Note the relationship to our loop gas diagrams discussed in section 18.1.
• **Move 1**: “Isotopy”. Meaning any smooth deformation of lines.

On the lattice:

![Diagram](image1)

Off the lattice:

![Diagram](image2)

• **Move 2**: Adding and Removing Contractable Loops.

On the lattice:

![Diagram](image3)

Off the lattice:

![Diagram](image4)

• **Move 3**: “Surgery”. Meaning reconnection of loops.

On the lattice:

![Diagram](image5)

Off the lattice:

![Diagram](image6)

**Fig. 28.3** The moves we can make by flipping over a plaquette on the lattice, and their interpretation in the continuum off the lattice. In the diagrams on the lattice, each move is achieved by flipping over the plaquette marked with $\times$. 
The allowed moves off the lattice remind us of the diagrammatic “skein”-like rules we’ve been working with, starting in chapter 2. Here we are considering planar diagrams only (no over- or under-crossings, as in chapter 12) and the rules can be summarized as

\[ \circ = 1 \]  
\[ \langle \rangle = 0 \]  

where we are assuming isotopy invariant diagrams (i.e., all smooth deformations of diagrams are allowed). In fact these diagrammatic rules are exactly the \( d = +1 \) version of the \( \mathbb{Z}_2 \) fusion rule planar diagram algebra discussed in section 15.1.1. As discussed there, the value of every valid loop diagram is unity, and this corresponds to the fact that all wavefunctions in Eq. 28.1 are added together with the same +1 coefficient.

We should note that in the definition of the wavefunction Eq. 28.1, we have not written a normalization constant as we did in Eq. 25.10. This is because in a continuum model it is hard to be precise about how many different diagrams one can write if one can deform a diagram infinitesimally to make a new diagram. One way to understand this is to accept that, while we draw a continuum diagram, we really mean that the diagrams should be understood to be on a lattice (which we don’t draw).

Another, perhaps better, way to interpret the sum in Eq. 28.1 is to think of all the terms that can be converted into each other via the three allowed moves to be an equivalence class of diagrams, and we never speak of the number of different diagrams in this class.

In the language of equivalence classes of diagrams there are exactly four equivalence class of (ground state) wavefunctions on a torus – corresponding to the evenness or oddness of the number of loops going around each cycle. Representative (or reference state) loop configurations are shown in Fig. 28.4. These correspond to the wavefunctions described in Eq. 25.11. Note that the parity of the number of loops going around a given cycle is not changed by any of the moves in Fig. 28.3.

![Fig. 28.4 Four representative (or reference state) loop diagrams corresponding to the four degenerate ground states of the toric code on a torus. Each figure represents a torus with the top and bottom edges identified and the left and right edges identified.](image)
28.1.1 Preview of Coming Attractions: Generalizations of the Toric Code

We have thus found a relation between the toric code, and the planar diagrammatic algebra that we spent so much time developing. In chapter 28.1 we will take the natural next step and, instead of using the simple $d = +1$ version of the $\mathbb{Z}_2$ planar diagram algebra, we will use more complicated diagram algebras. We will consider models where each edge can be labeled, not just with two possibilities (colored or not colored) as in the toric code, but rather with many different quantum numbers. We will then consider planar diagram algebras described by any consistent solution of the pentagon equation. Each diagram we draw in the plane can then be evaluated and turned into a number $W(\text{diagram})$, and we can then write ground state wavefunctions (or we can interpret them as error correcting codes!) in the form

$$|\psi\rangle \sim \sum_{\text{all diagrams that can be obtained from a reference diagram via allowed moves}} W(\text{diagram}) |\text{diagram}\rangle$$

(28.4)

Using this approach we will be able to generate a very wide range of topological phases of matter.

28.2 The Tube Algebra and Quasiparticle Excitations

We would now like to study the properties of the quasiparticles that occur in the toric code, but within the language of a loop gas without reference to an underlying lattice. While we know what the answers are here (we have classified all of the quasiparticles in chapter 26!) there is quite a lot to be learned by trying to develop a general technique which will work for other continuum diagrammatic models as well.

28.2.1 Operators on the Annulus

We would like to describe the quantum number of some particular region of our system — i.e., determine the net fusion product of all of the quasiparticles in the region. By our general locality principles (section 8.2 for example), we should be able to determine the total quantum number of a region only by examining a loop around that region$^1$. As such we will focus on an annulus surrounding the region of interest as shown in Fig. 28.5. We assume that there are no quasiparticles in the annular region (grey region), so that the annulus is in one of its possible ground states. However, we allow for the possibility that there are quasiparticles in the center of the annulus (inner white region of Fig. 28.5).

In Fig. 28.6 we show three different topologically equivalent representations of the annulus. The one on the right is a cylinder, or tube, which

---

$^1$For example, if we measure only single blue line emanating from the enclosed region, we know that somewhere in the enclosed region there is an end of the blue line, hence a quasiparticle defect of some sort.

---

Fig. 28.5 States of the annulus (grey region) can determine the total fusion channel of any quasiparticles surrounded by the annulus (white region).
will give the name “tube algebra” to the algebra of states we develop for the region of this annulus.

As with our discussion of states on the torus, the possible states of the annulus break up into sectors which cannot be converted into each other by the three moves given in Fig. 28.3. As in that case, it will be sufficient to describe one representative, or reference, configuration from each equivalence class.

The four equivalence classes we consider will notate as \( |n, m\rangle \) with \( n, m \in \{0, 1\} \). The meaning of this notation is that the diagram on the annulus has \( n \) lines going from the inner edge of the annulus to the outer edge, and has \( m \) lines going around the annulus. While it may appear that we need to consider other possibilities (for example lines going multiple times around the annulus), it will turn out that all diagrams can be reduced to one of these four possibilities as we will see.

Strictly speaking to describe the states on the annulus we must first describe the states on the boundary of the annulus (i.e., do any blue lines intersect the boundary of the annulus, and if so, at which points?). However, it turns out that we can simplify our lives by considering only the simplest possible boundary conditions — the case where a single quantum number (in this case either a blue line or no blue line) is emitted from each boundary at one convenient fixed point. Thus, in the toric code case, we need only consider the cases where either a single blue line, or no blue line, intersects each boundary.

No Boundary Intersections

The first case to consider is the case where no lines intersect either boundary. There are then two possible equivalence classes of states as shown in Fig. 28.7. The first, we call \( |0, 0\rangle \) meaning the empty state, the second we call \( |0, 1\rangle \), meaning there is a blue line going around the annulus.

\[
|0, 0\rangle = \begin{array}{c}
\text{annulus} \\
\end{array} = \begin{array}{c}
\text{square with sides identified} \\
\end{array} = \begin{array}{c}
\text{second state} \\
\end{array} = |0, 1\rangle
\]

\[
|0, 1\rangle = \begin{array}{c}
\text{annulus} \\
\end{array} = \begin{array}{c}
\text{same reference state drawn on a square with the orange sides identified} \\
\end{array} = \begin{array}{c}
\text{second state} \\
\end{array}
\]

Fig. 28.7 The two equivalence classes of states on the annulus where no lines intersect the boundary. The pictures on the left are the reference state. The picture in the middle is the same reference state drawn on a square with the orange sides identified. The picture on the right shows another state of the same equivalence class that can be made into the reference state by application of the moves in Fig. 28.3.

2 Qualitatively we can think of this as having grouped up all quantum numbers emitted from the boundary and fused them into a single quantum number.

3 Further, since there are no quasiparticles in the (gray) annular region, the parity of the number of blue lines is conserved going from the inner boundary to the outer boundary, so we cannot have a case where the number of blue lines intersecting the inner boundary has a different parity from the number of blue lines intersecting the outer boundary.

4 It may seem that restricting our attention to cases with only a single quantum number emitted from the edge at one point is too drastic a simplification. One (reasonably valid!) excuse we can use is that if put our system back on a lattice, our results should be independent of the particular lattice structure, so we can use an exceedingly simple lattice. In particular, we can choose to use a lattice which has only one edge pointing into the inner boundary and one edge pointing to the outer boundary (i.e., we are using a minimal “skeleton” decomposition of the annulus). A good (but not particularly easy) discussion of this type of simplification issue, along with a more rigorous argument, is given by ****.

Fig. 28.6 Three different representations of (things topologically equivalent to) the annulus. In the middle figure the two orange sides are identified with each other.
We might think about putting more than one loop around the cycle. However, we can always apply the moves from Fig. 28.3 to reduce the diagram to either zero loops or one loop around the cycle. An example of this is shown in Fig. 28.8.

![Fig. 28.8](image)

Two loops around the annulus can be converted to no loops by using the moves in Fig. 28.3. In the first step a surgery move is used at the position of the arrow. In the second step we have isotopy (smooth deformation). Finally in the last step we have loop removal.

**One Blue Line Intersecting Each Boundary**

The only other case to consider is when one blue line intersects each boundary. We will choose the intersections to be at the 6-o’clock direction (south direction) of the boundary circles. There are again two equivalence classes of diagrams, which are shown in Fig 28.9. The diagram that has a single line going in the vertical direction in the square representation of the annulus we call $|1, 0\rangle$, and the diagram which loops around both directions we call $|1, 1\rangle$. In each figure, on the right we show additional pictures in the same equivalence class, which can be reached by using moves from Fig. 28.3. To see that the double-twist figure on the far right of the first line, $|1, 0\rangle$, is the same as the figure on the far left, one needs to perform surgery at the two points marked with the arrow in the third picture on that line. Similarly to reach the figure on the far right of the second line, $|1, 1\rangle$, one performs surgery at the arrow.

![Fig. 28.9](image)

The two equivalence classes of states on the annulus where one blue line intersects each boundary. The pictures on the left are the reference state. The picture in the middle is the same reference state drawn on a square with the orange sides identified. The pictures further right shows other states of the same equivalence class that can be made into the reference state by application of the moves in Fig. 28.3. The arrows indicate places where we make surgery moves.
Despite the fact that $|1,0\rangle$ and $|1,1\rangle$ have blue lines intersecting the boundary of the annulus, this does not imply a quasiparticle at the boundary. If we revert back to the lattice, we still require that all vertices are in the low energy state with an even number blue edges coming into them, all the way up to the boundary.

### 28.2.2 Composition of States

We can define a type of multiplication, or composition, of states on the annulus which corresponds to placing one annulus inside of another, and we denote this multiplication with a $\circ$. (Here we are implicitly using the fact that we can stretch the annulus since we are considering a topological theory!). For example, if we write $|1,1\rangle \circ |1,0\rangle$ we mean we should put the $|1,1\rangle$ diagram on the inside annulus and the $|1,0\rangle$ on the outside annulus, as shown in Fig. 28.10 and we conclude that $|1,1\rangle \circ |1,0\rangle = |1,1\rangle$.

![Composition of annulus states. ($|1,0\rangle$ is put on the outside and $|1,1\rangle$ is put on the inside. The result shows ($|1,1\rangle \circ |1,0\rangle = |1,1\rangle$).](image)

It is important to note that to multiply states, they must match on their boundaries\(^5\). Both $|0,0\rangle$ and $|0,1\rangle$ (Fig. 28.7) have no blue lines intersecting the boundaries, so they can be composed with themselves or with each other. On the other hand both $|1,0\rangle$ and $|1,1\rangle$ (Fig. 28.9) have one blue line intersecting each boundary, so they can be composed with themselves or with each other. However, one cannot compose $|0,0\rangle$ or $|0,1\rangle$ with $|1,0\rangle$ or $|1,1\rangle$.

We can build up a multiplication table for this type of composition, which we write out in Table 28.1. We mention in particular the two most nontrivial results. First, $|0,1\rangle \circ |0,1\rangle = |0,0\rangle$. This is just the statement that two loops around the cycle can be annihilated to no loops as in Fig. 28.8. The second, $|1,1\rangle \circ |1,1\rangle = |1,0\rangle$ is a bit more complicated. We can see this in several ways: if we compose the counterclockwise directed wrapping shown the furthest left in the lower line of Fig. 28.9 with the clockwise directed wrapping furthest right we obviously get a result with no net wrapping which is $|1,0\rangle$. Another way to see this is to compose two wrappings in the same direction, which will give a diagram like the far right of the top line of Fig. 28.9 which is also $|1,0\rangle$.

A way to summarize the multiplication table is via the formula\(^6\)

\[
\begin{array}{c|cccc}
\circ & |0,0\rangle & |0,1\rangle & |1,0\rangle & |1,1\rangle \\
|0,0\rangle & |0,0\rangle & |0,1\rangle & • & • \\
|0,1\rangle & |0,1\rangle & |0,0\rangle & • & • \\
|1,0\rangle & • & • & |1,0\rangle & |1,1\rangle \\
|1,1\rangle & • & • & |1,1\rangle & |1,0\rangle \\
\end{array}
\]

Table 28.1 The multiplication table for composing annuli for the toric code. The • indicates a composition which is not allowed due to non-matching boundary conditions. This table is summarized by Eq. 28.10.

\(^5\)Strictly speaking we need to sum over all possible boundary conditions that can occur at the boundaries that have been joined (matching the boundary condition from the inner to the outer annulus). However, the excuse used in note \(^3\) above allows us to ignore this complication and just work with the one reference configuration.

\(^6\)The Kronecker delta assures that we only compose states that are compatible on their boundaries.
Abstracting the Toric Code: Introduction to Tube Algebra

\[ |n, m\rangle \circ |n', p\rangle = \delta_{n,n'} \ |n, (m + p) \text{mod} 2\rangle \]  \hspace{1cm} (28.5)

It is worth quickly mentioning how compositions look if we work with rectangular diagrams or tube diagrams. Both such diagrams are simply stacked on top of each other to create new diagrams. An example of composing rectangular diagrams and composing tube diagrams is given in Fig. 28.11.

Fig. 28.11 The composition \(|1, 1\rangle \circ |1, 0\rangle = |1, 1\rangle\), as in Fig. 28.10 but shown using rectangle notation (top) where the orange edges are identified, or using tube notation (bottom).

The algebra associated with this type of composition of diagrams is known as Ocneanu’s Tube Algebra, and although the multiplication is fairly trivial in the case of the toric code that we are considering here, it generalizes to much more complex algebras as we will see later in chapters **.

28.2.3 Quasiparticle Projectors

Our intent is to develop a way to determine the total quantum number of the quasiparticles enclosed in region. The way we now do this is to use our tube algebra to form orthogonal projectors \(O_a\) — where we define one projector corresponding to each quasiparticle type \(a\). The physical interpretation is the following. We imagine starting with a disk which contains some overall quasiparticles whose total quantum number is \(a\). If we surround this region with our annular projector \(O_b\), we want the projector to return zero if \(b \neq a\), and if \(b = a\) then it returns the same wavefunction on a slightly larger disk — i.e., leaves the system topologically unchanged (same total quantum number, and still topologically a disc). I.e., \(O_a\) has a value of 1 or 0 depending on whether we are surrounding a region with quantum number \(a\) or not\(^7\). To define these projectors we want to find linear combinations of our

\(^7\) Here we are defining operators which add a small annulus outside of a disk. I.e., they take as an input a wavefunction on a disk and give an output a wavefunction on a slightly larger disk. It is sometimes useful to define a different operators which act on an annulus in (perhaps the bulk of) an existing system not near any edge, without adding any new region. We would want such operators to project to a particular quasiparticle type inside the region. I.e., the operator would give unity if it detects a particular type of quasiparticle inside the region and give zero otherwise. We can imagine such operators as being \(\sum_k |k\rangle\langle k|\) where the sum over state is only over states in a given topological class. We will not work with such operators here because it simply adds a bit more notational complexity.
annular states \((n, m)\) such that
\[
O_a \circ O_b = \delta_{ab} O_a
\]  

(28.6)

Recall, however, that the space of states we are considering on the annulus breaks up into sectors with different boundary conditions, we cannot even talk about multiplying states together with \(\circ\) unless the boundary conditions match (i.e, \(|n, m\rangle\) can only multiply \(|a, b\rangle\) if \(a = n\)).

For the case of the toric code, as discussed above in section 28.2.1, there are two classes of boundary conditions, so we want to find a set of orthogonal projectors within each boundary condition class. Given the multiplication table \(28.1\) it is easy to construct these projectors. In the class where no blue lines intersect the boundary we have\(^8\)
\[
O_I = \frac{1}{2} (|0,0\rangle + |0,1\rangle) = \frac{1}{2} \begin{pmatrix} \bigcirc & \bigcirc \\ \bigcirc & \bigcirc \end{pmatrix} \quad (28.7)
\]
\[
O_p = \frac{1}{2} (|0,0\rangle - |0,1\rangle) = \frac{1}{2} \begin{pmatrix} \bigcirc & \bigcirc \\ \bigcirc & \bigcirc \end{pmatrix} \quad (28.8)
\]

and in the class where one blue line intersects the inner boundary and one blue line intersects the outer boundary, we have
\[
O_v = \frac{1}{2} (|1,0\rangle + |1,1\rangle) = \frac{1}{2} \begin{pmatrix} \bigcirc & \bigcirc \\ \bigcirc & \bigcirc \end{pmatrix} \quad (28.9)
\]
\[
O_f = \frac{1}{2} (|1,0\rangle - |1,1\rangle) = \frac{1}{2} \begin{pmatrix} \bigcirc & \bigcirc \\ \bigcirc & \bigcirc \end{pmatrix} \quad (28.10)
\]

It is easy enough to use the multiplication table \(28.1\) to check that these satisfy Eq. \(28.6\).

Here, with a bit of foresight, we have labeled these projectors \(I, p, v, f\) corresponding to their particle types (identity, plaquette defect, vertex defect, and fermion). Let us think a bit about how we make this connection between projectors and particle types.

First, we realize that the vertex defect \(v\) and the \(f\) particle both involve ends of blue strings (i.e., defects of the vertex term of the Hamiltonian) so if one wraps an annulus around one of these particles, there will be an (odd number) of blue strings going through the annulus. Thus the \(v\) and \(f\) projectors should be made from the \(|1,0\rangle\) and \(|1,1\rangle\) states on the annulus. Analogously, the ground state \(I\) and the plaquette \(p\) defect involve no vertex defects and therefore should be made from the \(|0,0\rangle\) and \(|0,1\rangle\) states.

Let us start with the \(|0,0\rangle\) and \(|0,1\rangle\) states — those without blue lines touching the boundary. If we think about a single plaquette, we know from section \(25.3\) that the ground state is always a superposition of flipped and unflipped plaquettes with a plus sign between the two pieces. In fact, it is easy to argue (by considering groups of plaquettes) that the ground state should be a superposition (with a plus sign) of flipped and unflipped regions. Thus the positive superposition of \(|0,0\rangle\) (no loops) and \(|0,1\rangle\) (one loop) must be a projector onto the ground state space —
the space with no quasiparticles. This is what we called $O_I$ in Eq. \ref{eq:28.7}
In the language of our diagrammatic algebra (the $d = +1$ version of the $\mathbb{Z}_2$ fusion rules from section 18.1.1), the operation of doing nothing to the region plus the operation of putting a blue line around the region is the same as putting a $\Omega$-loop around the region (See section 17.5 and Fig. 17.10) Here $D = \sqrt{2}$ since there are two particle types in $\mathbb{Z}_2$ fusion rules: the trivial particle and the nontrivial particle.)

If we return to the lattice Hamiltonian Eq. \ref{eq:26.1} we could also make the plaquette term of the Hamiltonian look like a $\Omega$-loop in the following way. Let us add a constant to the the plaquette term in the Hamiltonian Eq. \ref{eq:26.1} (which doesn’t change the ground state, just the ground state energy)

$$P_\beta \rightarrow 1 + P_\beta$$

and then we realize that this combination is also just an $\Omega$-loop around a plaquette:

$$1 + P_\beta = \text{unflipped} + \text{flipped} \sim \Omega\text{-loop} \sim O_I$$

The principle that the ground state projector, and similarly the plaquette term in the Hamiltonian, is just an $\Omega$-loop is something that will hold for all of the generalized loop gas models we will meet in chapters ***-*** as well.

Once we have determined the projector $O_I$ the other orthogonal projector within the space of $|0,0\rangle$ and $|0,1\rangle$ is fully defined and is given by Eq. \ref{eq:28.8}. However the plaquette defect projector $O_p$ can also be understood in a way similar to the argument for $O_I$. Recall from section 25.4.2 that the defect on a plaquette is the difference of flipped and unflipped. By grouping together plaquettes, it is easy to see that the annular projector around a region with a magnetic quasiparticle must be a superposition of $|0,0\rangle$ (no loops) and $|0,1\rangle$ (one loop) with a minus sign.

We now turn to the states with blue lines touching the boundaries — i.e., the $|1,0\rangle$ and $|1,1\rangle$ states. It may look a bit unexpected that the vertex defect projector $O_v$ needs to be a superposition of the $|1,0\rangle$ diagram and the $|1,1\rangle$ diagram as shown in Eq. \ref{eq:28.9} However, we should think of the quasiparticle that we call the vertex defect, not only being the vertex defect, but rather a vertex defect with the absence of a plaquette defect, whereas the $f$ particle is the vertex defect in the presence of a plaquette defect. The superposition of the two diagrams $|1,0\rangle$ and $|1,1\rangle$ is arranged to assure that the $O_v$ particle does not include a plaquette defect whereas the $O_I$ particle does. To see how this works, note that the $|1,0\rangle$ state can be created from the $|0,0\rangle$ state (no blue lines) by flipping over a string of spins connecting the inner and outer annulus without going around the cycle, i.e., by adding the one vertical blue line. If we perform the same operation on the $|0,1\rangle$ state, as shown in Fig. 28.12 we obtain the $|1,1\rangle$ state — a line that connects the two annuli and also goes around the cycle. Thus the superposition $O_v$ is the same as the superposition $O_I$ with the one string added between
annuli, whereas the superposition $O_f$ is the same as $O_p$ with the one string added.

It is worth recalling that the $d = +1$ version $\mathbb{Z}_2$ planar diagram algebra (the diagram algebra we are using here!) has two possible solutions to the hexagon equation as discussed in section 18.1.1 boson and fermion (neither solution is modular). It is not a coincidence that the particle types we have found arising from this diagrammatic construction are also bosons and fermions. We will see that this is a general principle: when we build a model based on a planar diagrammatic algebra, the particle types that arise must include all of the possible solutions to the hexagon equation for that planar algebra.

**Quasiparticle Basis for Torus States**

It is worth noting that once we have our basis for quasiparticles enclosed in an annulus, we can also glue the two boundaries of the annulus to each other to form a closed torus, and we have a basis for the degenerate ground state of the torus. This basis is a rather special basis since if you cut the handle of the torus, the boundary that you reveal has the quantum number of the given quasiparticle type. Recall that in the discussion of section 7.3.1 we defined exactly this basis on the surface of a torus by stating that a quasiparticle of a given type is running around the torus handle.

### 28.3 Twists and $T$-matrix

With the tube algebra one can determine the twist factor $\theta$ for each particle type. To do this, we imagine deforming the local environment of the particle around in circle by 360 degrees as depicted in the top of Fig. 28.13. This is equivalent to the twist operation shown in the lower part of Fig. 28.13.

Twist deforming either the $\phi$ state or the $h$ state leaves the state unchanged, so we trivially have $\hat{\theta}O_I = O_I$ and $\hat{\theta}O_p = O_p$ implying trivial twist factors for the $I$ particle and the plaquette defect particle, $\theta_I = \theta_p = 1$. I.e., these are both bosons, as we expected.

However, twist deforming $|1,0\rangle$ and $|1,1\rangle$ is nontrivial as shown in Fig. 28.14. These diagrams tell us that $\hat{\theta}|1,0\rangle = |1,1\rangle$ and $\hat{\theta}|1,1\rangle = |1,0\rangle$.

Note that a general formula for the twist operator on the states $(n,m)$ is given by

$$\hat{\theta}|n,m\rangle = |n,(n+m)\text{mod }2\rangle$$  \hspace{1cm} (28.11)

Using Eq. 28.9 we obtain $\hat{\theta}O_v = O_v$ confirming that $\theta_v = 1$, i.e., the vertex defect particle is a boson. Similarly using Eq. 28.10 we obtain $\hat{\theta}O_f = -O_f$ confirming that $\theta_f = -1$, the $f$ particle is a fermion. Note that it is a nontrivial result that the particle projectors we have found are eigenstates of the twist operator. It isn’t obvious that this should have to be true, although we certainly suspected it should be true considering our understanding of the structure of anyon theories!
If we connect the inside of the annulus to the outside to form a torus, as discussed at the end of section 28.2.3 we realize that the twist transformation is exactly the $T$ transformation discussed in section 17.3.2 (see Fig. 17.6) or Dehn twist as discussed in section 7.4.

### 28.4 $S$-matrix

We can also derive the modular $S$ matrix diagrammatically by exchanging the two directions of the torus as discussed in section 17.3.2 and 7.3.1 (see Fig. 17.6).

Let us abuse notation a bit and denote by $|0,0\rangle$, $|0,1\rangle$, $|1,0\rangle$, $|1,1\rangle$ the four basis states obtained from the annular states $|0,0\rangle$, $|0,1\rangle$, $|1,0\rangle$, $|1,1\rangle$ by connecting the inner boundary to the outer boundary to obtain a torus. Exchanging the two directions on the torus (making an $S$ transform as in section 17.3.2) we have transform on basis states for our torus

\[
\begin{align*}
S(0,0) &= \begin{array}{c}
\text{[\ ]}
\end{array} = \begin{array}{c}
\text{[\ ]}
\end{array} = |0,0\rangle \\
S(0,1) &= \begin{array}{c}
\text{[\ ]}
\end{array} = \begin{array}{c}
\text{[\ ]}
\end{array} = |1,0\rangle \\
S(1,0) &= \begin{array}{c}
\text{[\ ]}
\end{array} = \begin{array}{c}
\text{[\ ]}
\end{array} = |1,0\rangle \\
S(1,1) &= \begin{array}{c}
\text{[\ ]}
\end{array} = \begin{array}{c}
\text{[\ ]}
\end{array} = |1,1\rangle
\end{align*}
\]
surgery for the latter equality between diagrams. We can write this transformation as a matrix defined as (with \( n, m = 0, 1 \))

\[
S |n, m\rangle = |m, n\rangle
\]  

(28.16)

since the indices \( n \) and \( m \) refer to the two different directions, which are switched by \( S \).

We would like to change this basis into the quasiparticle basis obtained by connecting the inner to outer boundary of the annulus for the four quasiparticle projectors \( O_I, O_p, O_v, O_f \). Making this change of basis as specified\(^{10}\) in Eqs. 28.7-28.10 then applying \( S \) then changing basis back now need \( 1/\sqrt{2} \) out front rather than \( 1/2 \) out front to be normalized as wavefunctions rather than the normalized projectors that we have been using.

### 28.5 Direct Calculation of Braiding and Fusion

A more direct calculation of the braiding properties of the quasiparticles, the \( R \)-matrix, is possible by considering a more complex geometry, a disk with two holes. With this geometry, we add strings attached to holes and around holes as in Eqs. 28.7-28.10. To calculate braiding, we need to drag one hole around the other. As an example here we will show that exchanging two \( f \) particles gives a minus sign. Since each \( f \) is a superposition of two terms (\(|1, 0\rangle\) and \(|1, 1\rangle\) as in Eq. 28.10), our system with two \( f \) particles (one in each hole) is now a sum of four terms, as shown in the top line of Fig. 28.15.

\[
\frac{1}{4} \left[ \begin{array}{c}
\begin{array}{c}
\begin{array}{c}
\dot{\circ}
\end{array}
\end{array} & - & \begin{array}{c}
\begin{array}{c}
\begin{array}{c}
\dot{\circ}
\end{array}
\end{array} & - & \begin{array}{c}
\begin{array}{c}
\begin{array}{c}
\circ
\end{array}
\end{array} & + & \begin{array}{c}
\begin{array}{c}
\begin{array}{c}
\circ
\end{array}
\end{array}
\end{array}
\end{array}
\end{array}
\end{array}
\right]
\]

\[
\downarrow \text{Exchange}
\]

\[
\frac{1}{4} \left[ \begin{array}{c}
\begin{array}{c}
\begin{array}{c}
\dot{\circ}
\end{array}
\end{array} & - & \begin{array}{c}
\begin{array}{c}
\begin{array}{c}
\dot{\circ}
\end{array}
\end{array} & - & \begin{array}{c}
\begin{array}{c}
\begin{array}{c}
\circ
\end{array}
\end{array} & + & \begin{array}{c}
\begin{array}{c}
\begin{array}{c}
\circ
\end{array}
\end{array}
\end{array}
\end{array}
\end{array}
\end{array}
\right]
\]

Fig. 28.15 Calculation that the phase accumulated from exchanging two \( f \) particles is \(-1\). The upper line represents two \( f \) quasiparticles, one in each hole (Compare Eq. 28.10). The holes are exchanged, braiding the blue lines in the process. Using surgery and loop addition/removal (the moves of Fig. 28.3), one can show the lower line is precisely minus the top line showing an exchange phase of \(-1\). For example, performing surgery on the first picture in the second line at the location of the arrow gives exactly the second picture on the top line, but missing the minus sign.

We then braid the holes around each other, stretching the blue lines in the process as shown in the lower line of Fig. 28.15. The resulting picture can then be related back to the top line of the figure using surgery and loop addition/removal (the moves of Fig. 28.3). It is an easy exercise to show that the lower line is exactly minus the upper line of the figure.
thus showing that the phase obtained in exchanging two $f$ particles is
$-1$. (See exercise 28.2.a)

Fusion relations can also be calculated graphically with our diagrammatic rules by essentially merging together the two holes. This is just
another way of saying that we draw an imaginary circle around both
holes and treat this imaginary circle as a single, bigger, hole. An exam-
ple of this is shown in Fig. 28.16 where we show the fusion of a vertex
defect (left) with an $f$ particle (right) giving a plaquette defect. The top
line of this figure is the four term superposition describing the vertex
defect and $f$ particle (see Eq. 28.9 and 28.10). Going to the second line
we have used the rules (Fig. 28.3) of our diagrammatic algebra. We th en
realize that we can write this second line as a plaquette projector $O_p$
around “something” — which guarantees that the total quantum num-
ber inside the large hole in the third line is simply that of the plaquette
defect.

$$\frac{1}{4} \left[ \begin{array}{ccc}
\text{something} & - & \text{something} \\
\text{something} & - & \text{something}
\end{array} \right]$$

Fig. 28.16 Calculation that (vertex defect) $\times f = \text{(plaquette defect)}$. The upper line
represents a vertex defect (left) quasiparticle and an $f$ (right) quasiparticle (Compare
Eq. 28.9 and 28.10). In the lower line the diagrammatic moves are used. The small
blue string at the bottom is pulled off the bottom of the picture and the remaining
diagram at the bottom is a plaquette projector $O_p$ around the inner region.

28.6 Generalization to $\mathbb{Z}_N$ Toric Code (Briefly)

The diagrammatic arguments we have used all generalize very naturally
to the case of the $\mathbb{Z}_N$ toric code discussed in sections 25.6 and 26.5.
Instead of having simple loop diagrams we now have a $\mathbb{Z}_N$ fusion al-
gebra. Our diagrammatic algebra now requires us to draw arrows on lines
(for $N > 2$) and label each line with an integer which is interpreted
modulo $N$. Reversing an arrow on a line takes a label $j$ to $N - j$ as
shown in Fig. 28.17. The label 0 plays the role of the identity, and such
lines can be added or removed from the diagram freely as in our other
diagrammatic algebras.

Vertices must follow the rule that the $Q_\alpha$ at each vertex, which is the
sum of label on incoming arrows minus the sum of labels on outgoing
arrows, must be zero modulo $N$. An example of such a diagram is such
Generalization to $\mathbb{Z}_N$ Toric Code (Briefly)

A diagram is shown in Fig. 28.18. Analogous to the three allowed moves in Fig. 28.3, we are allowed three moves in order to manipulate diagrams into equivalence classes. These moves are shown in Fig. 28.19.

- **Move 1:** “Isotopy”. Meaning any smooth deformation of lines.

- **Move 2:** Adding and Removing Loops. A loop with any label can be added or removed.

- **Move 3:** Reconnecting (trivial $F$) move.

The first move is similar to what we had in the case of the toric code. The second move, that we can add or remove loops, is also similar to the case of the toric code, except that the loops have arrows as well as labels. The third move is “new” but should look familiar from our discussion of $F$-moves. However, here the $F$-coefficient is simply unity — and there is no sum over states on the right hand side (compare to Fig. 16.3). This type of $F$-matrix (which is not a matrix but simply a scalar) is what we called the trivial cocycle for the $\mathbb{Z}_N$ fusion algebra in sections 20.1.1 and 20.4.

We would now like to build basis states for our tube algebra analogous to what we did for in section 28.2. Let us define the tube state $|n, m\rangle$ to be the state with quantum number $n$ going between the inner and outer edge of the annulus, and quantum number $m$ going counterclockwise around the annulus, as shown in Fig. 28.20.

Composition of these basis states is performed by placing one annulus state inside the other as in section 28.2.2. The multiplication law is the natural generalization of Eq. 28.5

$$|n, m\rangle \circ |n', p\rangle = \delta_{n,n'} |n, (m+p) \text{mod} N\rangle$$

which we illustrate graphically in Fig. 28.21 (See exercise 28.4a). We
Abstracting the Toric Code: Introduction to Tube Algebra

\[ |n, m\rangle = m_n + m = n + m \]

Fig. 28.20 Basis states for the tube algebra of the \( \mathbb{Z}_N \) toric code. The quantities \( n, m, \) and \( n + m \) are interpreted mod \( N \). In the far right the two orange sides are identified.

\[ n \circ n' = \delta_{n,n'} \]

Fig. 28.21 Graphical depiction of the composition rule Eq. [28.17] All of the indices \( n, n', m, p, (m + p) \) are interpreted mod \( N \).

can then define \( N^2 \) projectors

\[ O^k_n = \frac{1}{N} \sum_{p=0}^{N-1} e^{2\pi ipk/N} |n,p\rangle \] (28.18)

which satisfy (See exercise 28.4.b)

\[ O^k_n \circ O^q_{n'} = \delta_{n,n'} \delta_{k,q} O^k_n \] (28.19)

as desired in Eq. [28.6]. These orthogonal projectors describe the quasiparticle types of the theory. We expect that the lower index, \( n \), which gives the total “charge” entering the annulus should match the \( Q_\alpha \) quantum number of the vertex operator for the \( \mathbb{Z}_N \) toric code that we discussed in section 26.5, whereas the upper index \( k \) should describe the \( e^{2\pi ik/N} \) eigenvalue of the \( A_\beta \) plaquette operator. In other words, we are describing the quasiparticle \( [n, k] \) from Eq. [26.8]

\[ [n, k] \leftrightarrow O^k_n \]

We can apply the twist operator to the basis states to obtain the analog of Eq. [28.11]

\[ \hat{\theta} |n, m\rangle = |n, (n + m) \text{mod} N \rangle \] (28.20)

which we show graphically in Fig. 28.22 (See exercise Eq. 28.4). This is the same twist transformation as shown in section 28.3 but using the rectangular representation of an annulus. See also Fig. 17.6.
28.6 Generalization to $\mathbb{Z}_N$ Toric Code (Briefly) 385

Applying the twist operator Eq. 28.20 to the quasiparticle projector $O_n^k$ in Eq. 28.18 we obtain

$$\hat{\theta} O_n^k = \frac{1}{N} \sum_p e^{2\pi ipk/N} \hat{\theta} |n, p\rangle = \frac{1}{N} \sum_p e^{2\pi ipk/N} |n, (p + n) \mathrm{mod} N\rangle$$

$$= e^{-2\pi ink/N \frac{1}{N} \sum_q e^{2\pi ikg/N} |k, q\rangle} = e^{-2\pi ink/N} O_n^k$$

Giving us a twist factor

$$\theta_{n,k} = e^{-2\pi ink/N}$$

In agreement with Eq. 26.10.

Finally, let us evaluate the $S$-matrix. Again our strategy is to exchange the two directions of the torus as discussed in section 17.3.2 and 7.3.1 (see Fig. 17.6). First, we should connect the inner and outer edges of the annulus in Fig. ?? to make a torus. By doing this we construct the basis states on a torus $|n, m\rangle$ that correspond to $|n, m\rangle$ on the annulus. The $S$ matrix applied to our basis states simply rotates the directions of the torus giving

$$S |n, p\rangle = |p, -n\rangle$$

(28.21)

as shown graphically in Fig. 28.23.

$$S$$

Fig. 28.23 The $S$ operation rotates the torus. Note in the second step we use a single $F$ move then reverse the quantum number on $n$ by reversing the arrow. All indices $n, p, -n$ are interpreted modulo $N$.

We then construct the quasiparticle superpositions on the torus corresponding to the projectors Eq. 28.18

$$| [n, k] \rangle = \frac{1}{\sqrt{N}} \sum_{p=0}^{N-1} e^{2\pi ipk/N} |n, p\rangle$$

(28.22)
Note the change in normalization of this wavefunction compared to the projector normalization used in Eq. 28.18. This is to assure the orthonormality
\[ \langle [n, k] | [n, l] \rangle = \delta_{k,l} \]

The S-matrix is then given by
\[
S_{[n',k'],[n,k]} = \langle [n', k'] | S | [n, k] \rangle = \frac{1}{N} \sum_{p,p'=0}^{N-1} e^{2\pi i (pk' - p'k) / N} \langle n', p' | S | n, p \rangle
\]
\[
\frac{1}{N} \sum_{p,p'=0}^{N-1} e^{2\pi i (pk' - p'k) / N} \langle n', p' | -n \rangle
\]
\[
\frac{1}{N} e^{2\pi i (n'k + nk') / N}
\]
which matches Eq. 26.11.

Exercises

Exercise 28.1 Toric Code S-matrix
Derive the S-matrix of the toric code (Eq. 26.5) by using the method described in section 28.4.11 (If you need help with this calculation look at the more general calculation given in section 28.6.)

Exercise 28.2 Braiding Quasiparticles in Toric Code Loop Gas
(a) Use the technique of section 28.5 to show that exchanging two f’s gives a minus sign (i.e., confirm the details of the argument given there).
(b) Use similar techniques to show that exchanging two e particles gives no sign and exchanging two m particles gives no sign.
(c) Show that braiding an e particle or an f particle all the way around an m particle gives a minus sign but braiding around the identity gives no sign.
(d) Show that braiding e all the way around f gives a minus sign.

Exercise 28.3 Fusing Quasiparticles in Toric Code Loop Gas
Use the technique of section 28.5 to deduce the full fusion table for the toric code.

Exercise 28.4 \( \mathbb{Z}_N \) Tube Algebra
(a) Derive the final equality in Fig. 28.21 from the three allowed moves in Fig. 28.19. You will also need to add and remove lines labeled with the identity (which is 0 here).
(b) Confirm Eq. 28.19
(c) Derive the final equality in Fig. 28.22
Part VI

More General Loop Gas and String Net Models
Kitaev Quantum Double Model

We have already seen the generalization of the conventional toric code to the more interesting $\mathbb{Z}_N$ toric code. In the next few chapters we will consider models that generalize the toric code further and in other ways. These generalizations all share many key properties: In short, we start with some planar diagram algebra and use that to build a Hamiltonian, which in turn produces a TQFT as its ground state known as the quantum double or Drinfel’d double of the input original diagram algebra. All of the calculations will hinge on diagrammatic manipulations.

29.1 Defining the Model

Perhaps the simplest generalization of the toric code is the generalization of the upspin/downspin (black/blue) assigned to each edge to group valued variables. Indeed, the $\mathbb{Z}_N$ toric code was just such a generalization where the group we used was the abelian group $\mathbb{Z}_N$ (integers mod $N$ under addition). This generalization to groups, which turns out to be essentially equivalent to a lattice gauge theory as we will discuss in section 29.7, is sometimes known as the Kitaev Model\(^1\) or more specifically the Kitaev Quantum Double Model\(^2\).

We start by choosing a group $G$ with elements $g \in G$. This group may be abelian or nonabelian. For this model we can work with any regular or irregular lattice\(^3\) like the one shown in Fig. 29.1 and we put arrows on each edge (we can fix them in arbitrary directions). Instead of labeling each edge with spin-up or spin-down we assign each edge a group element $g$ as shown in the figure. The set of all possible group elements assigned to all possible edges serves as an orthonormal and complete basis for the Hilbert space.

\(^1\)The phrase Kitaev Model is not sufficiently specific, since Kitaev has introduced many models and several very different models all go by this same name.

\(^2\)Caution: The presentation I give here is on the dual lattice compared to Kitaev [1997]. Be warned that almost all presentations of this material follow Kitaev’s original work, except us. We will switch to the same presentation as Kitaev in section 29.7. I have chosen this approach so as to emphasize the importance of the planar diagram algebra.

\(^3\)....which should not be called a lattice.

Fig. 29.1 Each edge of the (irregular) lattice is labeled with a group element $g \in G$. 

Medium Material
We make a definition that reversing the arrow on an edge inverts the element of the group, as shown in Fig. 29.2.

29.1.1 Vertex and Plaquette Operators

We can now define vertex operators \( \hat{V}_\alpha \) at each vertex \( \alpha \) as follows:

\[
\hat{V}_\alpha = \begin{bmatrix} g_1 & g_2 & g_3 & g_4 & e \\ g_1^{-1} & g_2^{-1} & g_3^{-1} & g_4^{-1} & e \end{bmatrix}
\]

(29.1)

In other words, to apply \( \hat{V}_\alpha \) at a vertex \( \alpha \) we orient all of the arrows on the edges so they point away from the vertex (using Fig. 29.2 if necessary). Then we multiply together the group elements on all of the edges of the vertex in a clockwise order around the vertex (independent of how many edges intersect the vertex\(^4\)). If the result gives the identity element \( e \) of the group, then \( \hat{V}_\alpha \) gives unity, otherwise the operator gives zero.

A few things to note about \( \hat{V}_\alpha \). First of all, it does not matter where we start multiplying around the vertex, since \( g_1 g_2 g_3 = e \) implies \( g_2 g_3 g_1 = g_3 g_1 g_2 = e \). So in essence the operator is rotationally invariant. Secondly, we note that \( \hat{V}_\alpha \) is a projector, meaning

\[
\hat{V}_\alpha^2 = \hat{V}_\alpha
\]

(29.2)

For each group element \( h \in G \) we now define an operator \( \hat{P}_\beta(h) \) on a plaquette as follows:

\[
\hat{P}_\beta(h) = \begin{bmatrix} g_3 & g_4 & g_5 \\ g_2 & g_1 & h^{-1}g_3 \\ h^{-1}g_4 & h^{-1}g_5 \\ h^{-1}g_1 & h^{-1}g_2 \\ h^{-1}g_5 & h^{-1}g_4 \\ h^{-1}g_2 & h^{-1}g_1 \\ h^{-1}g_5 & h^{-1}g_4 \\ h^{-1}g_2 & h^{-1}g_1 \\ h^{-1}g_5 & h^{-1}g_4 \\ h^{-1}g_2 & h^{-1}g_1 \end{bmatrix}
\]

(29.3)

In other words, we first orient the arrows around the plaquette in a counterclockwise manner (using Fig. 29.2 if necessary). The action of \( \hat{P}(h) \) is to premultiply each edge label by the group element \( h \).

\[
\hat{P}_\beta(h')\hat{P}_\beta(h) = \hat{P}_\beta(h'h)
\]

(29.4)

I.e., if you premultiply by \( h \) then premultiply by \( h' \) this is the same as premultiplying by \( h'h \).

We then construct the total plaquette operator for the plaquette \( \beta \) as

\[
\hat{P}_\beta = \frac{1}{|G|} \sum_{h \in G} \hat{P}_\beta(h)
\]

(29.5)

where \( |G| \) is the number of elements in the group \( G \).
From Eq. 29.4 and Eq. 29.5 it is easy to show that
\[ \hat{P}_\beta \hat{P}_\beta(h) = \hat{P}_\beta(h) \hat{P}_\beta = \hat{P}_\beta \]
and from this we show that \( \hat{P}_\beta \) is a projector, meaning
\[ \hat{P}_\beta^2 = \hat{P}_\beta \]
so that it has eigenvalues 0 and 1 only.

**Commuting Operators**

It is worth noting that, analogous to the case of the toric code, the operators \( \hat{V}_\alpha \) and \( \hat{P}_\beta \) all commute with each other. It should be fairly obvious that all of the \( \hat{V}_\alpha \) operators commute with each other. It is perhaps a bit less obvious that the \( \hat{P}_\beta \) operators commute with each other when they share an edge as in Fig. 29.3. The key here is to realize that the edge arrow needs to be reversed to apply the plaquette operator on the two different plaquettes. In the figure, the arrow on the shared edge is oriented counterclockwise with respect to \( \beta \) so that the \( \hat{P}_\beta(h) \) operator may be applied on plaquette \( \beta \), turning the shared edge from \( g \) to \( hg \). To apply the plaquette operator \( \hat{P}_\beta(h') \) on plaquette \( \beta' \) we need to first flip the edge arrow, turning \( g \) to \( g^{-1} \). Then applying \( \hat{P}_\beta(h') \) turns the edge into \( h'g^{-1} \), and we can then reverse the arrow again to put it in the original position giving the edge value \( (h'g^{-1})^{-1} = g(h')^{-1} \). Thus one of the plaquette operators premultiplies the edge element and the other plaquette operator postmultiplies the edge element — and hence these commute with each other.

It is also not immediately obvious that the the plaquette operators commute with the adjacent vertex operators. Consider Fig. 29.4 To apply the vertex operator at \( \alpha \), we must first reverse the arrow on the edge labeled \( g_1 \) so all edges point away from the vertex. With the reversed arrow, the edge is then labeled \( g_1^{-1} \). The vertex operator \( \hat{V}_\alpha \) now measures the product \( g_1^{-1}g_2g_3 \). Now instead consider applying the plaquette operator \( \hat{P}_\beta(h) \) on the plaquette \( \beta \) before reversing the direction of the arrow. This premultiplies both \( g_1 \) and \( g_2 \) by \( h \), yielding \( hg_1 \) and \( hg_2 \) with the arrows oriented as they are in the Figure. If we try applying the vertex operator after the plaquette operator we would now measure \( (hg_1^{-1})^{-1}(hg_2)g_3 = (g_1^{-1}h^{-1})(hg_2)g_3 = g_1^{-1}g_2g_3 \) which is unchanged. Thus we conclude that the plaquette and vertex operators commute.

### 29.1.2 Code Space

We can use the Kitaev model as a quantum memory (a error correcting code) analogous to the toric code. For a group with \( |G| \) elements, one need to have a \( |G| \)-state quantum system\(^6\) on each edge, instead of just having a single spin-\( \frac{1}{2} \) two-state system on each edge as in the conventional toric code.

Similar to the toric code, the code space for the Kitaev model will

---

\(^6\)A qubit is a two state quantum system. Sometimes people call a three state quantum system a “qutrit”. For a \( d \)-state quantum system one often says one has a “qudit”.
be the set of states where all $V_\alpha = +1$ and all $P_\beta = +1$. This code space, analogous to the toric code, will turn out to be closely related to a TQFT.

The Kitaev model built from a non-abelian group $G$ has a significant advantage over the conventional toric code: Not only can one store quantum information, but by braiding, fusing, and measuring defects one can also perform computation (See Kitaev [1997]; Preskill [2004], for example).

It is interesting to note that if one builds the Kitaev model from a group $G_1$ and then another Kitaev model from a group $G_2$, the product of the code spaces of these two will be the code space of a Kitaev model built from a group $G_1 \times G_2$.

**Abelian Case: Relation to the $Z_N$ Toric Code**

The $Z_N$ toric code is a special case of the Kitaev model\(^7\) where we take the group to be $G = Z_N$ (the integers modulo $N$ with addition as the group operation). As discussed in section 41.2, any abelian groups can be written as $G = Z_{N_1} \times Z_{N_2} \times \ldots \times Z_{N_p}$ for some number of factors $p$. Thus all Kitaev models built from abelian groups are equivalent to a product of some number of some ($Z_N$) toric codes.

\(29.1.3\) **Hamiltonian**

Instead of thinking about the Kitaev model as an error correcting code, we can think of it as a phase of matter analogous to our discussion in chapter 26. We can now write simple Hamiltonian for our system

$$H_{\text{Kitaev model}} = -\Delta_v \sum_{\text{vertices } \alpha} \hat{V}_\alpha - \Delta_p \sum_{\text{plaquettes } \beta} \hat{P}_\beta$$

(29.8)

where here $\Delta_v > 0$ and $\Delta_p > 0$. The ground state space will thus be the space where every vertex operator $\hat{V}_\alpha$ and every plaquette operator $\hat{P}_\beta$ is in the $+1$ eigenstate. I.e., the ground state is the code space. Any vertex or plaquette in the other eigenstate (having eigenvalue 0) then corresponds to an excitation, and has energy cost $\Delta_v$ or $\Delta_p$ respectively.

The Hamiltonian Eq. (29.8) is precisely the Hamiltonian we used for the $Z_N$ toric code in Eq. (26.6) previously.

**29.2 How Kitaev Model Generalizes Toric Code**

In the case of the toric code, the condition of being in the lower energy eigenstate of all of the vertex operators (section 25.3) was interpreted as allowing only loop configurations (See also section ??). We can think of the blue lines as carrying some $Z_2$ valued in-plane “flux” that is conserved at each vertex: if a blue line comes into the vertex another blue line must go out of the vertex. This was generalized for the $Z_N$ toric code.
code where we think of the edges as carrying $\mathbb{Z}_N$ flux: edges have arrows and integer (modulo $N$) on them, and the total of the arrow pointing in minus out should add up to zero modulo $N$. Now, in the more general Kitaev model based on a group $G$, we think of each arrow as carrying some in-plane $G$-valued "flux" which must be conserved at each vertex in order to satisfy the vertex condition of the code space.

In the toric code, the plaquette operator flips the quantum numbers around a plaquette while staying within the space of loop configurations. Similarly in the Kitaev model, the plaquette operator changes the labels around the edges of the plaquette, but in such a way as to not disturb the vertex operators.

Analogous to the toric code, we may write the states in the ground state space (the code space) of the Kitaev model as

$$|\psi\rangle = \mathcal{N}^{-1/2} \sum_{\text{all edge labelings that can be obtained from a reference edge labeling via application of any } P_\beta(h)} |\text{labeling}\rangle$$

(29.9)

where $\mathcal{N}$ is the total number of terms in the sum. The analogy to flipping a plaquette in the toric code is the applications of the operators $P_\beta(h)$ for any group element $h \in G$ to any plaquette $\beta$. Note that from Eq. 29.8 that the eigenvalue of $P_\beta$ is not changed by the application of $P_\beta(h)$, nor is the eigenvalue of any $V_\alpha$.

As with the toric code the vertex and plaquette operators will provide just enough constraints so that the ground state on a spherical surface is unique and the ground state on a higher genus manifold will have a degeneracy that depends on the topology of the system, but does not depend on the number of lattice points we use in our lattice. I.e., the ground state space is described by a TQFT.

The TQFT that results is known as the quantum double or Drinfel’d double of the group $G$. Although the detailed study of the properties of the quantum double can get complicated (See references at the end of the chapter) we can introduce much of the key physics with fairly simple\(^8\) diagrammatic reasoning.

\(^8\) well, not too complicated!

29.3 Kitaev Ground State is Topological

Here we will show that the ground state of Kitaev model is topological, that is, that it is independent of the detailed geometry of the lattice, but depends only on the topology of the manifold. Our strategy is to show that we can locally reconfigure the structure of the lattice (keeping the topology fixed) and uniquely map a ground state wavefunction on one lattice structure to a ground state wavefunction with another lattice structure.\(^9\) This will then imply that, for example, the ground state degeneracy is independent of the lattice that we choose to use, depending only on the underlying topology of the manifold.

\(^9\)This mapping gives an isomorphism between ground state spaces.
To restructure our lattice, we need only two restructuring moves, which are shown in Figs. 29.5 and 29.6. Using these two moves any lattice on a surface may be mutated into any other lattice on the surface, so long as the overall topology of the surface remains unchanged (See exercise 29.3). It is worth noting that non-contractable loops on a surface cannot be removed by these moves.

We must determine how the ground state wavefunction is changed under these two moves. First, let us consider the vertex merging/splitting move shown in Fig. 29.5. In splitting the single vertex into two vertices the added edge (ab in the lower part of the figure) has its quantum number completely fixed by the vertex condition. Thus in adding this edge there is no additional degree of freedom added, thus we have a unique mapping of the wavefunctions between the two structures.

The first move to consider is the vertex splitting/merging move shown in Fig. 29.5. In general in such a move, two neighboring vertices with coordination $n$ and $n'$ may be moved to a single point, eliminating the intervening edge, and creating a vertex of coordination $n + n' - 2$.

In a ground state, it is fairly easy to see that such a vertex merging/splitting move does not alter any of the important physics. In particular an orthogonal basis of ground state wavefunctions before the merging move can be mapped uniquely to another basis for the ground state wavefunctions after the merging move. The key here is to realize that the quantum number (the group label) on the intervening edge that is removed in the merging process (labeled $ab$ in the left of Fig. 29.5) is not an independent degree of freedom in the ground state. Due to the vertex condition, the label on this intervening edge is completely determined by the labels on the other edges intersecting the two vertices. Thus there is a trivial one-to-one mapping between the wavefunctions before and after merging.

The second move we need to consider is the plaquette addition move shown in Fig. 29.6. Here a vertex is inserted into an edge and “tadpole” is added to the the new vertex. This tadpole has a stem (labeled $k$) and a loop (labeled $h$) in the figure. The loop splits the plaquette ($\beta$) into two plaquettes ($\beta_1$ and $\beta_2$ in the figure).

Again we would like to to show that the ground states of the two lattices are in one-to-one correspondence. The fusion rules require that the stem of the tadpole can only be labeled by the identity element, so (although the edge is still there) diagrammatically we can ignore the stem altogether. If $|\Psi\rangle$ is a ground state wavefunction of the entire system on the lattice before splitting the plaquette, we thus construct a
wavefunction after splitting as

$$|\Psi'\rangle = |\Psi\rangle \otimes \frac{1}{|G|} \sum_{h \in G} |h\rangle$$  \hspace{1cm} (29.10)$$

The sum over $h$ now assures that $\hat{P}_{\beta_2} = 1$, i.e., it puts $\beta_2$ in the unique ground state. Since the stem is labeled only with the identity element, the operator $\hat{P}_{\beta_1}$ in the right figure then acts on the outer edges of the larger plaquette exactly the same as $\hat{P}_\beta$ did in the left figure. Thus we have an isomorphism between the ground states on the two lattices.

Once we have established this independence of geometry, we can find the ground state degeneracy for various topologies by using a minimal lattice decomposition. Let us consider the case of a sphere. (We will discuss the torus case in section 29.5 below). The minimal decomposition of a sphere is shown in Fig. 29.7. This consists of a single vertex on the equator, a single edge running around the equator and two plaquettes — one covering the north hemisphere and one covering the south hemisphere. The edge running around the equator is labeled with $g$ and the vertex condition is automatically satisfied. The two plaquette operators are both equal to

$$\hat{P}_{\beta N} = \hat{P}_{\beta S} = \frac{1}{|G|} \sum_{g \in G} \hat{P}_{\beta N}(g)$$

Thus the unique +1 eigenstate of this operator is given by

$$|\Psi\rangle = \frac{1}{|G|} \sum_{g \in G} |g\rangle$$

and is the unique ground state of the system.

### 29.4 Continuum model

As with the toric code, we are not tied to any particular lattice. We can further dispense with the lattice altogether and consider diagrams in the absence of the lattice as shown, for example in Fig. 29.8.
In this language we can write the wavefunction analogously as
\[ |\psi\rangle \sim \sum_{\text{all diagrams that can be obtained from a reference diagram via allowed moves}} |\text{diagram}\rangle \quad (29.11) \]

(Compare this to Eq. (28.1) for the toric code.)

The “allowed moves” between such diagrams are of the three types shown in Fig. 29.9.

- **Move 1:** “Isotopy”. Meaning any smooth deformation of lines.

- **Move 2:** Adding and Removing Contractable Loops. A contractable loop with any label can be added or removed.

- **Move 3:** Reconnecting (trivial \( F \)) move.

![Fig. 29.9 Allowed diagrammatic moves in the Kitaev model. Compare Fig. 28.3](image)

The first move is similar to what we had in the case of the toric code. The second move, that we can add or remove loops, is also similar to the case of the toric code, except that the loops have arrows as well as group valued labels. The third move is “new” but should look familiar from our discussion of \( F \)-moves. However, here the \( F \)-coefficient is simply unity — and there is no sum over states on the right hand side (compare to Fig. 16.3). This type of \( F \)-matrix (which is not a matrix but simply a scalar) is what we called the trivial cocycle in section 20.1.1.

In addition to these three moves one can reverse arrows by inverting the group element (as in Fig. 29.2), and one can freely add or remove lines labeled with the identity element of the group.

Using these moves, we can derive a number of useful (and perhaps unsurprising) lemmas such as those shown in Fig. 29.10 and Fig. 29.11. (See exercise 29.2)

It is sometimes useful to merge the language of quantum numbers on
the lattice with the language of continuum diagrams off the lattice. A very convenient example of this is the statement that the operator $\hat{P}_\beta(h)$ from Eq. 29.3 can be thought of as inserting a loop labeled $h$ inside of a plaquette and then merging it into the edge variables as shown in Fig. 29.12.

With this notation we can write the plaquette operator diagrammatically as

$$
\hat{P}_\beta \left[ \begin{array}{c}
\beta \\
\times \Omega
\end{array} \right] = \left[ \begin{array}{c}
\times \Omega
\end{array} \right] \quad (29.12)
$$

where here the purple line is an $\tilde{\Omega}$ loop (compare Fig. 17.10) defined in this case as

$$
\tilde{\Omega} = \frac{1}{|G|} \sum_{h \in G} h \quad (29.13)
$$

The $\times$ in the middle of the loop in Eq. 29.12 indicates that the $\tilde{\Omega}$ loop is not contractable to a point so that Move 2 is not allowed (we can think of the $\times$ as being a puncture in the plane), but instead the loop must be pushed into the edges as in Fig. 29.12 to achieve exactly the operation defined in Eq. 29.5. This principle, that the plaquette operator is exactly a $\tilde{\Omega}$ loop is true of all of our generalizations of the toric code.
29.4.1 Brief Comments on Twisted Kitaev Theory

Once we have made the connection between the Kitaev model and a planar diagrammatic algebra it becomes natural to ask whether we can generalize the diagrammatic algebra and build more general topological models. Keeping with the idea of basing the model on the properties of a group $G$, we can construct other consistent planar diagram algebra by generalizing the $F$-move (move 3 in Fig. 29.9). Indeed as discussed in section 20.1.1 any so-called 3-cocycle provides a consistent scalar $F$-matrix for a planar diagrammatic algebra. The 3-cocycle used by the Kitaev model is just the trivial cocycle (i.e., $F$, or $\omega$ in the notation of section 20.1.1 is just unity). However, one can build a version of the Kitaev model that uses nontrivial cocycles (nontrivial $F$’s) as well. This is known as twisting the Kitaev model (or sometimes known as a quasi-quantum double). The word “twisting” here has nothing to do with the twists introduced in chapter 15. The language comes from its original usage in the context of discrete gauge theories. As we will discuss in section 29.7, the Kitaev model is essentially a discrete gauge theory. The twisted Kitaev model corresponds to what is known as twisted gauge theories. The twisted Kitaev models are also equivalent to the Dijkgraaf-Witten models introduced in section 21.4 with the un-twisted Kitaev model corresponding to the case of the trivial cocycle. The difference between (possibly twisted) Kitaev and Dijkgraaf-Witten is that the Kitaev model is a two dimensional model with a Hamiltonian and continuous time, whereas Dijkgraaf-Witten calculates an action for a discretization of a three-dimensional space-time.

29.5 Ground State Degeneracy on a Torus

We now turn to calculate the ground state degeneracy on a torus. We can use the minimal lattice decompositions of the torus as shown in Fig. 29.13 with a single vertex two edges and a single plaquette. We denote by $|b, a\rangle$ the state on the torus with the edge going around the vertical cycle labeled by $b$ and the edge going around the horizontal cycle labeled by $a$. 

\[ |b, a\rangle = \begin{array}{c} \includegraphics[width=0.3\textwidth]{torus_decomposition} \end{array} = \begin{array}{c} \includegraphics[width=0.3\textwidth]{trivalent_decomposition} \end{array} \sim \begin{array}{c} \includegraphics[width=0.3\textwidth]{more_trivalent_decomposition} \end{array} \]

Fig. 29.13 Decomposition of a torus into (left and middle) one vertex, two edges, and one plaquette. (Right) A slightly more complicated decomposition with two vertices, three edges, and one plaquette. This adds no additional degrees of freedom since $c$ is completely fixed by $a$ and $b$. However, this trivalent picture fits better with our discussion of diagrammatic algebras where vertices are always trivalent.

The vertex condition at the single vertex is $bab^{-1}a^{-1} = e$ with $e$ the
On the far right of Fig. 29.13 we show an alternative decomposition of the torus with two vertices and three edges and a single plaquette. This picture fits better with our continuum diagrammatic algebra where we insist on vertices that are trivalent\textsuperscript{10}. There is no added degree of freedom from the added edge, since the quantum number \(a\) is completely defined by \(a\) and \(b\). One vertex condition imposes \(c = ab\) and the other imposes \(c = ba\), so again \(a\) and \(b\) must commute.

\[
\hat{P}_\beta(h)|b, a\rangle = \begin{pmatrix}
|b, h, h, h, a, b, h, h\rangle = \begin{pmatrix}
h & b & h & h & a & h & h & h
\end{pmatrix}
\end{pmatrix}
\]

\[
hah^{-1} = |hbh^{-1}, hah^{-1}\rangle
\]

Fig. 29.14 Application of the plaquette operator \(\hat{P}_\beta(h)\) on the single plaquette conjugates both edge variables.

Let us now consider the plaquette operator \(\hat{P}_\beta(h)\) for the single plaquette. To apply this operator we push a loop labeled \(h\) into the edges (analogous to Fig. 29.12) thus conjugating both edge variable by \(h\) as shown in Fig. 29.14. Note that if \(a\) commutes with \(b\), then \(hhb^{-1}\) commutes with \(hah^{-1}\), so if the vertex condition is satisfied (if the two edges commute) the action of \(\hat{P}_\beta(h)\) does not change this.

We would now like to find the ground state or code space of our system which must be made of our basis states \(|b, a\rangle\) with \(b\) and \(a\) commuting but must be a superposition so that we obtain a +1 eigenvalue of the plaquette operator \(\hat{P}_\beta\) (Eq. 29.14). Given a state \(|b, a\rangle\) as above satisfying the vertex condition (i.e., with \(b\) and \(a\) commuting) we construct the state

\[
|\Psi_{b,a}\rangle \propto \hat{P}_\beta|b, a\rangle \propto \sum_{h \in G} |hbh^{-1}, hah^{-1}\rangle
\]

and here we do not keep track of a normalizing prefactor of the wavefunction. It is easy to check that this is indeed a +1 eigenstate of \(\hat{P}_\beta\) and is therefore a ground state. However, it is also true that not all sets of commuting \(b\) and \(a\) generate distinct wavefunctions. In particular, if we define the orbit of \(|b, a\rangle\) to be all of the states of the form \(|hbh^{-1}, hah^{-1}\rangle\) for any \(h \in G\), we realize that each distinct orbit represents a different orthogonal ground state wavefunction. I.e., Eq. 29.14 tells us to simply add up all of the wavefunctions within an orbit to make a ground state wavefunction.

**Example: \(S_3\)**

Let us consider the nonabelian group \(S_3\), the permutation group on three elements, also known as the symmetries of a triangle. (See section 41.2.1). This group has two generators \(X\) and \(R\) with the properties that \(X^2 = R^2 = e\) with \(e\) the identity element and \(XR = R^2X = R^{-1}X\). There are a total of six elements of the group (6 = 3! permutations)

\[
\Psi_{e,e} = |e, e\rangle
\]
\[
\Psi_{R,e} = |R, e\rangle + |R^2, e\rangle
\]
\[
\Psi_{e,R} = |e, R\rangle + |e, R^2\rangle
\]
\[
\Psi_{R,R} = |R, R\rangle + |R^2, R\rangle
\]
\[
\Psi_{e,X} = |e, X\rangle + |e, XR\rangle + |e, X^2\rangle
\]
\[
\Psi_{X,e} = |X, e\rangle + |XR, e\rangle + |XR^2, e\rangle
\]
\[
\Psi_{X,X} = |X, X\rangle + |XR, X\rangle + |XR^2, X^2\rangle
\]

Table 29.1 The eight wavefunctions generated by Eq. 29.14 Note that we have chosen a different normalization of these wavefunctions where we write each ket exactly once with a coefficient of unity.
which we list as $e, R, R^2, X, XR, XR^2$. We find eight orthogonal wavefunctions generated by Eq. 29.14 which are given in table 29.1. Note that these wavefunctions are not properly normalized.

General Case

For general nonabelian group it may seem complicated to try to figure out the dimension of the ground state space. In fact, there is a beautiful theorem from group theory, known as Burnside’s Lemma

\[ \text{number of orbits} = \frac{1}{|G|} \sum_{h \in G} \left( \text{number of elements of } X \text{ that are left unchanged by } h \right) \]

In our case, our set is the set of all elements $|b, a\rangle$ where $b$ and $a$ are elements of $G$ and commute, and the action of the group is conjugation as in Fig. 29.14. I.e., $h$ acts on $|b, a\rangle$ to give $|hbb^{-1}, hah^{-1}\rangle$. An element $|a, b\rangle$ is unchanged by $h$ if $h$ commutes with both $a$ and $b$. Thus we obtain the following general result

\[ \text{ground state degeneracy of Kitaev model on torus} = \text{number of orbits} = \frac{1}{|G|} \left( \text{Number of triples } a, b, h \text{ that all commute with each other} \right) \] (29.15)

29.6 Quasiparticles

Being that we know the ground state degeneracy on the torus, we know the number of quasiparticle types. In this section we will discuss the properties of these quasiparticles. Although this is a bit more complicated than what we did for the (abelian) case of the toric code, most of the development recapitulates the discussions of chapter 28 and particularly section 28.6.

Recall in the case of the toric code, and more generally for the $\mathbb{Z}_N$ toric code, the vertex defect involved the ends of strings. We can examine the system locally and determine the quantum number of this string end, even if we do not know where the string is. In the $\mathbb{Z}_N$ case we measure the charge of the vertex defect (see Eq. 25.18). However in the case of the Kitaev model with a nonabelian group, the “charge” at the end of a string is not as easily defined. To see this, consider Fig. 29.14. On the left we have a diagram with a string end labeled with $h$. Move 2 from Fig. 29.14 allows us to create the loop labeled $g$ on the left. This loop may then be pulled over the string-end which conjugates the label

\[ A \text{ group } G \text{ acting on a set } X \text{ here means a mapping } f(g, x) \text{ with } g \in G \text{ and } x \in X \text{ which gives an output in } X \text{ such that } f(e, x) = x \text{ and } f(g, f(h, x)) = f(gh, x). \]
of the string end to $ghg^{-1}$. Since the wavefunction of the Kitaev model should be in a superposition of diagrams from all allowed moves, the wavefunction must be in a superposition of having a loop end labeled $h$ and $ghg^{-1}$ for any possible group element $g$. Thus the string ends should not be labeled by their group element, but rather by a \textit{conjugacy class}.$^{13}$ When we do diagrammatic manipulations, we may choose a particular representative element of the conjugacy class (such as $h$ or some particular value of $ghg^{-1}$). However, we should remember that the group value of the string end is actually as superposition of all elements in the class.

With this knowledge we now would like to follow the tube algebra approach we introduced in section 28.2 and 28.6 in order to deduce the properties of the quasiparticles. We might start by introducing a basis of the form shown in Fig. 29.16 In the rectangular (right) representation of the annulus, $h$ comes in the bottom and $ghg^{-1}$ goes out the top (these are the inner and outer edges of the annulus). Both $h$ and $ghg^{-1}$ are in the same conjugacy class, but unlike the case of the toric code, they may not be the same group element.

$$|h, g\rangle = \begin{array}{c}
\includegraphics[width=0.5\textwidth]{fig29_16.png}
\end{array}$$

\textbf{Fig. 29.16} The basis we will work with for the tube algebra of the Kitaev model. The blue line going to the inner edge of the annulus (or off the top on the right) is labeled $ghg^{-1}$ to satisfy the vertex condition at the intersection.

We could work with the notation shown in Fig. 29.16 However, it turns out to be more convenient to work with a different notation. For each conjugacy class $C$ we choose one representative element which we call $h_C$ (it does not matter which element we choose). The remaining elements of conjugacy class can all be written as $p h_C p^{-1}$ for some$^{14}$ element $p$. We can then rewrite the basis state $|h, g\rangle$ in the manner shown in Fig. 29.17 In particular instead of writing $|h, g\rangle$ we write

$^{13}$To remind the reader (see section 11.2.2), the conjugacy class of $h$ is the set of all group elements that can be written as $ghg^{-1}$ for any $g$ in the group.

$^{14}$In fact for each element of the conjugacy class $h_i$ we should choose a particular $p_i$ such that $h_i = p_i h_C p_i^-$ and we should always use this value of $p_i$ to represent $h_i$. 
\[ |h,g\rangle = hghg^{-1} = hCz \]

\[ g = p'zp^{-1} \]

\[ \equiv |C,z;p',p\rangle \]

**Fig. 29.17** Graphically defining the \(|h,g\rangle\) basis in terms of the \(|h,g\rangle\). Here \(hC\) is the representative element of the conjugacy class \(C\) and \(zhC = hCz\) is required by the vertex condition.

\[ h = p^2 C p^{-1} \]

\[ ghg^{-1} = p' hC p'^{-1} \]

\[ g = p'zp^{-1} \]

\[ \equiv |C,z;p',p\rangle \]

**Fig. 29.18** Graphical demonstration of the composition law Eq. 29.17.

\[ |C, z; p', p\rangle \text{ where } C \text{ is a conjugacy class with representative element } h_C, \]

and we see from the figure that

\[ h = p^2 C p^{-1} ; \quad ghg^{-1} = p' hC p'^{-1} ; \quad g = p'zp^{-1} \]  

(29.16)

Note crucially that the vertex condition \(h_Czh^{-1}z^{-1} = e\) in the center of the figure requires that \(z\) commutes with \(h_C\), or we say that \(z\) is in the center of \(h_C\). I.e., \(z \in Z(h_C)\).

Our original basis \(|g,h\rangle\) had \(|G|^2\) different elements, and our new basis \(|C,z;p',p\rangle\) also has \(|G|^2\) different elements, so we are still spanning the same space\(^{15}\).

In terms of this new representation of the basis, the composition rule for such basis states is similar to the case for the \(Z_N\) toric code in Eq. 28.17 and is now given by

\[ |C_1, z_1; p'_1, p_1\rangle \circ |C_2, z_2; p'_2, p_2\rangle = \delta_{C_1C_2}\delta_{p_1p_2'} |C_1, z_1z_2; p'_1, p_2\rangle \]  

(29.17)

which we show graphically in Fig. 29.18.
We would now like to construct projectors from these basis states analogous to Eqs. \[28.18\] for the toric code. In that equation, we are essentially Fourier transforming to get from the basis to the projector. Here we need the non-abelian generalization of the Fourier transform, which is done with group representations. We write the following operator

\[
O^R_C(p', n'; p, n) = \frac{d_R}{|Z(h_C)|} \sum_{z \in Z(h_C)} \rho^R_{n', n}(z) |C, z, p', p\rangle
\]

where \(Z(h_C)\) is the centralizer of \(h_C\), the representative element of the conjugacy class \(C\). (Recall that the centralizer \(Z(h_C)\) is the group of elements that commutes with \(h_C\), and it has \(|Z(h_C)|\) elements). Here \(R\) is an irreducible representation of \(Z(h_C)\) of dimension \(d_R\) and \(\rho^R\) is the representation matrix (See section \[41.2.4\]).

Again, our original basis \([g, h]\) had \(|G|^2\) elements in it, and our new basis \(O^R_C(p', n'; p, n)\) also has \(|G|^2\) elements in it so we are still spanning the whole space\(^{16}\).

Using the grand orthogonality theorem Eq. \[41.6\] it is easy to prove (see exercise \[29.6\]) that these objects obey the projector law

\[
O^R_{C_1}(p'_1, n'_1; p_1, n_1) \circ O^R_{C_2}(p'_2, n'_2; p_2, n_2) = \delta_{C_1, C_2} \delta_{R_1, R_2} \delta_{p_1, p_2} \delta_{n_1, n_2} O^R_{C_1}(p'_1, n'_1; p_2, n_2) \tag{29.19}
\]

This is somewhat more complicated than the projector equation we had for the toric code Eq. \[28.19\]. Here \(p\) and \(n\) are simply additional boundary degrees of freedom that need to match when we sew together the annuli. The \(C\) and \(R\) degrees of freedom, however, are unchanged when we glue annuli together, and it is these indices that describe the quasiparticle types. Thus we have isolated the annulus states corresponding to the different quasiparticle types in the Kitaev model.

To emphasize: A quasiparticle type is described by a conjugacy class \(C\), and an irreducible representation \(R\) of the centralizer of a representative element \(h_C\) of the conjugacy class.

It is conventional to call the conjugacy class the magnetic charge and the representation \(R\) the electric charge (despite the fact that we are here thinking of the conjugacy class as being a vertex defect!). The origin of these names will be discussed in section \[29.7\] when we discuss the relationship to gauge theory.

Note that in the case of the \(\mathbb{Z}_N\) toric code, which is just the Kitaev model with the group \(G = \mathbb{Z}_N\), each group element is its own conjugacy class so there are \(|G|\) conjugacy classes, and the centralizer of any element is the entire group. The number of irreducible representations of the group is also \(|G|\) (number of irreps is always the number of conjugacy classes), so the total number of particle species is \(|G|^2\) in agreement with what we found in section \[28.6\].
As per footnote 14 we should sum over values of \( p \) such that we generate all possible \( h = ph_Cp^{-1} \) in the conjugacy class once. However, summing over the entire group is simpler and counts each \( h \) a fixed number of times, thus only changing the normalization.

\[
\langle \[C, R\] | \[C', R'\] \rangle = \delta_{CC'}\delta_{RR'}
\]

(29.21)

Again, in Eq. (29.20), we choose one representative value of \( h_C \) from each conjugacy class \( C \) and \( R \) is a representation of \( Z(h_C) \). If these make a basis for the states on a torus we must then have

\[
\text{ground state degeneracy of Kitaev model on torus} = \sum_C \text{(number of irreducible reps of } Z(h_C))
\]

(29.22)

where the sum is over conjugacy classes \( C, h_C \) is a representative element of the conjugacy class, and \( Z(h_C) \) is the centralizer of \( h_C \), i.e., the group of elements that commute with \( h_C \). It is not immediately obvious that this expression for the ground state degeneracy should match Eq. (29.15). Indeed, these two expressions do match (with a bit of group theory!). This is shown appendix 29.8.

**Example \( S_3 \) Again**

Lets return to the example of the group \( S_3 \) as we did in section 29.5. The group has three conjugacy classes \( C_e = \{e\} \), and \( C_R = \{R, R^2\} \), and \( C_X = \{X, XR, XR^2\} \). Let us choose the first element in each class as its representative elements. These have centralizers \( Z(e) = S_3 \) (the whole group, since everything commutes with \( e \)), \( Z(X) = Z_3 = \{e, X\} \) and \( Z(R) = Z_3 = \{e, R, R^2\} \). There are three irreducible representations of \( S_3 \) (See the table 20.1), which we call \( I, S, V \). For \( Z_2 \) and \( Z_3 \) there are 2 and 3 irreps respectively (see Eq. 41.5). The two irreps of \( Z_2 \) we call \( \{1, -1\} \) and the three irreps of \( Z_3 \) we call \( \{1, \omega, \omega^2\} \).
where $\omega = e^{2\pi i/3}$. Thus there are a total of eight quasiparticle types $[e, I]$, $[e, S]$, $[e, V]$, $[X, +1]$, $[X, -1]$, $[R, 1]$, $[R, \omega]$, $[R, \omega^2]$ matching the eight ground states we found in Table 29.1.

The characters of $S_3$ are given in Table 20.1 whereas the characters of the $Z_N$ are just the one-dimensional representation matrices themselves since the representations are scalar (See Eq. 41.5). Using Eq. 29.20 we obtain the eight explicit ground state wavefunctions shown in Table 29.2.

$$|e, I \rangle \sim \psi_{ee} + \psi_{Re} + \psi_{Xe}$$
$$|e, S \rangle \sim \psi_{ee} + \psi_{Re} - \psi_{Xe}$$
$$|e, S \rangle \sim 2\psi_{ee} - \psi_{Re}$$
$$|X, +1 \rangle \sim \psi_{XX} + \psi_{XX}$$
$$|X, -1 \rangle \sim \psi_{XX} - \psi_{XX}$$
$$|R, 1 \rangle \sim \psi_{RR} + \psi_{RR} + \omega^2 \psi_{RR}$$
$$|R, \omega \rangle \sim \psi_{RR} + \omega \psi_{RR} + \omega^2 \psi_{RR}$$
$$|R, \omega^2 \rangle \sim \psi_{RR} + \omega \psi_{RR} + \omega^2 \psi_{RR}$$

Table 29.2 The eight wavefunctions generated by Eq. 29.20. We have written these using the (unnormalized) basis states given in Table 29.1. Here $\omega = e^{2\pi i/3}$. The normalization of each wavefunction is simply a prefactor of $1/\sqrt{6}$ which we have not written for lack of margin space.

29.6.2 Quasiparticle Ribbon Operators on a Lattice

In the case of the toric code, we were able to create quasiparticles and pull them apart from each other by applying operations on a “string” of edges. For the vertex defects, we applied operations on a string of edges connecting two defects (Fig. 25.10) whereas for the plaquette defects we applied operations to a set of edges dual to a line connecting...
Fig. 29.19 A ribbon operator generally will operate on both the dark green edges and the red edges, and will create defects only at its endpoints.

The defects (Fig. 29.14). These strings of operators were defined by the fact that they were only detectable at their endpoints — the actual path the string takes could not be measured. Further if the strings formed contractable closed loops they act trivially on the ground state. One can construct similar objects, known as ribbon operators for quasiparticles of the Kitaev model as shown in Fig. 29.19. Most generally a ribbon operator will act on a string of edges on the lattice (red edges in Fig. 29.19) and also on a string of dual edges (dark green edges in Fig. 29.19).

Let us start by considering plaquette defects, which are fairly simple. As in the toric code case, the relevant operator will act on a string of edges (dark green in Fig. 29.19) dual to a path (bright green) between defects. As in the figure, we reorient all of the edges so that the arrows point toward the neighboring red path. Consider now measuring the ordered product of the group elements on all of these edges $x_1 x_2 x_3 \ldots$.

We write an operator

$$\hat{F}_p(\text{path}) = \delta_{g, x_1 x_2 x_3 \ldots x_N}$$

which measures whether this product equals $g$. Here we have assumed there are $N + 1$ plaquettes along the path (See Fig. 29.19 where $N = 5$). The subscript $p$ here stands for plaquette. This operator clearly commutes with the vertex operators, since both do not change the value of the group element along any edge. In addition this operator commutes with every plaquette operator except the plaquette at the start and the end of the path. To see this, consider the example shown in Fig. 29.20 where $P_\beta(a)$ is applied between to the plaquette between $x_1$ and $x_2$. This takes $x_1 \rightarrow x_1 a^{-1}$ but also takes $x_2 \rightarrow a x_2$ so that the product $x_1 x_2 x_3 \ldots$ remains unchanged. Thus this is a good candidate for a string operator that only makes plaquette defects at its ends.
To modify the values of the red edges, we insert a blue string labeled $h$ into the diagram and pull this string through the green edges as shown on the left. As it gets pulled through the green edge $x_1$ its value is conjugated to $x_1^{-1}hx_1$ and then when it gets pulled through $x_2$ its value is conjugated again to $x_2^{-1}hx_1x_2$. On the right, the blue line is then fused into the red lines. $y_1 \rightarrow y_1h$, $y_2 \rightarrow y_2h$, $y_3 \rightarrow y_2x_1^{-1}hx_1$, and so forth. Note that this operation commutes with all of the vertex and plaquette operators except at the endpoints of the ribbon.

The other operator we want to consider, the analog of the vertex defect, is a bit more complicated. As with the vertex defect string for the toric code, the general idea will be to put a string on top of a line (the red line) of edges. However the procedure for doing this is a bit more complicated. Consider adding a string with quantum number $h$ starting in the first plaquette and ending in the last plaquette as shown in Fig. 29.21. If we think about our diagrammatic algebra, when we pull this string through the dark green edges, its value gets conjugated by the value of the green edge. So for example, on the left of the figure, the blue string starts out labeled $h$ but after passing through $x_1$ it is labeled $x_1^{-1}hx_1$ analogous to Fig. 29.15, and then after also passing through $x_2$ it is labeled $x_2^{-1}x_1^{-1}hx_1x_2$ and so forth. Once we have this blue string in the diagram, we can then fuse it into the red edges to obtain the picture on the right of Fig. 29.21. Let us call this operator

$$F^h_v(path) = \text{pull } h \text{ string along path and fuse into edge}$$

with the subscript $v$ indicating the analog of a vertex defect (although actually this operator can create plaquette excitations at its ends as well as vertex excitations). We note that this operator commutes with all vertex and plaquette operators except at the end points of the ribbon — i.e., it does not create any defects except at the ends of the ribbon.

Let us first consider the vertex operators. As an example, examine the vertex with $q_1$ incoming in Fig. 29.21. On the left the vertex condition is that $q_1^{-1}y_1x_2^{-1}y_2^{-1} = e$. On the right, after fusing the blue $h$-line into the red edges the condition becomes $q_1^{-1}(y_3x_1^{-1}hx_1)x_1^{-1}(y_2h)^{-1} = e$ which is the same condition. It is thus easy to see that if the vertex condition is satisfied before fusing the blue line into the edge, it is also satisfied after. (Note that unlike the toric code, one generically does not have ribbon operators that violate the vertices without also violating the plaquettes as well.)
Now let us consider the plaquette operator $P_3(g)$ applied to the plaquette between $x_1$ and $x_2$ (same as in Fig. 29.20). The type of edge we need to focus on is an edge like $y_3$ in Fig. 29.21. If we first apply $F_h^v$ to the path as shown in Fig. 29.21 we take $y_3 \to y_3x_1^{-1}h x_1$. Then if we apply $P_3(a)$ this edge becomes $y_3x_1^{-1}h x_1 a^{-1}$. On the other hand if we apply $P_3(a)$ first, this edge becomes $y_3x_1^{-1}a^{-1}$. But at the same time the edge $x_1$ becomes $x_1 a^{-1}$. When we then pull the $h$ string through this $x_1 a^{-1}$ edge it becomes $(x_1 a^{-1})^{-1}h(x_1 a^{-1})$ and fusing the $h$ string into the $y_3 a^{-1}$ edge we then get $y_3 a^{-1}(x_1 a^{-1})^{-1}h(x_1 a^{-1}) = y_3 x_1 h x_1 a^{-1}$ which is the same result, thus showing that this operator only creates defects at the endpoints of the string.

We have constructed two types of operators $F_g^p$ and $F_h^v$ for quasiparticle ribbon operators. Analogous to the case of the toric code, these are only detectable at their endpoints, and string operators that form contractable closed loops act trivially on the ground state.

The most general type of operator we can construct is a combination of the two types of strings we have found

$$\hat{F}^{g,h}(\text{path}) = \hat{F}_v^h(\text{path})\hat{F}_p^g(\text{path})$$

It does not matter which order we put the operators on the right as they commute with each other. Thus we have $|G|^2$ types of ribbon operators. An appropriate linear superposition of these operators will create the quasiparticle types that we constructed in Eq. 29.18. Indeed it is quite simple to figure out what the correct superposition needs to be. To do this, use the strategy of section ?? and mutate the lattice step by step.

We can take any ribbon operator between two end points and convert it into a picture like Fig. 29.16 where $g$ is measured around a loop and $h$ is applied between the inner and outer edges of the annulus. (This requires stretching one of the $x_i$ edges in Fig. 29.19 all the way around in a loop around the endpoint of the ribbon). We then conclude that our quasiparticle ribbon operators can be written as

$$\hat{F}^{[R,C]}_{p,n,p',n'}(\text{path}) \sim \sum_{z \in \mathbb{Z}(h c)} \rho_{R,n}^{p,n'}(z) \hat{F}^{g,h}(\text{path})$$

where $g$ and $h$ are derived from $z, p, p'$ and $h_C$ via Eqs. 29.16

### 29.7 Relation to Gauge Theory

The Kitaev model is essentially a discrete gauge theory with gauge group $G$. To see this relation, let us begin with a quick review of the basics of gauge theory.

In chapters [4] and [5] we already worked with gauge theories. Recall that the fundamental quantity we needed to keep track of was the Wilson loop operator (See Eq. 5.11), which is the generalization of the Aharonov-Bohm phase\(^\text{19}\).
29.7 Relation to Gauge Theory

\[ W_{\text{loop}} = \text{Tr}_R \left[ P \left\{ \exp \left( i \oint_{\text{loop}} dl^\mu A_\mu \right) \right\} \right] \]

where \( P \) means we should path order the integral in the case where the gauge field \( A_\mu \) takes its value in a (generally nonabelian) Lie algebra. Here the trace is taken in some representation. I.e., the value inside the curly brackets \( \{ \} \) is an element \( g \) of the Lie group, and one then takes \( \text{Tr}[\rho^R(g)] \) where \( R \) is a representation and \( \rho \) is the representation matrix.

Let us now discretize space, introducing vertices which are connected together by edges as in the right of Fig. 29.22. (Note crucially, this is not going to be the same lattice as the one in the Kitaev model, but will rather be dual to that lattice.) For a directed edge between vertex \( i \) and \( j \) we can assign a group value

\[ g_{i,j} = P \left[ \exp \left( i \int_{\mathbf{r}_i}^{\mathbf{r}_j} dl^\mu A_\mu \right) \right] \]

Traversing the edge in the opposite direction gives the inverse group value \( g_{j,i} = g_{i,j}^{-1} \). A Wilson loop which visits sites \( i_1, i_2, i_3, \ldots, i_N, i_1 \) is then given by

\[ W_{\text{loop}} = \text{Tr}_R [ g_{i_1,i_2} g_{i_2,i_3} \ldots g_{i_{N-1},i_N} g_{i_N,i_1} ] . \quad (29.28) \]

For a discrete gauge theory, all of the values of \( g_{ij} \) in the system are assumed to take values in a discrete group \( G \). We can then choose to work with the representations of the discrete group.

Now in gauge theories we are allowed to make gauge transformations, which change the gauge field, but leave every Wilson loop unchanged (We can think of this as leaving the “magnetic flux” through every loop unchanged). The way we do this on a discrete lattice is to assign a group element \( u_i \in G \) to each site \( i \). The gauge transformation then takes

\[ g_{ij} \rightarrow u_i \ g_{ij} \ u_j^{-1} \]

Compare this to the gauge transform of Eq. 5.19 which we discussed for Chern-Simons theory, which has exactly the same structure. Once this is plugged into Eq. 29.28 and using the cyclic invariance of the trace, it is clear that the flux becomes gauge invariant.

We say that a gauge field configuration is flat if the flux through any loop is zero, or the group element in Eq. 29.28 before taking the trace, gives the identity.

We now write a Hamiltonian using the following two physical principle:

1. Gauge invariance is imposed energetically rather than as a strict symmetry principle.

2. Flat configurations should be energetically favored.

While principle (2) is typical of gauge theories (even in electromagnetism the Hamiltonian is minimized for zero flux), principle (1) is somewhat unusual. The ground state of our system will maintain gauge in-
Recall we can always flip the direction of an arrow and invert the corresponding group element. I.e., $g_{i,j} = g_{j,i}^{-1}$.

To implement principle (1) in a Hamiltonian, we define a gauge transform operator $A^h_i$ at site $i$ with $h \in G$ to premultiply all $g_{i,j}$ by $h$ and postmultiply all $g_{j,i}$ by $h^{-1}$. Thus we have for example

$$A^h_i = \begin{pmatrix} g_{i,j} & g_{i,k} \\ g_{j,i} & g_{k,i} \end{pmatrix}$$

where $h \in G$. To enforce gauge invariance we write a term in the Hamiltonian at each site $i$

$$A_i = \frac{1}{|G|} \sum_{h \in G} A^h_i$$

(29.29)

Now if the wavefunction is gauge invariant, it will be unchanged by the application of this operator. Further, it is easy to check that $A_i A_i = A_i$, so that $A_i$ is a projector, having eigenvalues of 0 and 1 only: Only the gauge-invariant part of a wavefunction survives when this operator is applied.

To implement principle 2, that we want flat, or flux-free, configurations to be energetically favorable, we define a projection operator that acts on plaquette such that it gives the identity if plaquette is flat (i.e., the group element in Eq. 29.28 is the identity) and otherwise gives zero. Such an operator acting on plaquette $p$ would then be, for example,

$$B_p = \begin{pmatrix} g_{i,j} & g_{j,k} & g_{k,l} & g_{l,i} \\ g_{k,i} & g_{l,j} & g_{i,k} & g_{j,l} \end{pmatrix}$$

Obviously $B_p$ is also a projector, that is $B_p B_p = B_p$, so it has eigenvalues of 0 and 1 only as well.

The full Hamiltonian, implementing both principles, is given by

$$H = -J_1 \sum_{\text{vertices } i} A_i - J_2 \sum_{\text{plaquettes } p} B_p$$

(29.30)

with positive $J_1$ and $J_2$. The ground state space is the space where all operators $A_i$ and $B_p$ have eigenvalue 1 on all vertices and plaquettes respectively.

The Hamiltonian Eq. 29.30 is precisely the one introduced by Kitaev [1997]. It is fairly easy to that it is also equivalent to the one we introduced in Eq. 29.8 but written on the dual lattice as shown in Fig 29.22. The operator $V_\alpha$ from our earlier construction (section 29.1.1) that acts
Fig. 29.22 Left: The lattice structure for the Hamiltonian Eq. 29.8 earlier in this chapter. We have labeled the plaquettes with black numbers. The directed edges \( g_{i,j} \) are labeled with \( i \) and \( j \) which are the plaquettes to the left and right of the edge respectively (facing the direction of the arrow). Right: The lattice we introduced for the Hamiltonian Eq. 29.30 is dual to the lattice on the left. Vertices are labeled and edges \( g_{i,j} \) are directed from vertex \( i \) to vertex \( j \). Middle: We have shown both the lattice in the left panel (black) and the lattice in the right panel (blue) overlaid on each other. For simplicity we have dropped the edge labels and the arrows — except for a single bond which is labeled to show that when two edges of the two lattices cross they are labeled with the same group element. A plaquette on the left becomes a vertex on the right and vice-versa.

on the vertices of the original model is identical to the operator \( B_p \) introduced here which acts on the plaquettes of this model, whereas the operator \( P_\beta \) that acts on the plaquettes of the original model is identical to the operator \( A_i \) which acts on the vertices here.

The plaquette defects in this model are plaquettes that violate \( B_p \) (i.e., have eigenvalue of \( B_p \) of 0 rather than 1). These defects are plaquettes that are not in “flat” configurations, and we can think of these plaquettes as having “magnetic flux” through them. This lends these quasiparticles the name “magnetic” as we mentioned in section 29.6. Correspondingly, violations of \( A_i \) are known as “electric”. Quasiparticles that violate both vertex and plaquette conditions are usually known as dyons.

29.8 Appendix: Two Expressions for Ground State Degeneracy on a Torus

Here we will show that Eq. 29.15 and Eq. 29.22 for the ground state degeneracy give the same result.

First, we use Burnside’s lemma (see section 29.5) to calculate the number of conjugacy classes of a group \( Z \). Let the set \( X \) be the group \( Z \) itself and the action of the group on this set is conjugation \( x \to hxh^{-1} \) with \( x \in X = Z \) and \( h \in Z \). The number of orbits of this action is the number of conjugacy classes which is just

\[
\left( \frac{\text{number of conjugacy classes of group } Z}{|Z|} \right) = \frac{1}{|Z|} \sum_{h \in Z} \left( \frac{\text{number of } x \in Z \text{ that commute with } h}{1} \right)
\]

(29.31)
We now consider the sum in Eq. 29.22. We can write this sum over classes as a sum over all group elements with each term in the sum weighted by the inverse of the size of the conjugacy class.

\[ \sum_C \rightarrow \sum_{g \in G} \frac{1}{|C_g|} \]

with \(|C_g|\) the number of elements in the conjugacy class of \(g\). Then using Eq. 29.31 along with the fact that the number of representations of a group is equal to the number of conjugacy classes in the group (See section 41.2.4), the sum in Eq. 29.22 may be written as

\[ \sum_{g \in G} \frac{1}{|C_g|} \frac{1}{|Z(g)|} \sum_{h \in Z(g)} \left( \text{ number of } x \in Z(g) \text{ that commute with } h \right) \] (29.32)

Finally we use a well known theorem (Eq. 41.3) that given an element \(g\) of a group \(G\), the number of elements \(|C_g|\) in the conjugacy class of \(g\) times the number of elements \(|Z(g)|\) that commute with \(g\) is equal to the number of elements in the group \(|G|\). I.e., \(|C_g| \cdot |Z(g)| = |G|\). Thus since \(Z(g)\) are the elements of \(G\) that commute with \(g\), Eq. 29.32 becomes

\[ \frac{1}{|G|} \left( \text{ number of triples } g, h, x \text{ which all commute with } g \right) \] (29.33)

in agreement with Eq. 29.15.

Further Reading

The original discussion of the Kitaev model is given by Kitaev [1997] and is still a fairly good discussion.

A nice physical description of the quantum double of a group is given by Preskill [2004]. Earlier work on the physics of the quantum double, and the twisted quantum double, is given by de Wild Propitius [1995].

A detailed analytic discussion of the properties of the quantum double of a group is given in Delcamp et al. [2017]; Delcamp [2017], including a detailed discussion of the tube algebra.

A detailed discussion of the Kitaev model and how one might implement it in experiment is given by Brennen et al. [2009].

A discussion of condensations from the the Kitaev model (in the sense of chapter 23) is given by Bombin and Martin-Delgado [2008] and also Beigi et al. [2011]. These are also good general references about the Kitaev model. Bais et al. [2003] describes condensations of the quantum double of a group in a more abstract approach.

For modular data (\(S\) and \(T\) matrices) of twisted quantum doubles, see Coste et al. [2000]. An explicit construction of a lattice model for twisted quantum doubles is given by Hu et al. [2013a].

Some details of using the Kitaev model as an error correcting code
are given by Cui et al. [2020].

The relationship between the Kitaev model and the Levin-Wen string net (chapter ***) is given in Buerschaper and Aguado [2009].

The theories discussed in this chapter (discrete gauge theories) can be generalized to 3+1 dimensions. This is discussed in some detail, including a discussion of the higher dimensional tube algebra, by Delcamp [2017]; Bullivant and Delcamp [2019].

Another generalization is to “higher” (or sometimes “higher-lattice”) discrete gauge theories in 2+1 and 3+1 dimensions. In such a generalization, one assigns not only gauge variables on edges, but also a “higher” gauge variable on the plaquettes. These are discussed by Bullivant et al. [2017, 2020] and less abstractly by Huxford and Simon [2021].

Exercises

Exercise 29.1 Product of Groups

It is stated in section 29.1.2 that the code space of a Kitaev model for a group $G = G_1 \times G_2$ is the tensor product of the code spaces for a Kitaev model built from group $G_1$ and the code space for a Kitaev model built from group $G_2$. Prove this statement.

Exercise 29.2 Diagram Manipulation in the Kitaev Model

(a) Using the moves given in Fig. 29.9, derive the result shown in Fig. 29.23 for the Kitaev model.

(b) Derive the result shown in Fig. 29.11.

(c) Show that the value of the tetrahedral figure in Fig. 29.24 is unity if all of the vertices are allowed, and is zero otherwise. In other words, show its value is $\delta_{dc^{-1},e}\delta_{bc,f^{-1}}\delta_{b^{-1},d_1,a}$.

Exercise 29.3 Mutating Lattices

(a) Show how to use the two moves given in Fig. 29.5 and 29.6 to achieve the following deformation of the lattice:

(b) Show that any lattice with a given topology may be mutated into any other lattice with the same topology using the two moves given in Fig. 29.5 and 29.6. Hint: Remove edges one at a time until you are left only with noncontractable loops.

Exercise 29.4 Kitaev Handle-Slide
Exercises

Use the diagrammatic rules to show the planar handle-slide identity shown in Fig. 29.25 where Ω is the Kitaev Kirby-strand defined in Eq. 29.13. Note that the × in the middle of the Ω loop should be thought of as a puncture in the plane that prevents this loop from being contractable.

Exercise 29.5 Ground State Degeneracy of $S_3$
Calculate the ground state degeneracy of the Kitaev model on a torus for the group $S_3$ by using Eq. 29.15.

Exercise 29.6 Kitaev Annular Projectors
Prove Eq. 29.19. Hint, use $\rho_R(ab) = \rho_R(a)\rho_R(b)$ along with the grand orthogonality theorem.

Exercise 29.7 Kitaev Model Twist Factor
Given the quasiparticle projection operator Eq. 29.18 and the twist operation on the $|h,g\rangle$ basis in Eq. 29.23, derive the twist factor in Eq. 29.25. Why is the expression for the twist factor of unit magnitude? Hint: (i) write the character $\chi_R$ as the trace of a matrix $\rho_R$ as in Eq. 41.7 then use $\rho_R(ab) = \rho_R(a)\rho_R(b)$. (ii) You will also need Schur’s second lemma (see section 41.2.4).

Exercise 29.8 Kitaev Model Quasiparticle Torus Basis
Prove Eq. 29.21. You will need the grand orthogonality theorem Eq. 41.6 as well as hint (ii) from exercise 29.7.

Exercise 29.9 Kitaev Model $S$-matrix
(a) [Easy] Using Eq. 29.28 give a simple expression for $S_{0,0} = 1/D$ for the Kitaev model for an arbitrary group $G$. Compare your result to that of exercise 21.5.
(a) [Harder] Derive Eq. 29.28.

Exercise 29.10 Kitaev Model for $S_3$
Calculate the $T$ matrix and the $S$ matrix for the group $S_3$. Hint: You can either use the explicit formulae (Eqs. 29.25 and 29.28) or you can use the wavefunctions explicitly in Eqs. 29.23 and 29.26.

Answers:

\[
\begin{array}{cccccccc}
\text{particle} & |e, I\rangle & |e, S\rangle & |e, V\rangle & |X, +1\rangle & |X, -1\rangle & |R, 1\rangle & |R, \omega\rangle & |R, \omega^{-1}\rangle \\
\theta & 1 & 1 & 1 & 1 & -1 & 1 & \omega^{-1} & \omega \\
\hline
1 & 1 & 2 & 3 & 3 & 2 & 2 & 2 & 2 \\
1 & 1 & 2 & -3 & -3 & 2 & 2 & 2 \\
2 & 2 & 4 & 0 & 0 & -2 & -2 & -2 \\
3 & -3 & 0 & 3 & -3 & 0 & 0 & 0 \\
3 & -3 & 0 & -3 & 3 & 0 & 0 & 0 \\
2 & 2 & -2 & 0 & 0 & 4 & -2 & -2 \\
2 & 2 & -2 & 0 & 0 & -2 & -2 & 4 \\
2 & 2 & -2 & 0 & 0 & -2 & 4 & -2 \\
\end{array}
\]

Exercise 29.11 Kitaev Model for the group $Q_8$
[Harder] Calculate the $T$ matrix and the $S$ matrix for the group $Q_8$. The group description and the character table is given in section ???. You can make a number of consistency checks on your answer using Eqs. 17.13, 17.20 and 17.24.
Doubled Semion Model

One of the simplest generalizations of the toric code is known as the **doubled semion model**. As with the toric code, it can be considered as a form of a quantum loop gas. However, additional signs are added to the loop gas to *twist* the theory in the sense discussed in 29.4.1 (indeed, the doubled semion model is the simplest case of a twisted Kitaev theory). The doubled semion model is also a simple example of the Levin-Wen string net model as we will discuss in chapter ***.

We managed to derive everything about the toric code by starting with the skein rules (Eqs. 28.2 and 28.3) of the $d = +1$ version of the $\mathbb{Z}_2$ fusion rule planar diagram algebra discussed in section 18.1.1. To obtain the doubled semion model, we instead use the $d = -1$ version of the $\mathbb{Z}_2$ fusion rule planar diagram algebra:

\[ \text{Here we are twisting (See section 29.4.1) the } \mathbb{Z}_2 \text{ fusion rules by using the nontrivial cocyle discussed in section 20.1.2).} \]

\[ (30.1) \]

\[ (30.2) \]

The diagrammatic rules are thus changed so that each loop removal or addition, and each surgery, incurs a minus sign. Note that these two minus signs are consistent with each other because each surgery changes the parity of the number of loops in the system. Note that these rules were precisely the skein rules we used for the Kauffman invariant when we considered semions (See section 18.1.2 for example). However, here we are not inputting any rules about braiding, just the planar diagram algebra.

From these rules we expect wavefunctions of the form

\[ |\psi\rangle = \sum_{\text{all loop configs that can be obtained from a reference loop config}} (-1)^{\text{number of loops}} |\text{loop config}\rangle \quad (30.3) \]

We can think of the prefactor $(-1)^{\text{number of loops}}$ as being the evaluation of the loop diagram exactly as in Eq. 18.14.

\[ \text{In the language of section 14.5 we are performing a non-unitary evaluation of this diagram, i.e., we are not using rule 0 which would give an additional minus sign for each cap. We will fix this issue in section 30.3 below. *** CHECK THIS} \]
As with the toric code, there should be four ground states on the torus corresponding to the different possible parities of blue strings around the two cycles.

### 30.1 A microscopic model

We would like to build a microscopic model that implements this $d = -1$ diagrammatic algebra analogous to the way the toric code Hamiltonian (Eq. 26.1) implements the $d = +1$ skein rules. We will once again work on a honeycomb lattice\(^3\) although any trivalent graph will work just as well. Again we put a spin on each edge, and color the edge blue if the spin is down. Following our procedure for the toric code, we will impose a vertex term (Eq. 25.1 in Eq. 26.1) that penalize any vertex with an odd number of blue edges, such that any ground state wavefunction must be made of closed loops. Again we want to have a plaquette term which flips the state of the plaquette (analogous to Eq. 25.3), but each flip must now come along with an appropriate number of minus signs. To determine the sign, it is useful to look at a few cases, as shown in Fig. 30.1.

\[\text{Loop addition: Sign} = -1\]

\[\text{Isotopy (deformation): Sign} = +1\]

\[\text{Surgery: Sign} = -1\]

\[\text{Double Surgery: Sign} = +1\]

**Fig. 30.1** When flipping over a plaquette, the sign is $-1$ if the number of loops in the system changes parity. A “leg” of the plaquette is one of the edges pointing out of the plaquette, i.e., having only one end on a vertex of the plaquette (these are drawn partially dotted in the figure). Let the number of blue legs of the plaquette be $L$, which is always even as long as we have no vertex defects. The sign associated with flipping the plaquette is $-1$ if $L/2$ is even and is $+1$ otherwise. This rule holds for plaquettes with any number of sides.

As shown in the figure, there is a simple rule for determining the sign accumulated for a plaquette flip. Given a plaquette which we want to flip, a leg of the plaquette is an edge with only one vertex on the
plaquette. The number of legs $L$ that are blue is always even as long as there are no vertex defects (loop ends) on the plaquette. The sign associated with flipping the plaquette is $-1$ when $L/2$ is even and is $+1$ otherwise. Another way to write this sign is as

$$\text{Sign} = -(1 - \sigma_z)^{\frac{1}{4}} \sum_{i \in \text{legs}} (1 - \sigma_z^{(i)}) = - \prod_{i \in \text{legs}} e^{\frac{\pi}{4}(1 - \sigma_z^{(i)})}$$

noting that $(1 - \sigma_z)$ gives 0 for spin up and 2 for spin down (i.e., a blue edge). With this information we can now write a plaquette operator for this model

$$P'_{\beta} = \left( \prod_{i \in \text{plaquette } \beta} \sigma_x^{(i)} \right) \prod_{i \in \text{legs of } \beta} e^{\frac{\pi}{4}(1 - \sigma_z^{(i)})}$$

where the first term is the same as that of the toric code (Eq. 25.3) and now we have included these additional signs.

The overall Hamiltonian for this model is then

$$H_{\text{doubled semion}} = -\frac{\Delta_L}{2} \sum_{\text{vertices } \alpha} V_{\alpha} - \frac{\Delta_P}{2} \sum_{\text{plaquettes } \beta} P'_{\beta} \quad (30.4)$$

which is identical to that of the toric code (Eq. 26.1) except for the modification of the plaquette term to include the appropriate signs.

### 30.1.1 Magnetic String Operator

In the case of the toric code we showed how we can pull vertex (e) defects apart from each other by successive application of a string of $\sigma_x$ (See section 25.4.1 and Fig. 25.10). Similarly we showed (See section [25.4.2](#)) that a (dual) string of $\sigma_z$ operators could be used pull plaquette operators apart from each other. Such strings are creatively known as “string operators”. The key property of such operators is that they only create excitations at the ends of the string, not along their length — and indeed, the actual path that the string operator follows is not measurable.

The analog of the magnetic string operator is quite straightforward. As with the toric code, one draws a path on the dual lattice (bright green in Fig. 30.2) and applies $\sigma_z$ to every edge that this dual string crosses (dark green in Fig. 30.2). Since, except at the end of the string, each plaquette has an even number of $\sigma_z$’s applied, this string operator commutes with the Hamiltonian, and hence does not create any excitations except at the string ends.

There are two more possible string operators corresponding to two other quasiparticle types (as mentioned above, there are four ground states on the torus, so we expect four quasiparticle types including the vacuum). These, however, are more complicated, and we defer their discussion to the more general case in chapter *** below.
30.2 Graphical Analysis

We can now follow the graphical methods of section 28.2-28.4 to determine the quasiparticle types and their properties. The graphical algebra is similar to that of the toric code, however, signs are incurred from loop addition/subtraction and surgery. We have four basis states on the annulus, given in Figs. 30.3 and 30.4 (which are equivalent to Figs. 28.7 and 28.9 for the toric code except for the added minus signs!).

\[ |0,0\rangle = \begin{array}{c}
\end{array} = \begin{array}{c}
\end{array} = \begin{array}{c}
\end{array} \]

\[ |0,1\rangle = \begin{array}{c}
\end{array} \]

Fig. 30.3 The two equivalence classes of states on the annulus where no lines intersect the boundary. In the top line, a minus sign is added with the loop addition going to the middle figure, then another minus sign is obtained from the surgery (marked by the arrow) going to the right picture.

\[ |1,0\rangle = \begin{array}{c}
\end{array} = \begin{array}{c}
\end{array} = \begin{array}{c}
\end{array} \]

\[ |1,1\rangle = \begin{array}{c}
\end{array} = \begin{array}{c}
\end{array} \]

Fig. 30.4 The two equivalence classes of states on the annulus where one blue line intersects each boundary. The two equivalence classes of states on the annulus where no lines intersect the boundary. In the top line, a minus sign is added with the loop addition going to the middle figure, two more minus signs are obtained from two surgeries going to the right picture at the position of the arrows. In the bottom line a minus sign is obtained from surgery at the position of the arrow.

We can now consider the multiplication or composition of states analogous to that discussed in section 28.2.2, that is, putting one annulus inside another. The multiplication table, Table 30.1, is entirely the same as that for the toric code (Table 28.1) except for one entry: \[ |1,1\rangle \circ |1,1\rangle = -|1,0\rangle. \] To understand this, we imagine putting one \[ |1,1\rangle \]
state inside another, and we obtain a picture like the far right of the top line of Fig. 30.4, which now differs from $|1,1\rangle$ by a minus sign.

With this multiplication table, we can construct the projectors to determine the four quasiparticle types:

$$O_I = \frac{1}{2}(|0,0\rangle - |0,1\rangle) = \frac{1}{2} (\begin{array}{c} \bullet \\ \bullet \end{array} - \begin{array}{c} \bullet \\ \circ \end{array})$$  
(30.5)

$$O_m = \frac{1}{2}(|0,0\rangle + |0,1\rangle) = \frac{1}{2} (\begin{array}{c} \bullet \\ \bullet \end{array} + \begin{array}{c} \bullet \\ \circ \end{array})$$  
(30.6)

$$O_{sR} = \frac{1}{2}(|1,0\rangle - i|1,1\rangle) = \frac{1}{2} (\begin{array}{c} \bullet \\ \circ \end{array} - i \begin{array}{c} \bullet \\ \bullet \end{array})$$  
(30.7)

$$O_{sL} = \frac{1}{2}(|1,0\rangle + i|1,1\rangle) = \frac{1}{2} (\begin{array}{c} \bullet \\ \circ \end{array} + i \begin{array}{c} \bullet \\ \bullet \end{array})$$  
(30.8)

Again we have predicted the names of these particle types. Since the ground state is a superposition of flipped and unflipped with a minus sign (since addition of a loop includes a sign), we identify Eq. 30.5 as the identity $I$. This superposition can again be thought of as a Kirby string $\Omega$ being put around the annulus$^4$.

Eq. 30.6 we will call “magnetic”, $m$, it is the projector orthogonal to the ground state within the space having no blue lines intersecting the boundaries of the annulus.

The two projectors with blue lines intersecting both boundaries (Eq. 30.7 and 30.8) are labeled $sR$ and $sL$ for “right handed semion” and “left handed semion” respectively.

We can proceed to calculate the twist factors for each of these particles. As in the case of the toric code (section 28.3), the twist operator $\hat{\theta}$ leaves $|0,0\rangle$ and $|0,1\rangle$ unchanged, implying that $\theta_I = \theta_m = 1$, i.e., these are bosons. Again as with the toric code $\hat{\theta}|1,0\rangle = |1,1\rangle$. However, in contrast to the bottom of Fig. ??, we have $\hat{\theta}|1,1\rangle = -|1,0\rangle$ as shown in Fig. 30.5.

$$\hat{\theta}|1,0\rangle = \hat{\theta} \begin{array}{c} \bullet \\ \bullet \end{array} = |1,1\rangle$$

$$\hat{\theta}|1,1\rangle = \hat{\theta} \begin{array}{c} \bullet \\ \circ \end{array} = - \begin{array}{c} \bullet \\ \bullet \end{array} = -|1,0\rangle$$

4There are two types of particles in the $\mathbb{Z}_2$ loop gas, the vacuum and the non-trivial particle (the blue string). We weight each by its value of $d/D^2$ with $D = \sqrt{2}$. Here we use $d$ rather than $|d| = d$ in the definition of the Kirby string $\Omega$ because we are performing a nonunitary evaluation of the loop diagram. See note 27 of chapter 17, as well as note 24 of this chapter.

Fig. 30.5 Twist deformations of the $|1,0\rangle$ and $|1,1\rangle$ states. The inner boundary is rotated counterclockwise while the outer boundary is held fixed, analogous to the deformation shown in Fig. 28.13. In the second line we have used the surgery equivalence as shown in Fig. 30.4.

| $\circ$ | $|0,0\rangle$ | $|0,1\rangle$ | $|1,0\rangle$ | $|1,1\rangle$ |
|-------|---------------|---------------|---------------|---------------|
| $|0,0\rangle$ | $\bullet$ | $\bullet$ |
| $|0,1\rangle$ | $\bullet$ | $\bullet$ |
| $|1,0\rangle$ | $\bullet$ | $\bullet$ |
| $|1,1\rangle$ | $\bullet$ | $\bullet$ |

Table 30.1 The multiplication table for composing annuli for the doubled semion model. The $\bullet$ indicates a composition which is not allowed due to non-matching boundary conditions.
Knowing now how $\hat{\theta}$ acts on $|1,0\rangle$ and $|1,1\rangle$ we can use this operator on $O_{sR}$ and $O_{sL}$ as given in Eq. 30.7 and 30.8 obtaining $\theta_{sR} = i$ and $\theta_{sL} = -i$.

We can further use the graphical techniques of section 28.5 to determine the fusion properties of the quasiparticles. Working out the details of this is assigned as Exercise 30.2, with the result being given in Table 30.2. We might hope that we can use similar graphical methods to derive the braiding properties and use the graphical (tube algebra) methods of section 28.4 to derive the $S$-matrix as well. However, here we run into a problem\(^5\). We have constructed our diagrammatic algebra using a $d = -1$ loop gas. As discussed section 14.5, this would correspond to a non-unitary diagrammatic algebra, since a loop has a negative value (See margin note 2 of this chapter). Having made this choice, the straightforward application of the graphical approach to calculating the $R$-matrix and $S$-matrix now gives some incorrect signs (Details are worked out in Exercise 30.1). However, as in section 14.5 there is a fairly easy way to repair this problem by counting caps. We work through this scheme in detail in section *** below. It is important to note, however, that the original Hamiltonian Eq. 30.4 is not faulty, and does indeed correspond to the ground state of a unitary TQFT.

Once we work out the details of the theory properly (in section ***), the resulting theory is simply the direct product of a left-handed semion theory and a right-handed semion theory. Recall from section 18.1.2 that a right handed semion theory has particle types $0_R$ and $1_R$, fusion rule $1_R \times 1_R = 0_R$ and $R_{0_R1_R}^{1_R1_R} = \theta_{1_R} = i$ and is equivalent to the Chern-Simons theory $SU(2)_1$. The left handed theory has particle types $0_L$ and $1_L$, fusion rule $1_L \times 1_L = 0_L$ and $R_{0_L1_L}^{1_L1_L} = \theta_{1_L} = -i$ and is equivalent to the Chern-Simons theory $\overline{SU(2)_1} = SU(2)_{-1}$. The doubled semion theory is thus

$$\text{Doubled Semion Theory} = SU(2)_1 \times \overline{SU(2)_1}$$

The product (See section 5.3) of the two theories means that each particle from the doubled semion theory is a combination of one particle from each of the constituent factors. Thus we have

$$I = (0_R, 0_L) \ ; \ sR = (1_R, 0_L) \ ; \ sL = (0_R, 1_L) \ ; \ m = (1_R, 1_L)$$

The fusion and braiding relations are inherited from the constituent theories as described in sections 5.3 ??, and ???. For example, it is easy to check that the simple $1 \times 1 = 0$ fusion rules of each factor gives the given the fusion Table 30.2 for the doubled model.

\(^5\)A problem which has caused quite a bit of confusion in the literature!
30.3 Gauge Choice and Unitary Diagram Algebra

The diagrammatic $d = -1$ version of the $Z_2$ loop gas that we chose to work with (Eqs. 30.1 and 30.2) corresponds to a nonunitary quantum theory. While this makes for a simple diagrammatic theory, it might seem that we might be better off working with a unitary diagrammatic algebra. Indeed, working with the unitary version of the algebra is a good way to avoid making mistakes! As discussed in section 16.1.3 (see also section 16.5) we can convert this diagrammatic algebra to a unitary theory with two steps:

1. Before evaluating a diagram, count the number of (blue) caps, and call it $n$. After fully evaluating the diagram (using the nonunitary rules with $d = -1$) multiply the final result by $(-1)^n$.

Here a “cap” is where a blue line experiences a maximum point (with respect to the chosen up direction). This is shown in Fig. 30.6 and 30.7.

We can thus modify our wavefunction Eq. 30.3 to the form of Eq. 30.9:

$$|\psi\rangle = \sum \text{all loop configs that can be obtained from a reference loop config} (-1)^{\text{number of loops} + \text{number of caps}} |\text{loop config}\rangle$$

(30.9)

Comparing this to Eq. 18.15 we see that the prefactor sign of the ket is just the unitary evaluation of the loop config diagram.

What does this change do to the ground state? Actually it is nothing more than a gauge change! It is a local rule that associates an extra minus sign to certain wavefunctions. This may seem odd that we are allowed to do this, being that in section 14.2 we concluded that we cannot generally remove minus signs from nonunitary theories with negative $d$ by gauge change. However, the context here is different. In section 14.2 we were concerned with theories in space-time (say, $1+1$ dimensions or $2+1$ dimension). In that case time is drawn vertically and it was crucial that a cup (right of Fig. 30.6) is the Hermitian conjugate of a cap (left of Fig. 30.6), this prevents us from making a gauge change of the cup and cap independently. Here, however, we are considering wavefunctions that live in a $2+0$ dimensional plane — no direction is meant to be time. We are thus free to associate a minus sign with the cap, but not the cup, and this is just a gauge change on the wavefunction!

We must now be careful to count the number of caps when we evaluate any diagram or we do any diagrammatic manipulation. Fortunately, for all the equalities drawn in Fig. 30.3 and 30.4 both sides of the equalities have the same parity of the number of caps, so we do not need to add any minus signs to the equations. This then implies that Eqs. 30.5-30.8 are still correct, and the deduced twist factors are still correct.

---

6 Note that if we represent the annulus as cylinder (or square with two sides identified) we might put the minus signs in different places compared to drawing an annulus in the more obvious way as in Figs. 30.4-30.5 for example. Any choice is allowed as long as we stick with it!

---

7 It may be convenient to add a factor of $-1$ to the $|0,1\rangle$ diagram and the $|1,1\rangle$ diagram since each one of these has a cap, but this is just a definitional change, and does not change the physical result.

---

Fig. 30.6 To unitarize the $d = -1$ version of the $Z_2$ loop gas, we assign a minus sign to each cap. A cap is a point where the blue line reaches a maximum such as on the left, but not the right.

Fig. 30.7 On the honeycomb, a cap configuration is as on the left. The right vertex can never be a cap no matter which edges are colored blue.
When we try to calculate the exchange phase, however, the minus signs will become important. As an example, let us try to counter-clockwise exchange two $sR$ particles as shown in Fig. 30.8. (This is analogous to the exchange of quasiparticles in Fig. 28.15 for the toric code.) Here we need to be very careful with keeping track of caps. In implementing the exchange operation, going from the first to second line of Fig. 30.8, we introduce exactly one cap in each figure, which is the reason for the overall minus sign we introduce out front. We then compare the second line to the first line to determine the overall braiding phase. Indeed, these two lines are equal up to a phase. To determine the phase, let us look, for example, at the left most picture in the second line of Fig. 30.8. Surgering where the small black arrow points picks up no net sign (this is shown carefully in Fig. 30.9), so the first figure on the bottom line is equivalent to the second figure on the top line, without the $-i$ prefactor and with the overall minus sign out front. This tells us that the exchange phase is $R_{sR,sR} = \theta_{sR} = i$ which agrees with Eq. 15.1 and 15.2, telling us that we have a right handed semion.

![Fig. 30.8](image)

**Fig. 30.8** Calculation that the phase accumulated from exchanging two $sR$ particles is $i$. The upper line represents two $sR$ quasiparticles, one in each hole (Compare Eq. 30.7). The holes are exchanged, braiding the blue lines in the process and creating a single cap which gives the overall minus sign out front. Using surgery and loop addition/removal (the moves of Eq. 30.1 and 30.2) as well as including a $-1$ for each cap, one can show the lower line is precisely $i$ times the top line showing an exchange phase of $i$. For example, the first picture in the second line is equivalent to the second picture in the first line (See Fig. 30.9) without the $-i$ prefactor and without the prefactor of $-1$.

We can similarly use graphical reasoning as in section 28.4 to calculate the $S$-matrix. We again define the operator that exchanges longitude and meridian on the torus. However, we must be careful to keep track of caps and signs. Since this is done more generally in section ..., we

![Fig. 30.9](image)

**Fig. 30.9** Close up of the first picture on the bottom row of Fig. 30.8. Reconnecting these blue strings requires first introduction of a cap, which gives a minus sign and then a surgery which gives another minus sign. The resulting figure is then identical to the second figure on the top line, without the $-i$ prefactor.
will leave this as an exercise for now.

### 30.4 Comments

It is interesting that we used the skein rules for a model of semions (Eqs. 30.1 and 30.2) to build our loop gas, and we got out both right and left handed semions. This is perhaps to be expected, since nowhere in our input rules did we ever break “time-reversal” or say whether the original theory was right or left handed — it comes out to be both!

This principle is very general. If we start with any theory of anyons and build a quantum loop gas from it (not putting in any of the braiding relations) we will get out the *doubled* theory, meaning it has both right and left handed versions of the theory. We will see this in more detail in chapter ***.

### Further Reading

Double Semion, first written by Levin-Wen; also Freedman et al.

### Exercises

**Exercise 30.1 Braiding Quasiparticles in the Doubled Semion Loop Gas**

(a) Use the technique of section 28.5 to show that counterclockwise exchanging two \( sL \) particles gives a phase of \( i \) whereas counter clockwise exchanging two \( sR \) particles gives a phase of \(-i\).

(b) Show that wrapping an \( sL \) all the way around an \( sR \) gives no phase.

(c) Show that wrapping an \( m \) particle around either an \( sL \) or an \( sR \) gives a phase of \(-1\).

(d) Show that wrapping an \( m \) particle around another \( m \) particle gives no phase.

**Exercise 30.2 Fusing Quasiparticles in the Doubled Semion Loop Gas**

Use the technique of section 28.6 to deduce the full fusion table for the toric code.

**Exercise 30.3 S-Matrix of Doubled Semion Model**

For this exercise it is perhaps easiest to rewrite all of the graphical diagrams for the annulus as squares with two opposite edges identified. Keep careful track of signs associated with caps.

Derive the S-matrix of the Doubled semion model by using the same technique as in section 28.4. Be careful of signs!
Levin-Wen String Net

The theme of the last few chapters has been to build up topological models from diagram algebras. One of the most general constructions of this type is the Levin-Wen\(^1\) string net model. The input information for building this model is a (unitary) planar diagram algebra as we discussed starting in chapter 12 and in more detail in chapter 14. More formally we say that we input a so-called spherical tensor category. The output of this model will be a quantum system described by a (modular) TQFT known as the Drinfel’d double or Drinfel’d center (or sometimes just the quantum double) of the input category.

The toric code (including the \(\mathbb{Z}_N\) toric code) and the doubled semion model are special simple cases of the Levin-Wen string net model. The Levin-Wen model can also be shown to be essentially equivalent to the Kitaev model as well, after a fairly simple transformation\(^2\). As such, this chapter brings together many of the ideas we have run into since chapter 25.

In the cases where the input diagram algebra can be given a braiding, such that resulting input theory plus the braiding is a modular anyon theory, the Drinfel’d double of this input theory is then simply two copies of the input modular anyon theory — one right handed copy, and one left handed copy. We saw this in the case of the doubled semion model in chapter 30. The planar algebra we put in could be given either a right-handed braiding to become a right-handed semion TQFT, or a left-handed braiding to become a left-handed semion TQFT. The Drinfel’d double of this planar algebra, the outcome of the string net construction, is then just the product of the left and right handed semion theories as we saw in section 30.2.

In the case of the toric code, however, while the input \(\mathbb{Z}_2\) planar diagram algebra can be given two possible braidings (either bosonic or fermionic, see section 15.1.1), neither braiding is modular. As such, the TQFT that arises from the Drinfel’d double, i.e., the toric code, does not factorize into a simple left-handed times right-handed theory. However, the two possible braidings, bosonic and fermionic, are two of the resulting particle types of the toric code.

More generally a planar diagram algebra (a spherical tensor category) may not have any braidings at all\(^3\) (i.e., no \(R\)-matrix satisfies the hexagon equation with the given \(F\)-matrices). Nonetheless, all such planar diagram algebras used as an input to a string-net model generate Drinfel’d doubles that are valid modular anyon theories.

Given a planar diagram algebra, such as those described in chapters 14 or 16 we would like our ground state wavefunction to be a sum over

\(^1\)This very general model is quite crucial to topological ideas. It was introduced by Levin and Wen [2005]. Michael Levin was a graduate student at the time. Xiao-Gang Wen is one of the founding figures in the field of topological condensed matter physics. He started his physics career as a student of Ed Witten.

\(^2\)The connection between Levin-Wen and Kitaev is explored in detail by Buerschaper and Aguado [2009]. In short, instead of describing edges by their group elements, one makes a transformation akin to a Fourier transform to describe the edge by its representations. The equivalence between these two descriptions of the same model is an example of a Morita Equivalence.

\(^3\)The simplest such examples are given by Hagge and Hong [2009]. These theories have three particle types \(I, x, y\) and fusion rules \(x \times x = 2x + y + I\) and \(x \times y = y \times x = x\) and \(y \times y = I\). There are several possible solutions of the pentagon equation, but none of these admit a solution of the hexagon equation.
all diagrams which are equivalent via the set of allowed diagrammatic manipulations. We thus intend ground state wavefunctions of the form
\[ |\psi\rangle \sim \sum_{\text{all diagrams that can be obtained from a reference diagram via allowed moves}} W(\text{diagram}) |\text{diagram}\rangle \]  
(31.1)

where \( W(\text{diagram}) \) is the weight of the diagram, which we determine simply by making an evaluation via the diagrammatic rules. So for example, if we have a theory with a particle type \( a \) having quantum dimension \( d_a \), if we add a contractable loop of type \( a \) to a diagram the weight \( W \) of the diagram is multiplied by \( 4 |d_a| \) as indicated in Eq. (31.1).

4Here we are assuming a unitary diagram algebra so the value of a loop is always positive (i.e. if \( d_a < 0 \) we apply rule 0 from section 14.3). The construction of the doubled semion model in chapter 30 shows us that, when we have \( d_a < 0 \), we could alternately choose to work in another gauge by not invoking rule 0 so that the value of a loop is negative and our planar diagrammatic algebra is not unitary. However, this is actually equivalent to working with a unitary diagram algebra. See in particular the discussion of section 30.3. Figs. 14.3 or 16.5.

31.1 Microscopic Hamiltonian

We would like to construct a microscopic Hamiltonian on a (regular or irregular) lattice which will implement the diagrammatic planar algebra – i.e., whose ground state space is described by wavefunctions of the form of Eq. 31.1. As usual, we here assume a trivalent lattice. For simplicity we might often assume a honeycomb, but any trivalent lattice, even a disorderd trivalent graph, works just as well. Edges are labeled with the possible quantum numbers of the planar diagram algebra. Unless a quantum number \( a \) is self dual (\( a = \bar{a} \)) then the edges must be labeled with arrows to indicate a direction. Reversing the direction of an arrow changes \( a \) to \( \bar{a} \) as shown in Fig. 31.1. If there are fusion multiplicities \( N_{ab}^c > 1 \) then vertices between three edges with labels \( a, b, c \) must also carry labels \( \mu \in 1, \ldots, N_{ab}^c \). For simplicity we will usually assume all \( N_{ab}^c \leq 1 \) so there are no vertex indices, although it is fairly easy to extend to the more general case.

31.1.1 Fat Lattice Construction

As we have discussed starting with the toric code, it is often useful to think on the continuum instead of thinking about degrees of freedom on the lattice. We were quite cavalier about doing this, and even mixed the lattice and continuum descriptions sometimes (such as in Fig. 29.12). Here we will make this idea a bit more formal with a construction due to Levin and Wen [2005], known as the “fat lattice”. Our microscopic model will be defined on a trivalent graph, and we add an \( x \) in the middle of each plaquette which we interpret as some sort of “puncture” in the manifold. We now allow ourselves to draw diagrams with particle lines that may or may not be on the edges of the graph. To interpret these lines for the physical system on the lattice, we deform the diagram, using our diagrammatic rules, until all of the lines lie on the lattice. (We sometimes say we “push” the diagram onto the lattice). In this process

\[ a \quad \Rightarrow \quad \bar{a} \]

Fig. 31.1 Reversing an arrow on an edge takes the particle type to its antiparticle.

5The construction is known as “fat lattice” because we think of the entire region between two marked punctures as being the result of taking the edge between the punctures and fattening it until it fills all the space between the punctures.
we use our diagrammatic equivalence rules, but we are not allowed to cross any lines over one of the marked ×’s. This procedure is well defined in the sense that there is a unique lattice configuration corresponding to each continuum diagram we draw. An example of this procedure is shown in Fig. 31.2.

### 31.1.2 Vertex and Plaquette Operators

We introduce a vertex operator at vertex \( \alpha \) as

\[
\hat{V}_\alpha = \delta(a, b, c) \left( \begin{array}{c} a \\ b \\ c \end{array} \right)
\]

(31.2)

where

\[
\delta(a, b, c) = \begin{cases} 1 & \text{if } (a, b, c) \text{ is an allowed vertex } (N_{abc} = N_{\bar{a}b} > 0) \\ 0 & \text{otherwise} \end{cases}
\]

Here \( \hat{V}_\alpha \) is a projector (it has eigenvalues of 0 and 1 only).

Analogous to the plaquette term in Eq. 29.12 for the Kitaev model, we graphically represent a plaquette operator as the introduction of an \( \Omega \) loop (See Fig. 17.10).

\[
\hat{P}_\beta = \left( \begin{array}{c} \beta \\ \times \\omega \end{array} \right)
\]

(31.3)

In the spirit of the fat lattice construction, the × in the middle of the
The loop should be thought of as a puncture so that the loop may not be contracted. To find the action of this operator of the edge variables, one should “push” the $\Omega$ strand into the edge, by using the diagram rules to fuse it into the edge variables. While one can write an explicit form of this operator in terms of the effect on the quantum numbers on the edges, it is not particularly enlightening to do so at this point, so we defer this exercise to Appendix 31.4.

Recall now the handle-slide property of the $\Omega$ strand shown in Fig. 31.3, which we originally derived in Fig. 22.10. This move allows us to quickly show that

$$\Omega \times a = \Omega \times a$$

where in the first step of Eq. 31.5 we use the handleslide (Fig. 31.3) and in the second step we use the fact that a contractable loop of $\Omega$ evaluates to unity (see Fig. 17.10). Eq. 31.3 indicates the $\hat{P}_\beta$ is a projector, having eigenvalues of 0 and 1 only.

Further because the planar diagram algebra is already known to be self-consistent (i.e., it does not matter in which order we do moves to evaluate a diagram) and that diagrammatic moves starting with an allowed fusion diagram never generate unallowed diagrams, we can quickly conclude that all of the $\hat{V}_\alpha$ and $\hat{P}_\beta$ operators commute with each other.

### 31.1.3 Levin-Wen Hamiltonian

Our Hamiltonian is then written in a now familiar form

$$H_{\text{Levin-Wen model}} = -\Delta_v \sum_{\text{vertices } \alpha} \hat{V}_\alpha - \Delta_p \sum_{\text{plaquettes } \beta} \hat{P}_\beta$$

with $\Delta_v$ and $\Delta_p$ both positive. The vertex term simply enforces the condition that all vertices have edges corresponding to allowed fusions of the planar diagrammatic algebra. The plaquette term allows the edge variables to fluctuate dynamically. The key here is that if we are in a ground state of the system then $\hat{P}_\beta = 1$ for all plaquettes, so that when we draw a diagram in the fat-lattice picture, we can surround each $\times$ with an $\Omega$ loop. However, due to the handle-slide identity (Fig. 31.3) our diagrammatic lines may now be freely deformed over the $\times$ punctures. As a result we may now use our diagrammatic rules entirely freely on the manifold ignoring the punctures entirely. This then enforces that the ground state wavefunctions must be of the form shown in Eq. 31.1.
31.1 Microscopic Hamiltonian

The reasoning here is subtle. We needed to puncture the plane so that we had a mapping from continuum diagrams on the fat-lattice to quantum numbers on the lattice. But then the plaquette operators tell us that we can freely deform over these punctures in a ground state wavefunction and may then apply all diagrammatic operations to our wavefunctions.

31.1.4 Ground State is Topological

As with the other diagrammatic models we have encountered, the toric code, the Kitaev model, and the doubled semion model, the ground state of this Hamiltonian is topological — meaning that it does not depend on the detailed geometry of the lattice, but only on the topology of the underlying manifold. The proof of this statement follows very closely the discussion of section 29.3 for the Kitaev model. We will briefly outline the argument here; a more detailed discussion is given by Hu et al. [2012]. The strategy here is to show that there is a simple one to one mapping between ground state wavefunctions on one lattice structure and those on a second lattice structure which has been altered (or “mutated”) locally without changing the overall topology of the surface.

The elementary moves we need to consider are shown in Figs. 31.4 and 31.5. Fig. 31.4 is simply an $F$-move (compare Fig. 16.3). Note that here we are not just changing the quantum numbers on the edge, but we are also restructuring the underlying lattice. Nonetheless it provides a unitary transformation between states on the left and right of the figure.

Fig. 31.5 is the splitting of a plaquette $\beta$ into two plaquettes $\beta_1$ and $\beta_2$. Unsurprisingly the stem of the tadpole (labeled $k$) can only be labeled
Levin-Wen String Net

The doubter might object that we have used a vertex with only two edges meeting it whereas we defined the model to have only trivalent vertices. If we want to be not cheat, we could use a decomposition of the sphere with two (trivalent) vertices, three edges, and three plaquettes. By using $F$ moves if necessary, this can be restructured into two connected tadpoles living on the surface of the sphere.

The connecting line in the middle must be labeled with the identity, leaving us just a bunch of $\Omega$ loops for the plaquettes, which have unique ground states.

Ground State on a Sphere

We can then establish, analogous to the case of the Kitaev model, that the ground state on a sphere is nondegenerate. To do this we choose the simplest lattice decomposition of the sphere (See Fig. 29.7 with a single vertex, a single edge forming the equator (connected to the single vertex at each end) and two plaquettes — one covering the north hemisphere and one covering the south hemisphere. The vertex condition is always satisfied (having the same quantum number going out as coming in) and the two plaquette operators are identical to each other. Thus the ground state is the unique state given by the +1 eigenstate of the plaquette operator which is just $\sim \sum s |s\rangle$. I.e., the wavefunction is a $\Omega$ loop around the single edge.\(^6\)

31.2 Braiding of the Input Category and String Operators of the Double

The general construction of quasiparticles for the Levin-Wen model can be quite complicated. However, in cases where the input category has a solution to the hexagon equation (i.e., has a braiding), certain quasiparticle string operators can be deduced trivially. Further in cases where the input category is modular all of the quasiparticle string operators can be deduced.

Recall that the definition of string operator is an operator that can be detected only at its ends — i.e., when operating on the ground state, it creates no defects except at its ends.

Let us suppose there exists a solution to the hexagon equation for the input category. This means we can give a diagrammatic meaning to over- and under-crossings. We construct a string operator graphically by imagining that we add a string from the input category either over or under the plane of the system. Then we operate on the edges of the physical system by fusing this string into the edge variables of lattice using the diagrammatic rules. An explicit example of turning this diagrammatic prescription into an operation on the lattice is given in Appendix 31.4.2. Here it is crucial to realize that the operator we are considering acts entirely on the two dimensional system — the diagram of a string above or below the lattice is simply a way to encode what operations we perform on the lattice variables.

Since the diagrammatic rules (including the braiding rules) are self-consistent, we are allowed to make diagrammatic transformations with-
out changing the value of the diagram. In particular we can freely deform the figure, by sliding the string around the lattice. This means it cannot cause any defects to either the vertex or plaquette operators along its length (This is the very definition of a string-operator – that it does not create defects except at its ends). A string operator that forms a closed contractable loop can be shrunk down to a very small loop and removed with the usual diagrammatic rule (Fig. 16.5), thus the closed loop acts trivially on the ground state (except possibly giving a constant depending on how we normalize the operator).

For a string operator which is a strand with ends, we must have a violation of the diagrammatic rules at the ends of the string. Recall that in diagrams if a particular quantum number comes into a region, that quantum number must also leave the region. (See for example, the locality constraint in Fig. 8.7). A string that comes to an abrupt end must violate this condition. Thus we expect that the system will locally not be in the ground state at the position of a string end.

We will now consider two examples of such string operators that we have already seen (albeit in different language).

### 31.2.1 Toric Code

In the toric code (section 28.1) the planar diagram algebra is the $Z_2$ loop gas with $d = +1$. There are two possible solutions of the hexagon, worked out in section 18.1.1, the boson and the fermion. It is not a coincidence that the quasiparticle types of the toric code include a boson and a fermion!

Let us consider the bosonic solution first. The boson braids trivially ($R^{[1]}_{01} = 1$ in section 18.1.1) so we only need to think about fusing this string operator into the lattice In Fig. 31.7 we show such a string operator (blue) above the toric code lattice (chosen to be a square lattice for simplicity). Fusing this blue line into the lattice at the position of the red path, simply flips the spins along this red path. This is exactly the string operator for the (bosonic!) vertex defects that we introduced in section 25.4.1.

We can also consider the fermionic solution of the hexagon. The fermion has braiding $R^{[1]}_{01} = -1$ meaning that we accumulate a minus sign every time the string (blue in Fig. 31.7) crosses over an edge of the lattice which itself is colored blue (meaning the spin on that edge is pointing down in the notation of chapter 25). Thus before fusing the blue string into the red path, we measure the state of the edges marked green in Fig. 31.7 and get a minus sign for each green edges where the spin is in the down position. This is precisely the action of the string operator that creates the plaquette defects that we introduced in section 25.4.2. One we have introduced the appropriate minus signs we must also fuse the string into the red path, which then flips the spins along this line. Thus the net effect of string operator is the product of the plaquette defect string and the vertex defect string, which we recall is the fermionic quasiparticle!

---

*Fig. 31.7* The blue string above the lattice is a string operator that we will fuse into the lattice at the position of the red line. If we choose the blue string to have bosonic braiding statistics then it simply flips the spins along the red line, creating vertex defects at the ends of the red line. If we choose the blue string to have fermionic braiding statistics, then it measures the spins along the green lines before flipping the spins along the red line. The measurement of the green spins creates plaquette defects, whereas the flipping of spins creates vertex defects. The combination of these two is a fermionic quasiparticle.

---

7 In this case it does not matter whether we think of the blue line as being above or below the lattice plane.
Thus the bosonic string operator corresponds to the vertex defect
boson, and the fermionic string operator corresponds to the fermionic
quasiparticle. The fusion of these two quasiparticles together would give
the plaquette defects (only measuring the spins on the green horizontal
edges). We have thus reconstructed all of the quasiparticle types of the
toric code with this graphical construction.

31.2.2 Doubled Semion

For the doubled semion model (chapter 30) the planar diagram algebras
is the $\mathbb{Z}_2$ loop gas with $d = +1$. There are two possible solutions of
the hexagon, worked out in section 18.1.2 corresponding to right and
left handed semions. Correspondingly, the quasiparticle types of the
doubled semion model include both a right and left hand semion.

Let us consider the case of $R_0^{11} = i$, the right-handed semion model.
We imagine placing a string of this type (drawn blue) either over or
under the lattice, as shown in Fig. 31.8. We would like to fuse this blue
line into the red marked path.

31.2.3 More Generally

31.3 Detailed Example: Doubled Fibonacci

It is worth working through one nontrivial example of a Levin-Wen
model in some detail to see how the Drinfel’d double arises. The simplest
examples, the toric code and the doubled semion model, were based on
the two possible $\mathbb{Z}_2$ planar fusion algebras as discussed in section 18.1
(in another language, these are the two possible cocycles, see section
20.1.2). However, these two are both abelian fusion rules so they are
perhaps a bit too simple to see what happens more generally.

Here, we will consider the Fibonacci planar diagram algebra (which
we ran into as far back as section 8.2.1) as an input to our Levin-Wen
model. This model is perhaps the simplest example with nonabelian
fusion rules, so it presents a good example for demonstrating how this
construction works. Note, however, that the Fibonacci planar diagram
algebra can be given a braiding, and the resulting theory is then modular.
As discussed above we then claim that the content of the Drinfel’d double
will be simply the product of a left and right handed copy of the modular
Fibonacci anyon theory.

$$\text{Doubled Fib} = (\text{Fib Anyons})_L \times (\text{Fib Anyons})_R$$

We will study modular theories more generally in section *** below.
However, it is still useful to work through this example explicitly to get
a feel for how these models work.

Recall that in the Fibonacci planar diagram algebra there is one non-
trivial particle type, which we usually call $\tau$, and the nontrivial fusion
Detailed Example: Doubled Fibonacci

The (unique unitary) diagrammatic algebra for these fusion rules are derived in section [31.2]. We summarize the important rules here:

\[ \tau \times \tau = I + \tau \]

Diagrams for such a model can be represented as a branching loop gas as shown in Fig. 31.9. In such diagrams lines, drawn without arrows, are allowed to branch at trivalent vertices, but cannot have endpoints.

The (unique unitary) diagrammatic algebra for these fusion rules are derived in section [31.2]. We summarize the important rules here:

\[ \phi = 1 + \frac{\sqrt{5}}{2} \]

We recognize the first two lines (Eqs. 31.7 and 31.8) as being simply the Fibonacci \( F \)-move (compare Fig. 18.8). Eq. 31.9 gives a loop the value of the quantum dimension, and Eq. 31.10 is the usual locality principle (See Fig. 16.7). From these equations along with the assumption of full isotopy invariance (any diagram may be smoothly deformed in any way in the plane without changing its value) these diagrams fully define the planar diagrammatic algebra for the Fibonacci model.

From these principles it is easy to derive a few very useful additional lemmas including (See exercise 31.4)

\[ \phi = \frac{1 + \sqrt{5}}{2} \]
31.3.1 Excitations

As with the double-semion model we should be able to determine the quasiparticle eigenstates by looking at how a single line can end in a defect. We claim that all possible line endings can be reduced, by $F$-moves, to one of the three possible endings shown in Fig. 31.10. Just as an example, consider the ending shown on the left of Fig. 31.11. By using an $F$-move, it is reduced to a combination of the three presented above.

![Fig. 31.10 Possible string endings in the doubled Fibonacci string net model.](image)

![Fig. 31.11 An example of reducing a more complicated string ending into one of the three endings shown in Fig. 31.10](image)

As in the case of the toric code and the double semion model, we can figure out the twist factors by rotating these diagrams as shown in Fig. 31.12 and then using $F$-matrices to reduce the result back to linear combinations of the same three possible endings.
31.3 Detailed Example: Doubled Fibonacci

Fig. 31.12 The rotation operator \( \hat{\Theta} \) applied to the possible string endings. Then using \( F \) matrices we reduce the results to linear combinations of the same endings.

We can write these diagrammatic equations more algebraically by

\[
\hat{\Theta}\begin{pmatrix} a \\ b \\ c \end{pmatrix} = \begin{pmatrix} 0 & \phi^{-1} & \phi^{-1/2} \\ 1 & 0 & 0 \\ 0 & \phi^{-1/2} & -\phi^{-1} \end{pmatrix}\begin{pmatrix} a \\ b \\ c \end{pmatrix}
\]

The eigenvectors of this matrix are the particle types with definite twist factors given by their eigenvalues under rotation.

With a bit of algebra it can be shown that the eigenvalues of this matrix are given by

\[
\theta = e^{i\pi/5}, \quad e^{-i\pi/5}, \quad 1,
\]

The first two correspond to the expected spin factors for a right-handed Fibonacci anyon \( \tau \) or left-handed Fibonacci anyon \( \tau^* \) (recall that we worked out the spin factor using the hexagon equation earlier. See [13.3]). The final possibility represents the fusion of these two objects \( \tau \times \tau^* \). Indeed, these are all of the possible particle types in the doubled-Fibonacci theory. Since the theory was based on a full anyon theory with braiding fully defined, we expected to get both a right- and left-handed copy of the Fibonacci model and indeed we did. (We never broke time reversal in the definition of the model so we should get both hands of the theory!).

31.3.2 Ground State Degeneracy

It is a bit tricky to figure out the ground state degeneracy here. Using the above skein rules, any configuration can be reduced to a linear combination of four simple configuration – corresponding to the possibilities...
Fig. 31.14 Collapsing a triangular bubble. This is derived in exercise ??.

of having a loop, or not having a loop, around each handle. An example of reducing two loops around a handle to a linear combination of zero and one loop is given in Fig. 31.13

Fig. 31.13 Reducing two loops around a handle to a linear combination of one loops and zero loops.

31.4 Appendix: Explicit Form of Operators

For simplicity here we will assume an isotopy invariant diagram algebra as in chapter [16] although the generalization to the more general case is not difficult (See for example Hahn and Wolf [2020]).

In this derivation we will use the identities shown in Fig. 31.15 and Fig. 31.14.

31.4.1 Plaquette Operator

Let us now define an operator $\hat{P}_\beta(s)$ (shown in Fig. 31.16) that merges in a loop labeled $s$ (drawn red in this figure for clarity) into a plaquette $\beta$.

In going from the first line to the second, we use Fig. 31.15 one time for each edge of the polygon (four times in this case). In going from the second line to the third, we use the triangle collapsing rule Fig. 31.14 one time for each corner and we cancel factors of $\sqrt{d}$.

Since the $\Omega$ strand (Fig. 17.10) is just a weighted sum of all of the
particle types the full plaquette operator defined in Eq. 31.3 is given by

\[
\hat{P}_\beta = \sum_s \frac{d_s}{D^2} \hat{P}_\beta (s) \tag{31.13}
\]

### 31.4.2 String Operator Example

Here we give an example of fusing a string operator into the lattice. The string operator is

We start with the figure on the left, with a string operator under the lattice (drawn red for clarity).

This pushing of the string into the edge is quite similar to what we did for the plaquette operator in section 31.4.1. The triangular bubbles can be collapsed using Fig. 31.14. The upper triangular bubble with edges \(i,j,s\) gives a factor of \(F_{ijs} \sqrt{d_id_j/d_i'}\). The lower triangular bubble including with edges \(g,h,s\) gives a factor of \(F_{ghs} \sqrt{d gd h /d g'}\).

Let us now focus on the part of the diagram with the undercrossing as shown in Fig. 31.19. Using the uncrossing rule shown in Fig. 31.18 we can then evaluate the remaining diagram on the right of Fig. 31.19 using the same triangle collapse law Fig. 31.14. The upper triangle including \(i,b,s\) gives a factor of \(F_{sib} \sqrt{d_s d_b /d s'}\) and then we collapse a lower triangle with edges \(h,z,s\) to give \(F_{zhb} \sqrt{d_h d_z /d h'}\).
Fig. 31.17 The string operator is drawn (red) as a string running under the lattice (blue). To determine the action of this operator, we use the diagrammatic rules to fuse the red string into the blue lattice. The first step is shown here.

\[
\sum_{g',h',i',j'} \sqrt{j'j} = \sum_{g',h'} \sum_{i',j'} \sqrt{i'j}
\]

Combining together all of these factor The final result can be written as Fig. 31.20 where the prefactor is given by

\[
K_{a,b,c}^{g,h,i,j,g',h',i',j',z} = \sum_{g',h',i',j'} \sum_{g,h,i,j} K_{a,b,c}^{g,h,i,j,g',h',i',j',z}
\]

This formula itself is not really that interesting. However, by working through how the red string is pushed into the edges, hopefully we have made the method clear!

### 31.5 Appendix: S-matrix for Fibonacci Anyons

Without doing much work, we can figure out the S-matrix for Fibonacci anyons. There are only 2 particles in the theory I and \( \tau \). Further we know that the quantum dimension of \( \tau \) is \( \phi = (1 + \sqrt{5})/2 \). Thus, the total quantum dimension is \( D^2 = 1 + \phi^2 = 2 + \phi \) and the S matrix must be of the form

\[
S = \frac{1}{D} \begin{pmatrix} 1 & \phi \\ \phi & y \end{pmatrix}
\]
where the constraint of unitarity immediately fixes $y = -1$.

We can check this by using $F$ and $R$ matrices to determine the value
of two linked rings explicitly as shown in Fig. 31.21.

\[
\begin{align*}
\phi^{-1} (R_{\tau \tau})^2 + \phi^{-1/2} (R_{\tau z})^2 &= \\
&= \phi^{-1} (R_{\tau \tau})^2 + \phi^{-1/2} (R_{\tau z})^2 \\
&= \left[ \phi^{-1} (R_{\tau \tau})^2 + \phi^{-1/2} (R_{\tau z})^2 \right] \\
&= \left( R_{\tau \tau}^2 + \phi (R_{\tau z})^2 \right) = -1
\end{align*}
\]

**Fig. 31.21** Calculating the nontrivial element of the Fibonacci anyon $S$-matrix.

---

**Exercises**

**Exercise 31.1 Fibonacci Diagram Algebra**

Use Eqs. 31.7 - 31.10 to derive the identities Eq. 31.11 and Eq. 31.12.

**Exercise 31.2 Double Fibonacci String Net**

(a) As discussed in lecture, the double Fibonacci model ground state can be viewed as a branching string net with graphical rules given by Fig. 31.22.

(Compare to the problem on Fibonacci pentagon relation) where $\phi^{-1} = (\sqrt{5} - 1)/2$. In the ground state no endpoints of strings are allowed, but branching is allowed.

To complete the graphical rules we must also use the rules shown in Fig. 31.23 for some values of the variables, $d$, $X$ and $T$.

(We did much of this in lecture. What was left out is proving that any $T \neq 0$ solution is not self-consistent. Hint: Try evaluating a circle with three legs coming out of it. That should enable you to derive a useful identity. Then see if you can use this identity to derive a contradiction when $T \neq 0$.

(b) Consider quasiparticles which are the ends of strings. The general form of a quasiparticle is as shown in Fig. 31.24 with coefficients $a, b, c$ that need to be determined. Find the eigenvalues/eigenvectors of the rotation operator to determine the quasiparticle types and their spins. (We did most of this in lecture except the explicit evaluation of the eigenvalue problem!) Compare your result to the result of the problem “Fibonacci Hexagon Equation".
Fig. 31.22 String net rules for the doubled Fibonacci model

Fig. 31.23 Additional string net rules for the doubled Fibonacci model

Fig. 31.24 Combination of defect types for the doubled Fibonacci model
Part VII

Symmetries and Entanglement
Anyon Permuting Symmetry

Some TQFTs have a symmetry under permutation of anyons. This situation is not generic, but when it does happen, the corresponding TQFT has some special properties that are worth discussing.

Conveniently, the toric code provides an excellent example of anyon permuting symmetry. In fact this symmetry is mentioned back in chapter 26. In the fusion table given in Fig. 26.1 it is noted that you can exchange all $m$'s for $e$'s and the fusion table would remain correct. Further, looking at Eq. 26.5 which gives the $S$- and $T$-matrices for the toric code, the second column and second row refer to the $e$ particle whereas the third column and third row refer to the $m$ particle. One can switch the second and third columns and rows of both matrices and the matrices remain unchanged. This shows that the toric code has a precise symmetry under permutation of $e$ with $m$.

More generally we say we have an anyon permuting symmetry whenever there is a permutation matrix $P$ such that

$$\begin{align*}
PSP^T &= S \\
PTP^T &= T
\end{align*} \tag{32.1}$$

where $S$ and $T$ are the modular $S$- and $T$-matrices (see chapter 17) and the superscript $^T$ means transpose. It may be the case that for a given TQFT there are multiple matrices $P$ that satisfy Eq. 32.1. Generally the set of such all $P$'s (including the identity matrix, which trivially satisfies Eq. 32.1) form a group known as the anyon permutation group (See exercise 32.2).

32.0.1 The Idea of a Symmetry Defect

When an anyon permuting symmetry exists, this opens the possibility that a theory can contain so-called “symmetry defects”. A symmetry defect is a point in a 2 dimensional system (thinking of 2+1 D systems, as we usually do) where the identity of a particle is changed (by one of the above described permutation matrices) if the particle is dragged around that point. For example, for the toric, an $m$ particle could be changed into an $e$ particle and vice versa, as shown in Fig. 32.1. We say that the $e$ and $m$ particles have been “permuted”. Note that this means while the two different types of excitations can be distinguished locally, globally they cannot be distinguished as one can be turned into the other by going around the defect.

Symmetry defects are not contained in the structure of TQFTs, but rather in a mathematical structure known as a $G$-crossed extension of

---

$^1$ A matrix $P$ is a permutation matrix if $P^TP = PP^T = 1$ and each row and each column of $P$ has only a single nonzero entry which is unity and all other entries are zero.
a TQFT (here $G$ represents the anyon permuting symmetry group). Rather interestingly these defects themselves can be treated as having certain anyon-like properties. While this more detailed mathematical structure is beyond our current discussion, interested readers can refer to the work of [Barkeshli et al. 2019] for details (see also references at the end of the chapter).

Microscopic models with anyon permuting symmetries and symmetry defects have been constructed in several ways. We will introduce here a particularly simple way to see the anyon permuting symmetry in the toric code.

### 32.1 Symmetric Form of the Toric Code

Let us try to make the toric code anyon permuting symmetry a bit more explicit, and a bit more physical. Recall from chapters 25 and 26 that the terms of the Hamiltonian for the toric code phase of matter (the “stabilizers” if we think of the toric code as actually being a code rather than a phase of matter) are the vertex operator (drawn as a green cross in the left of Fig. 32.2), and the plaquette operator (drawn as a pink square in the left of Fig. 32.2). To remind the reader, the vertex operator (originally defined in Eq. 25.1) is a product of four $\sigma_z$ operators, and the plaquette operator (originally defined in Eq. 25.3), is a product of four $\sigma_x$ operators around a plaquette.

$$V_\alpha = \prod_{i \in \text{vertex } \alpha} \sigma_z^{(i)}$$

$$P_\beta = \prod_{i \in \text{plaquette } \beta} \sigma_x^{(i)}$$

Violations of the vertex terms (if $V_\alpha$ is not in its $+1$ eigenstate) were the particles that we previously called “electric” or $e$ and violations of the plaquette terms (if $P_\beta$ is not in its $+1$ eigenstate) we previously called “magnetic” or $m$.

In the left of Fig. 32.2 these two types of terms look fairly different from each other. However, with nothing more than a notational change, we can draw the same system as on the right of Fig. 32.2. Here we simply draw the reference lines diagonally (not changing any of the physical spins, or any of the terms in the Hamiltonian) so that the system now looks like it is a tiling of diamonds. In this case both vertex and plaquette operators now act on a diamond shaped plaquette, and this starts to look a bit more symmetric between the electric and magnetic particles. However still the $V_\alpha$ operators, which act on the green diamonds is a product of four $\sigma_z$’s whereas the $P_\beta$ operators, which act on the pink diamonds is a product of four $\sigma_x$’s.

We can make a further slight transformation on the model to make it look even more symmetric between the two types of terms in the Hamiltonian. Let us make a unitary transformation on all the spins that are on the east and west corners of the green diamonds. The unitary transformation we choose is to pre- and post-multiply each spin by $\sigma_y$. 
Such a transformation converts

\[
\begin{align*}
\sigma_x & \rightarrow \sigma_z \\
\sigma_z & \rightarrow \sigma_x \\
\sigma_y & \rightarrow \sigma_y
\end{align*}
\]

As a result, what we previously called the vertex operator (a product of four \(\sigma_z\) operators around a green diamond) is now a product of two \(\sigma_z\) operators on the north and south of the green diamond times a product of two \(\sigma_x\) operators on the east and west of the green diamond. Similarly what we previously called the plaquette operator (a product of four \(\sigma_x\) operators around a pink diamond) is now a product of two \(\sigma_x\) operators on the east and west of the pink diamond times a product of two \(\sigma_z\) operators on the north and south of the pink diamond. So after this unitary transformation all of the operators on all of the diamonds have exactly the same form, as shown in Fig. 32.3. In this form, the toric code Hamiltonian can be written as

\[H_{\text{toric code}} = -\frac{\Delta}{2} \sum_{\text{diamonds } q} \sigma_z^{q_N} \sigma_x^{q_E} \sigma_z^{q_S} \sigma_x^{q_W} \]  \hspace{1cm} (32.2)

where here \(q_N\) indicates the spin in the north corner of diamond \(q\) (and analogously for \(q_S\), \(q_E\), and \(q_W\)). This Hamiltonian is physically identical to that of Eq. 26.1 where we have set \(\Delta_n = \Delta_y = \Delta\) (we have done nothing more than make a unitary transformation on some of the spins). It is easy to check that all the terms in the Hamiltonian commute with each other as in the usual toric code.

As mentioned above, in the original formulation of the toric code we had two different types of terms, the vertices, whose violations we called \(e\) and the plaquettes, whose violations we called \(m\). Now we have transformed both vertex and plaquette terms of the original toric code Hamiltonian to all look like identical diamonds. Nonetheless, some of

\[\text{Fig. 32.2} \hspace{1cm} \text{Left: In our prior formulation of the toric code, the vertex operators acted on crosses (green), and the plaquette operators acted on squares (pink). By simply drawing the reference lines diagonally, both types of operators act on a diamond plaquette — the vertex operators acting on the green diamonds and the plaquette operators acting on the pink diamonds.}\]

\[\text{Fig. 32.3} \hspace{1cm} \text{The terms in the Hamiltonian, Eq. 32.2, on every diamond (diamonds of either color in the right of Fig. 32.2) are a product of } \sigma_x \text{ on the north and south vertices, and } \sigma_z \text{ on the east and west vertices.}\]
Fig. 32.4 Lattice dislocations are marked with the red dots. Walking around a dislocation, one comes back to the opposite sublattice. As shown in the figure, two steps north, two steps east, two steps south, and two steps west does not return to the same point, but rather returns to a neighboring diamond on the opposite sublattice.

Starting in the ground state, if you apply a $\sigma_x$ operator on a vertex this creates a pair of excitations on the neighboring diamonds to the north and south of that vertex, whereas applying $\sigma_z$ to a vertex creates excitations on the neighboring diamonds to the east and west. $\sigma_y = i\sigma_x\sigma_z$ does both.

The discovery that dislocations create symmetry defects, and the investigation of the implications was made by Bombin [2010].

If one generates this diamond model by starting with a conventional toric code and converting it to diamonds as in Fig. 32.2 one always obtains a model where walking around a cycle along edges requires an even number of steps. However, one can generalize the model by removing a single row of spins, such that one obtains an odd number of steps around a cycle (and a model that cannot be obtained from a conventional toric code as in the mapping of Fig. 32.2). In such a model, when one goes around the cycle with an odd number of steps, one switches sublattices, so there is then no global distinction between the pink sublattice and the green sublattice. This is similar to what happens in section 32.2. Correspondingly, the ground state degeneracy becomes two rather than four.

32.2 Symmetry Defects in the Toric Code

Presenting the toric code in this more symmetric form allows one to naturally consider symmetry defects. In this picture, the symmetry defect is realized by a lattice dislocation\(^5\). In Fig. 32.4 we show two disloca-

\(^3\)Starting in the ground state, if you apply a $\sigma_x$ operator on a vertex this creates a pair of excitations on the neighboring diamonds to the north and south of that vertex, whereas applying $\sigma_z$ to a vertex creates excitations on the neighboring diamonds to the east and west. $\sigma_y = i\sigma_x\sigma_z$ does both.

\(^5\)The discovery that dislocations create symmetry defects, and the investigation of the implications was made by Bombin [2010].

\(^4\)If one generates this diamond model by starting with a conventional toric code and converting it to diamonds as in Fig. 32.2 one always obtains a model where walking around a cycle along edges requires an even number of steps. However, one can generalize the model by removing a single row of spins, such that one obtains an odd number of steps around a cycle (and a model that cannot be obtained from a conventional toric code as in the mapping of Fig. 32.2). In such a model, when one goes around the cycle with an odd number of steps, one switches sublattices, so there is then no global distinction between the pink sublattice and the green sublattice. This is similar to what happens in section 32.2. Correspondingly, the ground state degeneracy becomes two rather than four.
tions (marked with red dots). We imagine moving an excitation around dislocation. As mentioned above, the excitation must jump between diamonds that share exactly a single vertex. Thus we can follow the path of the green arrows marked in the figure: north two steps, east two steps, south two steps, and west two steps. At the end of this procedure we have returned to the opposite sublattice (with sublattices defined locally). Hence if we had started with an $e$ particle we now have an $m$ particle and vice-versa. Thus we have implemented anyon permutation — the dislocation is a symmetry defect!

The careful reader will notice from Fig. 32.4 that at the position of the defect the plaquette has five vertices rather than four. In the Hamiltonian our rule was that the operator for each plaquette has $\sigma_z$ for the north and south vertices and $\sigma_x$ for the east and west vertices. For the 5-vertex plaquette we maintain this rule, but we also add the rule that for the additional vertex (the one that is 3-valent rather than 4-valent) we will include a factor of $\sigma_y$. This is shown in Fig. 32.5. The operator we will use for this plaquette will then be

$$\sigma_y^N \sigma_x^E \sigma_z^S \sigma_y^{3\text{-valent}}$$

It is easy to check that this operator has eigenvalues $\pm 1$ and commutes with all of the other plaquette operators from Eq. 32.2. Thus we still have a commuting projector model which is easily solved analytically.

We may then consider a system, say, on a torus, with some number of regular plaquettes and some number of dislocations. As we did in Section 25.3, we can count up the number of degrees of freedom to determine the ground state degeneracy. First, we notice that the product of all plaquette terms over the entire lattice is always $+1$. This is analogous to the constraints in Eq. 25.8 and Eq. 25.9. For a lattice with no dislocations, the two sublattices (green and pink) in Fig. 32.2 generate two different constraints. However, with dislocations there are not two distinct sublattices (since going around the dislocation takes you from one local sublattice to the other), and thus there is only a single constraint.

Let the number of spins be $N$, and the total number of plaquettes be $N_p$. It is not too hard an exercise (see Exercise *** ) to show that the number $n$ of dislocations must be

$$n/2 = N - N_p$$

showing that the number of dislocations must be even.

Let us now count degrees of freedom. We start with $N$ qubits. There are $N_p - 1$ independent plaquette operators (the single constraint mentioned above accounts for the $-1$). Thus putting all of the plaquettes in the ground state leaves $n/2 + 1$ qubits still undetermined. If there are no dislocations, this means the system can store a single qubit of information — or there is a ground state degeneracy of 2. For each pair of dislocations, there is one more qubit unassigned, or the ground state...
degeneracy doubles. This means that each dislocation is associated with half of a qubit.

How can it happen that a dislocation is associated with half of a qubit? Each dislocation traps a Majorana zero mode! We have run into Majorana zero modes before in our study of Ising anyons (See Exercise 5.3.9.7 and the comments in section 8.2.2). In short these are localized operators, two of which constitute a qubit. Thus the dislocations themselves have anyon-like properties!

Further Reading

- The original paper by Bombin [2010] gives a much more detailed study of the properties of the dislocations in the toric code model.
- You and Wen [2012] Originated the degrees-of-freedom argument at the end of this chapter. This paper also extended the work to \( Z_p \) toric codes.
- more her

Exercises

Exercise 32.1 Permutation Symmetry of the Toric Code
[Easy] What is the permutation matrix for Eq. 32.1 for the symmetry of the toric code.

Exercise 32.2 Permutation Symmetry Group
[Easy] If two different permutation matrices \( P_1 \) and \( P_2 \) both satisfy Eq. 32.1 show that the product \( P_1 P_2 \) also satisfies Eq. 32.1. Hence conclude that the set of possible permutation matrices form a group.

Exercise 32.3 Counting Degrees of Freedom
[Easy] Consider a lattice on a torus like that of Fig. 32.4 with some number of dislocations inserted into the lattice. Each normal plaquette has four sides, and each dislocation plaquette has five sides. Prove Eq. 32.3. Hint: use the Euler characteristic for a torus, faces+vertices=edges.

Exercise 32.4 \( Z_N \) Toric Code Symmetry Defects
[Hard] The entire construction performed in sections ?? and 32.2 can be extended to the \( Z_N \) toric code introduced in section 25.6 and 26.5. Show that each pair of dislocations harbors an \( N \) state system. We say that each dislocation is associated with a \( Z_N \) parafermion. (Note: These are not the same type of parafermion as discussed in chapter ***. This is an unfortunate over-use of the same name).
Topological Entanglement

We might wonder what is special about the ground states of the models we have been studying. What is it that makes them topological? Another way of asking the same question is: how is it that anyone can arise from simple degrees of freedom such as a system of simple spin-1/2 (like we have for the toric code). One interesting answer to these questions is that TQFT ground states are special because they have a special type of long ranged quantum entanglement. To understand this, we will study entanglement in topological systems.

First, let us review the idea of entanglement. Consider a Hilbert space $\mathcal{H}$ for some quantum mechanical system of interest. We then partition this Hilbert space into two pieces

$$
\mathcal{H} = \mathcal{H}_A \otimes \mathcal{H}_B
$$

If the overall system is in a pure state with normalized wavefunction, $|\psi\rangle$, this wavefunction can always be written as a so-called Schmidt decomposition

$$
|\psi\rangle = \sum_n \lambda_n |\psi_n^A\rangle \otimes |\psi_n^B\rangle
$$

where on the right-hand side, the wavefunctions $|\psi_n^A\rangle$ are an orthonormal set of wavefunctions spanning the Hilbert space $\mathcal{H}_A$ and the wavefunctions $|\psi_n^B\rangle$ are an orthonormal set of wavefunctions spanning the Hilbert space $\mathcal{H}_B$. The so-called “Schmidt weights”, $\lambda_n$, are non-negative real numbers. If more than one Schmidt weight is nonzero, we say that $|\psi\rangle$ is entangled between subsystems $A$ and $B$. A quantitative measure of the entanglement is given by the von Neumann entanglement entropy

$$
S_{A,B} = -\sum_n |\lambda_n|^2 \log(|\lambda_n|^2)
$$

Often these Schmidt weights are written as $\lambda_n = e^{-\xi_n^2/2}$ with the values $\xi_n$ called the entanglement energies. If one divides a $D$-dimensional system spatially into two pieces, under fairly general conditions these entanglement energies look like the spectrum of an effective Hamiltonian that lives along the $(D-1)$-dimensional cut. See for example, Li and Haldane (2008); Qi et al. (2013); [Dubail et al. (2012); Swingle and Senthil (2012).

There are other measures of entanglement entropy that can be considered, such as the Renyi entanglement entropy $S_{A,B}^{\alpha} = \frac{1}{1-\alpha} \log(\sum_n |\lambda_n|^{2\alpha})$ which matches the von Neumann entropy in the limit $\alpha \to 1$. As far as topological properties are concerned, Renyi entropies behave similarly to the von Neumann entropy. See for example, [Fidkowski et al. (2009)].
Topological Entanglement

where we have assumed that $|\psi\rangle$ is normalized such that we have

$$\sum_n |\lambda_n|^2 = 1$$

We are usually concerned with studying cases where the subsystems $A$ and $B$ are spatially local\(^5\). I.e., Hilbert space $\mathcal{H}_A$ describes degrees of freedom in region $A$ and Hilbert space $\mathcal{H}_B$ describes degrees of freedom in region $B$ such that the regions $A$ and $B$ partition a larger system, as suggested in Fig. 33.1.

It is reasonable to expect that there will be some entanglement between regions $A$ and $B$ that is proportional to the length of the boundary — some degrees of freedom near the boundary on the $A$ side interact with some degrees of freedom near the boundary on the $B$ side, and some entanglement develops between them. This type of entanglement is known as “short-ranged” since it occurs between degrees of freedom that are not physically far apart from each other. However, topologically ordered phases of matter have an additional piece of entanglement that is not short-ranged, and occurs only due to their special topological properites. To illustrate this so-called topological entanglement entropy\(^6\), we will consider the toric code as an example although the results generalize to any topologically ordered matter.

### 33.1 Entanglement in the Toric Code

Recall from chapters\(^{26, 28}\) that the ground state of the toric code can be thought of as a superposition of loop gas wavefunctions. Let us consider a particular loop gas configuration as in Fig. 33.2. The correlations (and hence the entanglement) for such a loop gas are non-local. To see this note that the number of blue lines that cross the boundary of a simply connected region (such as the orange boundary in the figure) must always be even — since if a blue line enters the region, it must also exit the region. Since the entry and exit may be very far apart from each other, this correlation is generally non-local. This correlation is essentially the origin of the non-local topological entanglement entropy.

Now let us try to evaluate the entanglement for the toric code more precisely. We will return to the lattice as in Fig. 33.3 instead of having abstract loops as in Fig. 33.2. Again the boundary between regions $A$ and $B$ is marked with an orange loop. Recall Eq. 25.14 that the total wavefunction is given by a sum over all loop configurations

$$|\psi\rangle = \sum_{\text{all loop configs}} N^{-1/2} |\text{loop config}\rangle$$

where $N$ is the total number of loop configs being summed over. For simplicity here, we have assumed that we are working on a spherical system so there is no ground state degeneracy (this will imply that some of our plaquettes are not square, but as discussed in section 25.3 this does...
not change any of the important physics). Note that the normalization is given by
\[ N = 2^{P-1} \]

where \( P \) is the number of plaquettes in the system, since flipping any plaquette gives a new loop configuration (hence \( 2^P \)), but flipping all of the plaquettes in the entire system (for a closed manifold) returns you to the original loop configuration (hence we have \( P - 1 \) rather than \( P \)).

We would now like to put Eq. 33.4 into the Schmidt form of Eq. 33.1 so that we may calculate the entanglement between the two regions. Let us cheat a bit here and first divide the system into three regions. We draw an orange line (not intersecting any vertices) to separate the system into two simply connected regions as shown in Fig. 33.3. Let region \( A \) be those edges that are entirely inside the orange line, and region \( B \) be those edges that are entirely outside of the orange line\(^7\). Here we also define a third region which we call the boundary region. The boundary region includes all of those edges that the orange line crosses through. Let the configuration of spins on this these boundary edges be called \( |\alpha\rangle \), the boundary state. If there are \( M \) boundary edges, each edge can take 2 possible states, so there are then \( 2^M \) possible boundary states \( |\alpha\rangle \).

Let us further consider the ground state wavefunction in region \( A \), given that the state of the boundary edges is given by the boundary state \( |\alpha\rangle \). We denote such a state of the region \( A \) as \( |\psi_A\rangle \alpha \), and similarly we denote the ground state wavefunction in region \( B \), having boundary state \( |\alpha\rangle \) as \( |\psi_B\rangle \). We can write these explicitly as
\[ |\psi_A\rangle = \frac{1}{\sqrt{N_A}} \sum \text{loop config} |\text{loop config}\rangle_{A,\alpha} \]

where \( N_A = 2^{P_A} \) the number of terms in the sum where \( P_A \) is the number of plaquettes entirely enclosed in region \( A \). The sum over loop configurations means that the blue lines can end only at the boundary of the region, and not in the interior (i.e., there should be no vertex defect in the interior of the region). We similarly write a wavefunction for region \( B \)
\[ |\psi_B\rangle = \frac{1}{\sqrt{N_B}} \sum \text{loop config} |\text{loop config}\rangle_{B,\alpha} \]

with \( N_B = 2^{P_B} \) the number of terms in the sum where \( P_B \) is the number of plaquettes entirely enclosed in region \( B \).

We can then write the full wavefunction as
\[ |\psi\rangle = \frac{1}{\sqrt{N}} \sum_{\text{allowed boundary states } \alpha} |\psi_A\rangle \alpha \otimes |\alpha\rangle \otimes |\psi_B\rangle \]

where \( N \) is the number of terms in the sum, which is the number of possible boundary states. The observant reader will note that on a sphere there is no well-defined “inside” or “outside”, so one arbitrarily defines one region to be called “inside”. It doesn’t matter which one.
Topological Entanglement

Eq. 33.5 is almost exactly the Schmidt decomposition form (Eq. 33.1) we would like, except that we have divided the degrees of freedom into three pieces rather than two (region A, region B and the boundary edges). It is just a matter of bookkeeping to fix this. Let us define the region $\tilde{B}$ to include both $B$ and the boundary region (all the boundary edges), we then define

$$|\psi_{\tilde{B}}\rangle = |\alpha\rangle \otimes |\psi_{\tilde{B}}\rangle$$

Region $A$ and $\tilde{B}$ now properly partition the system into two parts. The Schmidt decomposition of the wavefunction on the entire system can then be written as

$$|\psi\rangle = \frac{N_{\text{allowed}}}{2} \sum_{\text{allowed boundary states } \alpha} |\psi_{A}\rangle \otimes |\psi_{\tilde{B}}\rangle$$

where $N_{\text{allowed}}$ is the number of terms in the sum, which is the number of different boundary states.

To calculate the entanglement entropy (now between regions $A$ and $\tilde{B}$ which partition the full system) we need to know how many boundary states are in the sum. As mentioned above, if there are $M$ spins in the boundary region, there are $2^M$ possible boundary states. However, only half of these boundary states are allowed in the ground state! Again, since the ground state consists of closed loops only, we can only have an even number of blue edges in the boundary region when the full system is in the ground state. So in fact the number of terms in the sum is actually

$$N_{\text{allowed}} = 2^{M-1}$$

Plugging this result into Eq. 33.3 gives us the final result for the von Neumann entanglement entropy between the two regions

$$S_{A,B} = (M - 1) \log 2$$

As we might have predicted, the entanglement between the two regions is proportional to the length of the cut (proportional to the number of edges in the boundary region). The interesting part of this result is that the result is $M - 1$ rather than $M$. It is this subleading term (the $-1$) that reflects the topological properties of the state — the fact that it is comprised of a loop gas and there is therefore a long-ranged constraint on the state of the boundary.

It is worth noting that if region $A$ has two boundaries as in Fig. 33.4 we will get instead

$$S_{A,B} = (M - 2) \log 2$$

where $M$ is now the total number of spins on both boundaries put together. The point here is that there are now two constraints, one on the outer boundary and one on the inner boundary.

---

Footnote 8: Here we have dropped the tilde from $B$. It is implied in this formula that the union of region $A$ and region $B$ includes all the spins in the system.

---

Fig. 33.4 Partitioning a system into two regions. In this case region $A$ is inside the circle and $B$ is outside of the circle.
33.1.1 Generalizing To Arbitrary TQFTs

This technique of counting possible boundary states ([Levin and Wen 2006]) can be generalized to more general lattice models, such as the Kitaev quantum double (chapter 29) or the Levin-Wen model (chapter 31). Since all of these models are some sort of generalized loop gas, we expect that there will be similar long range correlations of the boundary states, and hence analogous topological entanglement entropy.

For any type of topological matter, if we have a smooth boundary of length $L$ between two regions $A$ and $B$, we should generically have an entanglement entropy of

$$S_{A,B} = \alpha L - \gamma + \ldots$$

for some constant $\alpha$ where $\gamma$ is the topological contributions\(^9\). For example, in Eq. 33.7 we have $\gamma = \log 2$. More generally we will obtain

$$\gamma = \log D \quad (33.9)$$

where $D$ is the total quantum dimension of the TQFT (Eq. 17.11)

$$D^2 = \sum_i d_i^2$$

with the sum over all particle species.

This is a rather remarkable result: If you are given the wavefunction of a system, by simply splitting the system into pieces and calculating the entanglement between the two pieces, you can deduce the topological properties of the system.

Two rather different proofs of the general result Eq. 33.9 are given by [Levin and Wen 2006; Kitaev and Preskill 2006]. We refer the reader to these works for the proofs.

33.2 Topological Entanglement Entropy is Robust

One might worry that the entanglement entropy we have found for the toric code is simply a particular property of the special toric code Hamiltonian we have used. Indeed, the entanglement (Eq. 33.7) will certainly change if we perturb the toric code Hamiltonian with some additional terms that change the local correlations between spins, thereby changing the prefactor $\alpha$ in Eq. 33.8. We might also wonder if, with a more general Hamiltonian, the details of the shape of the boundary (whether it is a rectangle or instead has many corners) may matter in addition to just the number of edges we cut. As emphasized in chapter 27 the topological properties of the ground state of a system should remain unchanged when the Hamiltonian is perturbed a little bit (so long as the excitation gap does not close), and this is true for the topological entanglement entropy as well. The long-ranged correlations that we discovered do re-
Topological Entanglement

Let us divide our sphere into four regions\(^{10}\) as shown in Fig. 33.5. We use the notation that \(AB\) means the union of the regions \(A\) and \(B\), and \(ABC\) is the union of regions \(A\), \(B\), and \(C\) and so forth. Now let us define the quantity

\[
S_{\text{top}} = (S_{A,BCD} + S_{B,ACD} + S_{C,ABD} + S_{D,ABC}) - (S_{AB,CD} + S_{AC,BD} + S_{AD,BC})
\] (33.10)

This looks a bit complicated, but actually it is easy to remember. The first line is the entanglements of each of the four regions \(A,B,C,D\) with their respective complementary regions (i.e., with the remainder of the sphere). The second line, added with a minus sign, are the entanglements of the union of two of these regions with the remaining two in all three possible combinations.

This combination of entanglement entropies is constructed so as to isolate only the topological term. We will find that

\[
S_{\text{top}} = -\gamma
\]

To see this let us first naively use Eq. 33.8 to calculate \(S_{\text{top}}\). Each time we calculate some \(S_{P,Q}\) we must calculate the length of the boundary between region \(P\) and \(Q\) to plug into Eq. 33.8. Looking at Fig. 33.5 there are six segments in this figure: the three straight segments pointing radially from the center (call their length \(L_{\text{straight}}\) and the three curved segments bounding region \(D\) (call their length \(L_{\text{curved}}\)). So using Eq. 33.8 we then have

\[
S_{A,BCD} = S_{B,ACD} = S_{C,ABD} = \alpha(L_{\text{curved}} + 2L_{\text{straight}}) - \gamma + \ldots
\]

\[
S_{D,ABC} = \alpha(3L_{\text{curved}}) - \gamma + \ldots
\]

\[
S_{AB,CD} = S_{AC,BD} = S_{AD,BC} = \alpha(2L_{\text{curved}} + 2L_{\text{straight}}) - \gamma + \ldots
\]

Plugging these quantities into Eq. 33.10 yields \(S_{\text{top}} = -\gamma\) as expected. All of the leading terms proportional to the length of the boundaries have cancelled. As we will see in a moment, this cancellation is extremely robust.

What happens if we alter the geometry? For example, suppose we add some wiggles in the middle of the segment forming the boundary between region \(A\) and region \(B\) as shown in Fig. 33.6 This change has no effect on regions \(C\) and \(D\), or any of the triple intersections. The change in \(S_{\text{top}}\) can be written as

\[
\Delta S_{\text{top}} = (\Delta S_{A,BCD} - \Delta S_{D,ABC}) + (\Delta S_{B,ACD} - \Delta S_{AC,BD})
\] (33.11)

It is not expected that \(S_{D,ABC}\) or \(S_{C,ABD}\) or \(S_{AB,CD}\) should change at all since in none of these cases is the entanglement measured across the main robust and are encoded into the ground state although they are a bit more tricky to see. We will now follow the procedure laid out by \(\text{[2006]}\) that described how to isolate the topological term in the entanglement entropy.

Let us divide our sphere into four regions\(^{10}\) as shown in Fig. 33.5. We use the argument and geometry of Kitaev and Preskill [2006]. A different geometry was used by Levin and Wen [2006].

\[^{10}\text{Here we use the argument and geometry of Kitaev and Preskill [2006]. A different geometry was used by Levin and Wen [2006].}\]
In Eq. 33.11 each term in parthenthesis should be zero. For example, in the first term we are comparing $S_{A,BCD}$ to $\Delta S_{AD,BC}$. These differ by whether $D$ is attached to $BC$ or attached to $A$. However, since $D$ is far away from the part of the $AB$ boundary that has been modified, the changes to the entanglement should be the same in both cases. Thus the total $\Delta S_{\text{top}}$ is zero.

We might worry that some problem could occur at the triple intersections of the three regions. However, we can argue similarly that deformation of this intersection cannot change $S_{\text{top}}$ as defined in Eq. 33.10. For example, suppose we change the geometry to that of Fig. 33.7. Again the boundaries with $D$ and the other three triple intersections are assumed to be unchanged. The total change in $S_{\text{top}}$ can be written as

$$\Delta S_{\text{top}} = (\Delta S_{A,BCD} - \Delta S_{AD,BC}) + (\Delta S_{B,ACD} - \Delta S_{AC,BD}) + (\Delta S_{C,ABD} - \Delta S_{AB,CD})$$

Again the two terms in each parenthesis differ from each other only by how region $D$ is attached, and this should not at all be changed by any deformation to boundaries between $A$, $B$, and $C$.

One can further argue that $S_{\text{top}}$ does not depend on the details of the Hamiltonian (so long as we remain in the ground state and do not close the gap to making excitations). If we change the Hamiltonian locally, this can only matter if we happen to be near a boundary. However, we just showed that we are free to move the boundaries. So we can move the boundary, change the Hamiltonian locally, and then move the boundary back to its original position, and $S_{\text{top}}$ must remain unchanged.

Thus the topological entanglement entropy is a robust property of a topologically ordered phase of matter which reflects its topological properties.

---

**Exercises**

**Exercise 33.1 Entanglement Entropy**

(a) Confirm that for a system in a pure state, the expression for the entanglement entropy $S_{AB} = -\text{Tr} [\rho_A \log \rho_A]$ given in footnote 4 agrees with Eq. 33.8.

(b) Confirm that for a system in a pure state, the expression for the Renyi entanglement entropy given in footnote 4 matches the von Neumann entropy in the limit that $\alpha \to 1$.

**Exercise 33.2 Renyi Entropy of the Toric Code**

Calculate the Renyi entanglement entropy for the toric code.

**Exercise 33.3 Entanglement Entropy of the $\mathbb{Z}_n$ Toric Code**

Generalize the calculation of section 33.1 to the case of the $\mathbb{Z}_n$ toric code. Confirm Eq. 33.9.
In chapter we explained that all topologically ordered systems have long-ranged topological entanglement. Gapped states of matter that do not have long ranged entanglement are (creatively) called short-ranged entangled states.

Recall that in section 27.2.3 we gave a rather general (and topologically motivated) definition of a topological phase of matter which we will repeat here:

**Definition of a Topological Phase of Matter:** Two (zero temperature) gapped states of matter are in the same topological phase of matter if and only if you can continuously deform the Hamiltonian to get from one state to the other without closing the excitation gap.

All (gapped) short-ranged entangled states are in the same topological phase of matter – they can all be deformed into each other without closing the excitation gap. In particular, they can all be deformed into the trivial state — a system with completely uncoupled degrees of freedom (say, spins) on each site. One might think that this makes short-ranged entangled states completely uninteresting. But in fact, the short-ranged entangled states have a beautiful connection to topology which becomes apparent once we generalize the above definition to include another aspect — symmetry.

### 34.1 Symmetry Protection

For nearly a century, symmetry has been the primary tool that physicists use to distinguish one type of matter from another. Introduced by Lev Landau\(^1\) ([Landau [1937]]), this approach has become known as the **Landau Paradigm**.

While it is too much of a digression to introduce Landau theory in detail, it is probably worth mentioning a canonical case: The ferromagnet/paramagnet transition. For example, consider a so-called “Heisenberg” magnet\(^2\). In the paramagnetic phase there is no net magnetization, whereas in the ferromagnetic phase, there is a nonzero magnetization vector. The symmetry of the system is different in the two cases: In the paramagnetic phase all directions of space are equivalent, whereas in the ferromagnetic phase there is one special direction set by the magnetization vector — we have broken the symmetry between all directions in

\(^1\)Landau was one of the most prominent physicists ever to live. Most physicists know of many of Landau’s important contributions, and due to modern specialization among fields, any given physicist is probably familiar with only a fraction of his very important works. As with so many of the greats, he was also a colorful, and difficult to control, character. Many have heard tale of how Landau crossed the Soviet establishment and ended up in prison. Fewer, perhaps, have heard that he espoused “free-love” rather than monogamy — something his wife was not so keen on.

\(^2\)The Heisenberg Hamiltonian is

\[ H = -\sum_{i,j} J_{ij} \mathbf{S}_i \cdot \mathbf{S}_j \]

with \(\mathbf{S}_i\) the spin vector for site \(i\). Here we take, say, \(J_{ij} > 0\) for neighboring spins \(i\) and \(j\) so that the energy is lower if two neighboring spins are aligned and we take \(J_{ij} = 0\) otherwise. In three dimensions there will be a finite temperature phase transition between a paramagnetic, high temperature, phase (where spins point randomly in every direction) and a ferromagnetic, low temperature, phase (where spins are mostly aligned).
space by choosing this one direction. Perhaps the most important characterization of a phase of matter is given by stating what symmetries it preserves and which symmetries it breaks.

The above definition of “Topological Phase of Matter” gives a method of classifying phases of matter, but unfortunately is blind to symmetries. An important generalization of the above definition is the idea of a symmetry protected topological phase of matter or “SPT” phase. Here we specify that a symmetry is present, and further when we deform our Hamiltonian (as in the above definition) we insist that this symmetry is never broken. This leads to a more refined classification of matter that considers both topology and symmetry.

**Definition of a Symmetry Protected Topological (SPT) Phase:** Two (zero temperature) gapped states of matter are in the same symmetry protected topological phase of matter if and only if you can continuously deform the Hamiltonian to get from one state to the other without closing the excitation gap or breaking the given symmetry.

Note that once we specify a particular symmetry we would like to consider, that two different symmetry protected topological phases (for this symmetry) typically can be smoothly deformed into each other without closing the gap if one is allowed to break the symmetry in the process.

As mentioned above, as topological phases of matter, all short-ranged entangled states are trivial (they are topologically equivalent to, or can be deformed into, the trivial state). However, once we consider symmetry protection the classification becomes more refined. There are many short-ranged entangled states that are non-trivial, i.e., that cannot be deformed into the trivial state. The cases that are non-trivial are known as SPT states, and have various interesting properties — particularly including protected edge modes that are either gapless or degenerate.

### 34.1.1 On-Site Symmetries

There are many types of symmetries one might be concerned with (and for each symmetry there is classification of phases of matter into symmetry protected topological phases). In chapter ** below, we will briefly discuss time-reversal symmetry, which is perhaps the most important example experimentally.

In this chapter instead we will consider so-called on-site symmetry. We imagine an $N$-state Hilbert space on each site (a $N$-state qudit). We can write the states in the Hilbert space at site $i$ as

$$|m\rangle_i \quad \text{for} \quad m \in 1 \ldots N$$

We can then consider unitary operations on each site

$$\hat{U}^{(i)} = \sum_{mn} |m\rangle_i U_{mn} |n\rangle_i$$

(34.1)
A global unitary symmetry operation is given by applying the same $\hat{U}^{(i)}$ at every site in the system

$$\hat{U}_{\text{global}} = \prod_j \hat{U}^{(i)}$$

If the ground state wave function is unchanged after application of this global unitary symmetry operation, then we say that $\hat{U}$ is a member of the symmetry group. Obviously if there are two different unitaries $\hat{U}$ and $\hat{U}'$ in the symmetry group, then their product is also in the symmetry group (hence we use the mathematical term “group” here). The $U$ matrices in Eq. 34.1 are formally representation matrices for a group which we will call $G$.

### 34.2 Example: Ising Symmetry

Let us consider the simplest possible example: Ising symmetry\(^3\). Recall that the Ising model is a model where spins are allowed to point either up or down only. Ising symmetry means a system is unchanged if you flip all up spins to down and all down spins to up. We will here consider the possible SPT phases of a system with Ising symmetry.

Consider a triangular lattice with a spin-1/2 at each vertex as shown in Fig. 34.1. Using Pauli spin operators, $\sigma_x^{(i)}$ flips over the spin at site $i$. We can thus consider our global symmetry operation to be

$$\hat{U}_{\text{global}} = \prod_i \sigma_x^{(i)}$$

This is the only symmetry we want to consider for the Ising case. If you square $\hat{U}_{\text{global}}$ you get the identity, so the symmetry group here is actually $\mathbb{Z}_2$.

We claim there are exactly two SPT phases possible that obey this Ising symmetry. The corresponding Hamiltonians are given by the following

$$H_{\text{trivial}} = -\sum_i \sigma_x^{(i)}$$

\(^3\)Caution: this has essentially nothing to do with the Ising TQFT.
Part VIII

Introduction to Quantum Hall Effects
Introduction to Quantum Hall
— The Integer Effect

The fractional quantum Hall effect is the best studied of all topologically ordered states of matter. In fact it is the only system which is extremely convincingly observed to be topologically ordered in experiment\(^1\). We will thus spend quite a bit of time discussing quantum Hall effects in detail. Before we can discuss fractional quantum Hall effect we need to discuss the basics, i.e., the integer quantum Hall effect.

35.1 Classical Hall Effect

In 1879 Edwin Hall discovered that when a current is run perpendicular to a magnetic field, a voltage is generated perpendicular to both field and current, and proportional to both (See Fig. 35.1). This voltage is now known as the Hall voltage. Drude theory, treating a metal as a gas of electrons, explains the Hall voltage as being a simple result of the Lorentz force on electrons.

![Fig. 35.1][1] Hall voltage \(V_H\) perpendicular to both magnetic field and current, and proportional to both. Also one measures a longitudinal voltage in the same direction as the current, roughly independent of magnetic field.

35.2 Two-Dimensional Electrons

In the late 1960s and early 70s semiconductor technology made it possible to do experiments with electrons that live in two dimensions. First

\(^{1}\)There are a good number of other contenders now. Probably the most convincing other case is \(^3\)HeA phase 2d films. Although very few experiments have actually been done on this. Other strong contenders include Majorana wires, certain exotic superconductors, and a few frustrated quantum spin systems.
MOSFETs\(^2\) and later quantum wells were used to provide a confining potential for electrons in one direction\(^3\), leaving motion only in the two remaining dimensions. As an example we will consider a quantum well structure, which is layered in the \(\hat{z}\) direction as shown in Fig. 35.2.

**Fig. 35.2** Top A quantum well structure is a quasi-two-dimensional layer of one semiconductor sandwiched between two other semiconductors. Bottom The potential felt by an electron is like a particle in a box. If the energy is low enough, the electron is stuck in the lowest particle-in-box wavefunction \(\varphi_0(z)\) giving a total wavefunction \(\Psi = \varphi_0(z) \psi(x, y)\) and having strictly two dimensional motion.

The electron moving in the \(z\)-direction experiences a strong confinement, such as the particle-in-box confinement shown in Fig. 35.2. The wavefunction of the electron then takes the form \(\varphi(z)\) in the \(z\)-direction. If the energy (i.e. the temperature and coulomb interaction) is very low compared to the gap between the particle-in-box states, then the electron is frozen in the lowest particle-in-box state \(\varphi_0(z)\) and the total wavefunction of the electron is \(\Psi(x, y, z) = \varphi_0(z) \psi(x, y)\) leaving only the \(x\) and \(y\) degrees of freedom. Thus we have a strictly two dimensional electron.

More recently two dimensional electronic systems have also been observed in single-layer atomic systems such as graphene. (Although even then, the same argument needs to be used — that the motion of the electron is “frozen” in the \(z\)-direction and only has freedom to move in \(x\) and \(y\).)
35.3 Phenomenology of Integer Quantum Hall Effect

In 1980 Klaus von Klitzing, having just left a postdoctoral position at Oxford, went to a new job at Grenoble carrying some new high mobility two dimensional electron samples grown by (now Sir) Michael Pepper at Cambridge. He put them in high magnetic field and cooled them down to a few degrees Kelvin temperature where he discovered something very different from what Hall had seen a hundred years earlier. An example of this type of experiment is shown in Fig. 35.3.

Fig. 35.3 An example of an Integer Quantum Hall experiment. The plateaus in $V_H$ are such that $V_H = (1/i)(h/e^2)I$ with $i$ the integer displayed over the plateau — where $h$ is Planck’s constant and $e$ is the electron charge. At the same magnetic field where a plateau occurs in $V_H$ the longitudinal voltage drops to zero. Note that at very low field, the Hall voltage is linear in $B$ and the longitudinal voltage is independent of $B$, as would be predicted by Drude theory.

At low magnetic field, the longitudinal voltage is relatively constant whereas the Hall voltage is linear in magnetic field — both of these are precisely what would be predicted by Drude theory. However, at high magnetic field, plateaus form in the Hall voltage with concomitant zeros of the longitudinal voltages. The plateaus have precisely the value

$$V_H = \frac{1}{i} \frac{h}{e^2} I$$

4 Meaning very clean
where $I$ is the current, $h$ is Planck’s constant and $e$ is the electron charge. Here $i$ is an integer as shown in the figure. Or equivalently we have

$$R_H = \frac{1}{i} \frac{h}{e^2} = 1/G_H \quad (35.1)$$

with $R_H$ the Hall resistance where $G_H$ the Hall conductance. Where we have plateaus in the Hall voltage, we have zeros in the longitudinal voltage and resistance

$$R_L = 0$$

which implies we have a dissipationless state — similar to a superfluid. These statements become increasingly precise as the temperature is lowered.

We should remember that conductivity and resistivities are both 2 by 2 matrices and are inverses of each other\(^5\). In this quantum Hall state, these matrices are both purely off-diagonal. Thus we have the interesting situation that both the diagonal part of the conductivity (the longitudinal conductivity) is zero, and the diagonal part of the resistivity (the longitudinal resistivity) is also zero.

The plateau $R_H = (1/i)(h/e^2)$ occurs near the magnetic field such that the so-called filling fraction ratio

$$\nu = \frac{n\phi_0}{B}$$

is roughly the integer $i$. Here $n$ is the 2d electron density and $\phi_0$ is the quantum of magnetic flux

$$\phi_0 = h/e$$

When von Klitzing discovered this effect he noticed mainly that the plateaus in the Hall resistance are extremely precisely given by Eq.\(^{35.1}\) and the plateaus are extremely flat. He submitted his manuscript to PRL claiming that this would be a useful way to make a new resistance standard\(^6\).\(^7\). In fact the result has been shown to be precise and reproducible to better than a part in $10^{10}$. This is like measuring the distance from London to Los Angeles to within a fraction of a millimeter. This accuracy should be extremely surprising. The samples are dirty, the electrical contacts are soldered on with big blobs of metal, and the shape of the sample is not very precisely defined.

\(^6\)The referee mentioned that at the time they already had resistance standards which were better than his initial measurement of one part in $10^{6}$, but proposed would be a uniquely good measurement of the ratio $h/e^2$. The paper was resubmitted proposing to use the effect as a precise measurement of the fine structure constant. The paper was accepted and the Nobel Prize for von Klitzing followed in 1985.

\(^7\)The quantum Hall effect is used as a metrological resistance standard, and it is proposed that the Ohm will soon be defined in terms of the result of quantum Hall experiments.

### 35.4 Transport in Zero Disorder

In strictly zero disorder it is easy to show that the longitudinal resistance is zero and the Hall resistance is precisely linear in the magnetic field. This is a simple result of Galilean/Lorentz invariance. Suppose we have a two dimensional disorder-free system of electrons in the $x,y$ plane and a magnetic field $\mathbf{B} = B\hat{z}$ in the $\hat{z}$-direction perpendicular to the plane.
The Lorentz force on an electron will be

\[ \mathbf{F} = -e(\mathbf{E} + \mathbf{v} \times \mathbf{B}) \]

If we then boost into a moving frame where

\[ \mathbf{v} = \frac{\mathbf{E} \times \mathbf{\hat{z}}}{|B|} \]

in this new frame we obtain \( \mathbf{F} = 0 \), so the ground state must be stationary in this frame.

Then we boost back into the lab frame, and we obtain a current

\[ \mathbf{j} = -e\mathbf{v} = -\frac{en \mathbf{E} \times \mathbf{\hat{z}}}{|B|} \]

thus giving us

\[ R_L = 0 \]
\[ R_H = \frac{B}{ne} \]

which is exactly the prediction that Drude would have made for a disorder free system.

While this calculation is rigorous even with the effects of quantum mechanics and interactions, it relies on having strictly zero disorder.

35.5 The Landau Problem

In order to understand quantum Hall effect, we should start by understanding the physics of a charge particle in a Magnetic field — a problem first studied by Landau. For simplicity we assume our electrons are spinless (indeed, the spins tend to be polarized by the magnetic field anyway.) We will consider an electron in the \( x,y \) plane, with a magnetic field of magnitude \( B \) in the \( z \) direction. We will assume the system is periodic in the \( y \) direction with length \( L_y \), but open in the \( x \) direction, with length \( L_y \) (i.e., we are working on a cylinder actually). We will eventually consider a small amount of disorder (as we showed above this is crucial!), but for now let us assume the system has no disorder.

The Hamiltonian is

\[ H_0 = \frac{(\mathbf{p} + e \mathbf{A})^2}{2m} \]

where \( e \) and \( m \) are the electron charge and mass, and \( \mathbf{A} \) is the vector potential. We then have to choose a particular gauge to work in. Later on we will want to work in symmetric gauge (there is a homework problem on this!) For now we will work in the so-called “Landau” gauge

\[ \mathbf{A} = B x \mathbf{\hat{y}} \]
which does indeed satisfy

$$B = \nabla \times A = B \hat{z}$$

as desired. The Hamiltonian is thus

$$H_0 = \frac{1}{2m} \left( (p_2^2 + (p_y + eBx)^2) \right)$$

where $p_j = -i\hbar \partial_j$.

The Hamiltonian is then translationally invariant in the $\hat{y}$ direction, so we can write the wavefunction as

$$\psi(x, y) = \phi_{k_y}(x) e^{ik_y y}$$

and due to the periodicity in the $y$-direction, we have

$$k_y = \frac{2\pi n}{L_y}$$

for some integer $n$. Plugging in this form gives a familiar Schrödinger equation

$$\left( \frac{p_x^2}{2m} + \frac{1}{2} m \omega_c^2 (k_y \ell^2 + x)^2 \right) \phi_{k_y}(x) = E \phi_{k_y}(x) \quad (35.2)$$

where $\ell$ is the so-called magnetic length

$$\ell = \sqrt{\hbar/(eB)}$$

and $\omega_c$ is the cyclotron frequency

$$\omega_c = eB/m.$$

We recognize this Schrödinger equation as being just a harmonic oscillator where the center of the harmonic potential is shifted to $x = -k_y \ell^2$. Thus the eigenenergies are of the usual harmonic oscillator form

$$E_p = \hbar \omega_c \left( p + \frac{1}{2} \right) \quad (35.3)$$

where $p$ is an integer. These quantized energy states are known as Landau levels. The form of the wavefunction will be harmonic oscillator on the $x$ direction and plane-wave in the $y$-direction as shown in Fig. 35.4.
Fixing the energy by fixing $p$ in Eq. 35.3, the value of $k_y$ is quantized in units of $2\pi/L_y$. Further, the position $x$ ranges over $L_x$, meaning that $k_y$ ranges over $L_x/\ell^2$. Thus the total number of possible values of $k_y$ is

$$\text{Number of states in a Landau level} = \frac{L_x L_y}{2\pi \ell^2} = \frac{\text{Area} \, B}{\phi_0}$$

where

$$\phi_0 = h/e$$

is the magnetic flux quantum. Thus, the number of states in a Landau level is equal to the number of magnetic flux quanta of magnetic field incident on the plane.

We can plot the density of states for electrons in a magnetic field, as shown in Fig. 35.5.

When there are multiple electrons present, we define the filling fraction to be the number of these Landau levels which are completely filled with electrons.

$$\nu = \frac{n\phi_0}{B}$$

where $n$ is the density of electrons. Or equivalently we can write a relationship between the number of electrons in the system, $N_e$ and the number of magnetic flux $N_\phi$

$$N_e = \nu N_\phi$$

**Incompressibility of Integer Number of Filled Landau Levels:**

When some integer number of Landau levels is filled, the chemical potential lies in the middle of the gap between the filled and unfilled states — analogous to a band insulator. In this case the the system is *incompressible*. This means there is a finite energy gap to creating any excitations.
Introduction to Quantum Hall — The Integer Effect

— i.e., all excitations must involve removing an electron from a filled Landau level, promoting it above the energy gap to place it in an empty state. In particular excitations which change the density (compressions) are gapped. Further, at this precise integer filling fraction, the longitudinal conductivity is zero, and the Hall conductivity is precisely the quantized value $R_H = ne/B = (1/i)(\hbar/e^2)$.

If we were to control the chemical potential in the experiment, we would have our answer as to why the Hall conductivity shows plateaus — for any value of the chemical potential except for the special values $\mu = (\hbar\omega_c)(p + 1/2)$ with integer $p$, the electron number is pinned to $N = N_0/i$ where $i$ is an integer, precisely $i$ Landau levels are filled, there is a gap to excitations, and the Hall conductivity would be precisely quantized. However, in real experiments, it is actually the density that is fixed — which means that generically the chemical potential $\mu$ sits in the degenerate band $\mu = (\hbar\omega_c)(p + 1/2)$ for some integer $p$ and generically the filling fraction is tuned continuously and is not quantized.

Thus the incompressible state is very fine tuned. It occurs only for a very precise (integer) value of the filling fraction — for all other values of the filling fraction, some Landau level is partially filled and (at least neglecting interactions) the system would be extremely compressible, as there are many zero energy excitations corresponding to rearrangements of the electrons (which orbitals are filled and which are empty) within the partially filled Landau level.

While the system does have a gap under fine tuning, we will need something that will preserve the special properties of the fine tuned state even when we move away from the filling fraction which is precisely an integer. What does this is actually disorder — it will provide a reservoir for excess electrons (or holes) added (or subtracted) from the integer filled state. With disorder, the special properties of the quantized state are made robust.

What Does Disorder Do?

As mentioned above, we will need to add disorder to the system in order to achieved quantized Hall effect. What is the effect of this disorder? Disorder will spread out the energies in the band by having some regions where the potential is higher than average and some regions where the potential is lower than average. This spreads the sharp peak in the density of states into a broader band, as shown in Fig. 35.6.

Since current tends to flow perpendicular to potential gradients (i.e., it is hall current), eigenstates tend to follow contours of constant potential. Thus many of the eigenstates at high and low energy will be trapped in local minima or maxima — isolated in a hill or valley and circling the peak or bottom. The result is that the eigenstates in the edge of the band experience localization, whereas (at least some) eigenstates near the center of the band as shown in Fig. 35.6.

When the chemical potential is anywhere in the localized states, then at low enough temperature, the electrons cannot move at all. Although

Fig. 35.6 The density of states for spin-polarized (or spinless) electrons in a magnetic field with disorder. The Landau bands are spread out, with localized eigenstates in the tails and extended eigenstates near the middle.
there are states at this energy, they are all localized and electrons cannot jump between them. Hence we expect in this case that the DC dissipative conductance goes to zero. (For dissipitive conductance to occur, an electron has to be excited up to the next delocalized band.) The state remains incompressible for filling fractions even away from the precise integer value of $\nu$.

What is not obvious is (a) that the Hall conductance should be precisely quantized, and (b) that we should have Hall conductance at all.

35.6 Laughlin’s Quantization Argument

In 1981, shortly after von Klitzing’s discovery of quantum Hall effect, Bob Laughlin\(^8\) presented an argument as to why the Hall conductance must be precisely quantized. The argument relies on gauge invariance. We first need to present a key theorem which comes from gauge invariance.

35.6.1 Byers and Yang Theorem

Consider any system (made of electrons and protons and neutrons) with a hole cut in it, as in Fig. 35.7. Now put some magnetic flux $\Phi$ through the hole in such a way that the flux does not touch any piece of the system, but just goes through the hole. By the Aharanov-Bohm effect, the charged particles in the system cannot detect the flux if it is an integer multiple of the flux quantum $\phi_0$. In fact the statement can be made stronger: The eigenspectrum of the system is precisely the same when an integer number of flux is inserted through the hole. This result is known as the Byers\(^9\)-Yang\(^10\) theorem (1961).

To prove this theorem we use gauge invariance. One is always free to make a gauge transformation

$$A'(r) = A(r) + (h/e)\nabla \chi(r)$$

$$\Psi'(r_1, \ldots, r_N) = \prod_{j=1}^{N} e^{i\chi(r_j)} \Psi(r_1, \ldots, r_N)$$

which leave the physical electromagentic field completely unchanged and changes the gauge of the wavefunction. The meaning of gauge invariance is that if we have a solution to the Schroedinger equation for $\Psi$ and $A$ at energy $E$, then we also have a solution at the same energy $E$ for $\Psi'$ and $A'$.

When the physical geometry we are concerned with is non-simply connected, we can make gauge transforms which are non-single-valued, such as

$$\chi(r) = m\theta(r)$$

where $\theta$ is the angle around the center. Making this gauge transform leaves the eigenspectrum of the system unchanged. However, the flux

---

\(^8\)Laughlin would later go on to win a Nobel Prize for his explanation of fractional quantum Hall effect, which we will start discussing in chapter ***.

\(^9\)Nina Byers was just starting as an assistant professor at UCLA when she proved this theorem. In the late 60s and early 70s she oscillated between Oxford (Somerville college) and UCLA, but eventually converged to UCLA. She told me personally that she regretted leaving Oxford. She passed away in 2014.

\(^10\)Yang is C.N.Yang, who won a Nobel Prize in 1957 along with T. D. Lee for his prediction of parity non-conservation of the weak interaction.
enclosed
\[ \Phi' = \oint A' \cdot dl = \oint A \cdot dl + 2\pi m\hbar/e = \Phi + m\phi_0 \]

has changed by an integer number of flux quanta.

### 35.6.2 Quantization of Hall Conductance

Laughlin’s argument applies the Byers-Yang theorem to the Quantum Hall case. Consider a two-dimensional electron system cut in an annulus\(^\text{11}\) as shown in Fig. 35.8. Here we put the entire system in a uniform magnetic field (so that we have Landau levels) and we arrange such that the chemical potential is in the localized part of the band so that at low enough temperature the longitudinal (dissipitive) conductivity is zero.

We then adiabatically insert an additional flux \(\Phi(t)\) through the center of the annulus and turn it on slowly from zero to one flux quantum. Due to the Faraday’s law, an EMF is generated around the annulus
\[ E = -\frac{d\Phi}{dt} = \oint dl \cdot E \]

If there is a Hall conductance, \(G_H\) then this generates a radial current
\[ J = G_H E \]

As we slowly increase the flux by an amount \(\Delta\Phi\) we have a total charge \(\Delta Q\) moved from the inside to the outside of the annulus given by
\[ \Delta Q = \int dt J(t) = G_H \int dt E(t) = -G_H \int dt \frac{d\Phi(t)}{dt} = -G_H \Delta \Phi \]

Now the key to the argument is the Byers-Yang theorem. If we choose \(\Delta \Phi = \phi_0\) a single flux quantum, then the final eigenstates of the system must be precisely the same as the initial eigenstates of the system. Since we have changed the system adiabatically (and there is a gap to excitations when the states at the chemical potential are localized due to disorder) the system must stay in the ground state\(^\text{12}\) and the insertion of the flux quantum must take us from the ground state back to the very same ground state. The only thing that might have changed during this process is that an integer number \(p\) of electrons may have been transferred from the inside of the annulus to the outside. Thus we have
\[ -pe = \Delta Q = -G_H \Delta \Phi = -G_H \phi_0 = -G_H (h/e) \]

Thus we obtain the quantized Hall conductance
\[ G_H = p(e^2/h) \]

with \(p\) an integer!

Thus we see that the Hall conductance experiment is really some sort

---

\(^{11}\)For studying current flow in magnetic fields, the annulus is known as "Corbino" geometry, after O. M. Corbino, who studied this in 1911.

\(^{12}\)There is a subtly here. With disorder, there are actually low energy excitations, but they require very long range hops of localized electrons which cannot be made. So the system is “locally” gapped.
of "spectroscopy" to measure the charge on the electron! (hence the precision of the effect).

Although we have shown the the Hall conductance must be quantized, what we have not shown is that it must be nonzero! Afterall, since the chemical potential is in a localized band, it looks like electrons simply can’t move at all. We will return to this issue in section 35.8 below.

### 35.7 Edge States

The bulk of a quantum Hall system is gapped, but on a finite system there are always low energy modes on the edges. (This is always true for any chiral topological system. Although achiral systems can have fully gapped edges). Even though the bulk is incompressible, the shape of the edge can be deformed as suggested in Fig. 35.9. Now let us think about the dynamics of a bump on the edge. On the edge of the system we always have an electric field (this is the potential that holds the electrons in the system — otherwise they would just leak out!). Since we have $E \times B$, we expect a drift velocity for all the electrons on the edge. Thus we expect edge dynamics to be basically just movement of charge along the edge.

#### 35.7.1 Landau Gauge Edge Picture for Integer Quantum Hall

Recall in Landau gauge (See section 35.5) the wavefunctions are plane waves in the $y$ direction, but are harmonic oscillator states in the $x$ direction. We now impose an additional confining potential in the $x$ direction near the edges of the system as shown in Fig. 35.10.

The addition of the confining potential $V(x)$ simply adds this potential to the 1-d Schroedinger equation 35.2. If the confining potential is fairly smooth, it simply increases the energy of the eigenstates when the position $x = -k_y \pi$ gets near the edge of the system as shown in Fig. 35.10.

In the case of the integer quantum Hall effect, all of the eigenstates of
some particular Landau level (the lowest Landau level in the figure) are filled within the bulk. At some point near the edge, the Landau level crosses through the chemical potential and this defines the position of the edge. Since the eigenstates are labeled by the quantum number $k_y$, it is possible to create a low energy excitation by moving an electron from a filled state near the edge just below the chemical potential to an empty state near the edge just above the chemical potential. The excitation will have momentum $\hbar \Delta k_y$. We thus have a 1-d system of fermions filled up to a chemical potential and they flow only in one direction along each edge — i.e., they are chiral fermions.

### 35.8 The Halperin Refinement of Laughlin’s Argument

A more careful version of Laughlin’s argument was made by Halperin immediately after Laughlin’s initial work. The key here is to think of a geometry where much of the system is free of disorder. In particular we consider the geometry shown in Fig. [35.11](#).

---

$^{13}$The change in energy will be

$$\Delta E = \frac{\partial V}{\partial x} \Delta x = \frac{\partial V}{\partial x} \ell^2 \Delta k_y$$

Thus the edge velocity is given by

$$v = \frac{1}{\hbar} \frac{\partial E}{\partial k} = \frac{1}{\hbar} \frac{\partial V}{\partial x} \ell^2$$

If the chemical potential along the one edge is raised by $\Delta \mu$, a range of $k$-states

$$\Delta k = \frac{\Delta \mu}{e^2 \ell^2}$$

will be filled. Since the spacing between adjacent $k$ states is $2\pi/L_y$ this corresponds to an increase in electrons per unit length along the edge of

$$\Delta n_{1d} = \frac{2\pi \Delta \mu}{e^2 \ell^2}$$

These then carry a net 1d electron current density

$$j = -ev\Delta n_{1d} = -e \left( \frac{1}{\hbar} \frac{\partial V}{\partial x} \right) \frac{2\pi \Delta \mu}{e^2 \ell^2 \frac{\partial V}{\partial x}} = -(e/\hbar) \Delta \mu$$

which is precisely the expected quantized Hall current flowing along the edge. ($\Delta \mu = -e\Delta V$).
Fig. 35.11  The Halperin geometry. The same as the Laughlin annulus geometry, except here we add disorder only in part of the annulus. We have also shown (dark blue) a single particle eigenstate in the clean region, which forms a circle (with a small gaussian cross-section).

Here, the disorder is confined to only part of the annulus, the inner-most and outer-most regions of the annulus being disorder-free. Within the clean regions we can solve for the eigenstates using symmetric gauge (this is a homework problem, but we will also discuss further in the next chapter). The eigenstates are indexed by their angular momentum $m$, and in the Lowest Landau level, for example, they are given by

$$\phi_m \sim z^m e^{-|z|^2/(4\ell^2)}$$

where $z = x + iy$ is the complex representation of the position. A radial cut of one of these eigenstates gives a gaussian wavepacket\(^{14}\) at radius $\ell\sqrt{2m}$— very similar to what we had in Landau gauge, but now these eigenstates are indexed by angular momenta instead of linear momenta, and they go around in circle instead of going straight.

Let us imagine the chemical potential above the middle of a Landau level (say above the middle of the lowest Landau level) until it sits in a localized piece (at least within the disordered region the wavefunctions are localized). Since this is above the middle of the Landau level, the Landau level is completely filled in the clean region. The only low energy excitations are the edge states!

Now, let us track what happens to the eigenstates as we change the flux through the hole. If the flux through the hole is an integer (in units of the flux quantum $\phi_0$), then the angular momentum is also an integer. However, if the flux through the hole is an integer plus some fraction $\alpha$, then the angular momentum quantum number must also be an integer plus $\alpha$. Thus, as we adiabatically increase the flux by one flux quantum, we adiabatically turn each $m$ eigenstate to $m + 1$. Thus

\(^{14}\)Just find the maximum of $|\psi_m|^2$. 

we are continuously pushing out electrons to the next further out radial wavefunction.

Now when we are in the disordered region of the annulus, we do not know any details of the shape of the eigenstates. All we know is that after insertion of a full flux quantum we must get back to the same many body eigenstate that we started with. However, we also know that an additional electron is being pushed into the disordered region from the clean region on the inside, whereas an electron is also being extracted into the clean region on the outside. Thus the disordered region must also convey exactly one electron (per Landau level) when a flux quantum is inserted adiabatically. An electron state is moved from one edge state on the inside to an edge state on the outside.

This argument pins down that the Hall conductance is not zero, but is $\frac{h}{e^2}$ times the number of Landau levels that are filled (in the clean regions).

## Exercises

**Exercise 35.1 Quantum Hall Conductivity vs Conductance**

Consider a two dimensional electron gas (2DEG) of arbitrary shape in the plane with four contacts (1,2,3,4) attached at its perimeter in a clockwise order as shown in Fig. 35.12. The conductivity tensor $\sigma_{ij}$ relates the electric field to the current via

$$j_i = \sigma_{ij} E_j$$

(35.4)

where indices $i$ and $j$ take values $\hat{x}$ and $\hat{y}$ (and sum over $j$ is implied). Assume that this is a quantized hall system with quantized hall conductance $s$. In other words, assume that

$$\sigma = \begin{pmatrix} 0 & s \\ -s & 0 \end{pmatrix}$$

(35.5)

Show that the following two statements are true independent of the shape of the sample.

(a) Suppose current $I$ is run from contact 1 to contact 2, show that the voltage measured between contact 3 and 4 is zero.

(b) Suppose current $I$ is run from contact 1 to contact 3, show that the
voltage measured between contact 2 and 4 is \( V = I/s \).

Note: The physical measurements proposed here measure the conductance of the sample, the microscopic quantity \( \sigma \) is the conductivity.

**Exercise 35.2 About the Lowest Landau Level**

If you have never before actually solved the problem of an electron in two dimensions in a magnetic field, it is worth doing. Even if you have done it before, it is worth doing again.

Consider a two dimensional plane with a perpendicular magnetic field \( \vec{B} \). Work in symmetric gauge \( \vec{A} = \frac{1}{2} \vec{r} \times \vec{B} \).

(a) (This is the hard part, see below for hints if you need them.) Show that the single electron Hamiltonian can be rewritten as

\[
H = \hbar \omega_c (a^\dagger a + \frac{1}{2})
\]

where \( \omega_c = eB/m \) and

\[
a = \sqrt{2} \ell \left( \bar{\partial} + \frac{1}{4\ell^2} \bar{z} \right)
\]

with \( z = x + iy \) and \( \bar{\partial} = \partial/\partial \bar{z} \) with the overbar meaning complex conjugation. Here \( \ell \) is the magnetic length \( \ell = \sqrt{\hbar/eB} \).

(b) Confirm that \( [a, a^\dagger] = 1 \)

and therefore that the energy spectrum is that of the harmonic oscillator

\[
E_n = \hbar \omega_c (n + \frac{1}{2})
\]

(c) Once you obtain Eq. (35.6) show that any wavefunction

\[
\psi = f(z)e^{-|z|^2/4\ell^2}
\]

with \( f \) any analytic function is an eigenstate with energy \( E_0 = \frac{1}{2}\hbar \omega_c \). Show that an orthogonal basis of wavefunctions in the lowest Landau level (i.e., with eigenenergy \( E_0 \)) is given by

\[
\psi_m = N_m z^m e^{-|z|^2/4\ell^2}
\]

where \( N_m \) is a normalization constant. Show that the maximum amplitude of the wavefunction \( \psi_m \) is a ring of radius \( |z| = \ell\sqrt{2m} \) and calculate roughly how the amplitude of the wavefunction decays as the radius is changed away from this value.

(d) Defining further

\[
b = \sqrt{2} \ell \left( \partial + \frac{1}{4\ell^2} \bar{z} \right)
\]

with \( \partial = \partial/\partial z \), Show that the operator \( b \) also has canonical commutations

\[
[b, b^\dagger] = 1
\]

but both \( b \) and \( b^\dagger \) commute with \( a \) and \( a^\dagger \). Conclude that applying \( b \) or \( b^\dagger \) to a wavefunction does not change the energy of the wavefunction.

(e) show that the \( \hat{z} \) component of angular momentum (angular momentum perpendicular to the plane) is given by

\[
L = \hat{z} \cdot (\vec{r} \times \vec{p}) = \hbar (b^\dagger b - a^\dagger a)
\]
Conclude that applying $b$ or $b^{\dagger}$ to a wavefunction changes its angular momentum, but not its energy.

(f) [Harder] Let us write an arbitrary wavefunction (not necessarily lowest Landau level) as a polynomial in $z$ and $\bar{z}$, times the usual gaussian factor. Show that projection of this wavefunction to the lowest Landau level can be performed by moving all of the $\bar{z}$ factors all the way to the left and replacing each $\bar{z}$ with $2\ell^2 \partial_x$.

Hints to part a: First, define the antisymmetric tensor $\epsilon_{ij}$, so that the vector potential may be written as $A_i = \frac{1}{2} B \epsilon_{ij} r_j$. We have variables $p_i$ and $r_i$ that have canonical commutations (four scalar variables total). It is useful to work with a new basis of variables. Consider the coordinates

\[
\pi_i^{(\alpha)} = p_i + \alpha \frac{\hbar}{2\ell^2} \epsilon_{ij} r_j
\]

and

\[
\frac{\hbar}{\ell^2} \epsilon_{ij} \xi_j
\]

defined for $\alpha = \pm 1$. Here $\alpha = +1$ gives the canonical momentum. Show that

\[
\left[ \pi_i^{(\alpha)}, \pi_j^{(\beta)} \right] = i\alpha \epsilon_{ij} \delta_{\alpha \beta} \frac{\hbar^2}{\ell^2}
\]

The Hamiltonian

\[
H = \frac{1}{2m} (p_i + eA_i) (p_i + eA_i)
\]

can then be rewritten as

\[
H = \frac{1}{2m} \pi_i^{(+1)} \pi_i^{(+1)}
\]

with a sum on $i = \hat{x}, \hat{y}$ implied. Finally use

\[
a = \frac{-\pi_y^{(+1)} + i\pi_x^{(+1)}}{\sqrt{2\hbar}} \ell
\]

and

\[
b = \frac{\pi_y^{(-1)} + i\pi_x^{(-1)}}{\sqrt{2\hbar}} \ell
\]

to confirm that $a$ and $b$ are given by Eqs. 35.7 and 35.12 respectively. Finally confirm Eq. 35.6 by rewriting Eq. 35.19 using Eqs. 35.20 and 35.21.

A typical place to get confused is the definition of $\partial$. Note that

\[
\partial z = \partial \bar{z} = 1
\]

\[
\partial \bar{z} = \partial z = 0
\]

Hints to part f: Rewrite the operators $a, a^{\dagger}, b, b^{\dagger}$ such that they operate on polynomials, but not on the Gaussian factor. Construct $\bar{z}$ in terms of these operators. Then project.
Aside: A Rapid Introduction to Topological Insulators

The integer quantum Hall effect is one of the simplest examples of what is now called a “topological insulator”. To explain what this is, and why it is interesting, let us review some basic facts about band structure and non-interacting electrons.¹

36.1 Topological Phases of Matter

We will consider systems of electrons in some periodic environment — which is what an electron would experience in a real material crystal². We can thus describe our system as some single electron kinetic energy and some periodic potential — or equivalently as some tight-binding model. Bloch’s theorem tells us that the eigenstates of such a periodic Hamiltonian can be written in the form

$$|\Psi^\alpha_k\rangle = e^{i\mathbf{k} \cdot \mathbf{r}} |u^\alpha_k\rangle$$

where $\alpha$ is the band index, and $u^\alpha_k(x)$ is a function periodic in the unit cell.

The eigen-spectrum breaks up into bands of electron states. If a (valence) band is completely filled and there is a gap to next (conduction) band which is empty, we generally call the system a band insulator. The conventional wisdom in most solid state physics books is that such band insulators carry no current. This wisdom, however, is not correct. A prime example of this is the integer quantum Hall effect! As we have just seen for the integer quantum Hall effect we have a filled band and a gap in the single electron spectrum. And while such a system carries no longitudinal current (and correspondingly has $\sigma_{xx} = 0$) it does carry Hall current with $\sigma_{xy} = ne^2/h$.

One might object that the integer quantum Hall effect is not really a valid example, because it does not have a periodic potential. However, it is certainly possible to add a very weak periodic potential to the quantum Hall system and maintain the gap.

It turns out that there is a topological distinction in the wavefunctions for the quantum Hall effect versus what we think of as a traditional band insulator. One way to describe this is to think of the band structure as being a mapping from the Brillouin zone (inequivalent values of $k$) to

¹In this chapter we are thinking about non-interacting electrons in periodic potentials!

²Some of the ideas discussed here do not depend too much on the system being precisely periodic.
the space of possible wavefunctions

\[ k \rightarrow u_k^\alpha(x). \] (36.1)

Once we have such a mapping we can ask about whether there are topologically different mappings, or whether one mapping can be continuously deformed to another.

An analogy is to consider a mapping from a circle \( S^1 \) to a circle \( S^1 \),

\[ e^{i\theta} \rightarrow e^{if(\theta)} \]

Here, one can topologically classify the mapping by its winding number. One such mapping cannot be continuously deformed into another if the two mappings have different winding numbers.

Similarly, we can define a “winding number” (known as a “Chern” number) of the band structure map Eq. 36.1 for two-dimensional systems. This integer topological quantity turns out to be precisely the quantized Hall conductance in units of \( e^2/h \). We give an explicit expression for this quantity in section *** below. Similar topological definitions of “winding numbers” of the map Eq. 36.1 can be given in any dimension.

If we imagine continuously changing the physical Hamiltonian, this Chern number, which must be an integer, cannot change continuously. It can only change by making it impossible to define a Chern number. This happens when if the system becomes a metal — i.e., if the gap between the filled and empty state closes. Thus we cannot deform between different topological classes without closing the gap.

Indeed, this general picture gives us a simple rule for topological classification:

**Definition of Topological Phase:** Two gapped states of matter are in the same topological phase of matter if and only if you can continuously deform the Hamiltonian to get from one state to the other without closing the excitation gap.

Although in this chapter we are concerned with non-interacting electrons only, this sort of definition can obviously be used much more generally to distinguish different phases of matter. Further, this definition fits with our intuition about topology

*Two objects are topologically equivalent if and only if you can continuously deform one to the other.*

In the context of noninteracting electron band structure, one can define topologically “trivial” phases of matter to be those that can be continuously deformed without closing the gap into individual atomic sites with electrons that do not hop between sites. (A “trivial” band structure). Phases of matter that cannot be continuously deformed to this trivial band structure without closing a gap are known as topologically nontrivial.
36.1.1 Gapless Edges

The existence of gapless edge states on the edge of integer quantum Hall samples is one of the fundamental properties of topologically nontrivial phases of matter (at least when one is considering topological properties of noninteracting electron band structure). We can give a rough argument about why edge states always come with topologically nontrivial phases.

Suppose we have a Hamiltonian that is almost periodic, but the potential is a very function of position, say in the $x$-direction. In other words if we move very far in the $x$-direction the Hamiltonian changes smoothly from $H(x_1)$ to $H(x_2)$, but locally both of these look like simple periodic Hamiltonians. If $H(x_1)$ and $H(x_2)$ are not in the same topological phase of matter, than for some $x$ between $x_1$ and $x_2$, we have $H(x)$ describing some gapless system — i.e., an edge state between the two phases.

For example, in the case of the integer quantum Hall effect, we can think of $H(x_1)$ as being the Hamiltonian of the system in the bulk which has nonzero Chern number, and $H(x_2)$ as being the Hamiltonian outside of the system, or the vacuum, which is topologically trivial and has zero Chern number. Somewhere between the two, the gap must close to give a metal where the Chern number changes. This is the edge state.

36.2 Curvature and Chern Number

The Gauss-Bonnet theorem gives a beautiful connection exists between topology and geometry. The statement of the theorem is that for any closed two dimensional orientable surface the integral of the Gaussian curvature $K$ over the surface gives $2\pi(2-2g)$ where $g$ is the number of handles of the surface. Or mathematically

$$2\pi(2-2g) = \int_M K dS$$

One can check, for example, with a sphere of radius $R$ we have $K = 1/R^2$ and $g = 0$, so that both sides give $4\pi$ independent of $R$. The interesting point here is that if you dent the sphere, you increase the curvature at some points, but you decrease it at other points such that the integral of the curvature over the surface remains the same. The only way to change this quantity is to rip the surface and add a handle!

It turns out that we can define a similar curvature that describes the topological index (the Chern-number) of the band structure. Let us define what is known as the Berry curvature of the $\alpha$th band

$$\mathcal{F}_\alpha(k) = i^{\delta j} \langle \partial_k u^\alpha_k | \partial_k u^\alpha_k \rangle$$

The topological Chern-number of the $\alpha$th filled band is then given by

$^{3}$The definition of Gaussian curvature $K$ at a point is $1/K = \pm r_{max}r_{min}$ where $r_{max}$ and $r_{min}$ are the maximum and minimum radii of curvatures of the surface at that point. The sign of $K$ is taken to be negative if the surface is saddle-like at that point rather than dome-like.
Aside: A Rapid Introduction to Topological Insulators

The realization that the Hall conductance is the topological Chern number in 1982 was made in a famous paper known as TKNN. This is one of key contributions that earned a Nobel Prize for David Thouless in 2016.

the integral of the Berry curvature over the Brillouin zone,

$$C^\alpha = \frac{1}{2\pi} \int_{BZ} d\mathbf{k} F^\alpha(\mathbf{k})$$

which is analogously quantized to be an integer.

In appendix *** we use the Kubo formula to calculate the Hall conductivity and we find that it is related to the Chern number by\(^4\)

$$\sigma_{xy} = \frac{e^2}{h} \sum_{\text{filled bands } \alpha} C^\alpha$$

Considering Laughlin’s proof that the Hall conductance is quantized, this might be considered a sufficient proof that the Chern number must be quantized as well. To see how this occurs mathematically, see appendix ***.

36.3 Symmetry Protection

Symmetry is one of the most fundamental ideas in modern physics. We often think about how physics changes when a symmetry is forced on a system. Considering the above definition of topological phases of matter in section 36.1, one may generalize this idea to systems with symmetry.

Definition of Symmetry Protected Topological Phase: Two gapped states of matter are in the same symmetry protected topological phase of matter if and only if you can continuously deform the Hamiltonian to get from one state to the other without closing the excitation gap or breaking the given symmetry.

The most interesting example of this is time reversal symmetry. Systems without magnetism and without magnetic impurities are time-reversal symmetric. In three dimensions, it turns out that there are no band structures that satisfy the above definition of a nontrivial topological phase of matter. In other words, all gapped periodic single-electron Hamiltonians can be deformed to a trivial Hamiltonian without closing the gap. However, if we enforce time reversal invariance, it turns out that there are band structures that cannot be deformed into the trivial band structure without closing the gap or breaking symmetry. These are known as “topological insulators” and are formally symmetry protected topological phases, where the symmetry is time reversal.

36.4 Appendix: Chern Number is Hall Conductivity

Here we calculate the Hall conductivity by simple time dependent perturbation theory and demonstrate that it is the same as the Chern number.
The general rule of time dependent perturbation theory is that if a system is exposed to a perturbation $\delta H(t)$ the expectation of an operator $O$ at some later time is given by

$$\langle O(t) \rangle = \frac{i}{\hbar} \int_{-\infty}^{t} dt' \langle [O(t), H(t')] \rangle$$

If we consider an electric field at frequency $\omega$ we write this in terms of the vector potential. Applying a perturbing vector potential we have

$$\delta H = \int d\mathbf{x} A(\mathbf{x}, t) \cdot \mathbf{j}(\mathbf{x}, t)$$

From perturbation theory we then have

$$\langle j_a(x, t) \rangle = \frac{i}{\hbar} \int_{-\infty}^{t} dt' \int d\mathbf{x}' \langle [j_a(x, t), j_b(x', t')] \rangle A_b(x', t')$$
Having determined that the quantum Hall effect is some sort of spectroscopy on the charge of the electron, it was particularly surprising in 1982 when Dan Tsui and Horst Stormer\(^1\) discovered quantum Hall plateaus at fractional values of the filling fraction
\[
\nu = \frac{p}{q}
\]
with Hall resistance
\[
R_H = \frac{\hbar}{\mu} \frac{q}{p}
\]
with \(p\) and \(q\) small integers. This effect is appropriately called the Fractional quantum Hall effect.

The first plateau observed was the \(\nu = 1/3\) plateau\(^2\), but soon thereafter many more plateaus were discovered\(^3\). The Nobel Prize for this discovery was awarded in 1998.

Given our prior gauge invariance argument that quantum Hall effect is measuring the charge of the electron — and that this is enforced by the principle of gauge invariance, it is hard to understand how the fractional effect can get around our prior calculation.

Two things must be true in order to have quantized Hall effect

(a) Charge must fractionalize into quasiparticles with charge \(e^* = e/q\), for example in the case of \(\nu = 1/q\).

(b) The ground state on an annulus must be degenerate, with \(q\) different ground states (in the case of \(\nu = 1/q\)) which cycle into each other by flux insertion through the annulus.

We should not lose sight of the fact that these things are surprising — even though the idea of degenerate ground states, and possibly even fractionalized charges, is something we have perhaps gotten used to in our studies of topological systems.

Given the Laughlin argument that inserting a flux through the annulus pumps an integer number of electrons from one side to the other, it is perhaps not surprising that fractional quantization of the Hall conductance must imply that a fractional charge has been pumped from one side of the annulus to the other (hence point (a) above). The way we get around the gauge invariance argument that implies the charge must be an integer is by having multiple degenerate ground states. In our

\(^1\)Stormer had recently invented the idea of “modulation doping” semiconductors, which is a technique to obtain extremely clean two dimensional electron systems — a prerequisite for observing fractional quantum Hall effect.

\(^2\)The legend is that Tsui very presciently looked at the data the moment it was taken and said “quarks!” realizing that the fractional plateau implied charge fractionalization!

\(^3\)Over 60 different fractional quantum Hall plateaus have been discovered!
argument for the Integer quantum hall effect we used adiabaticity, and the existence of a gap, to argue that we must stay in the ground state. However when there are multiple ground states (point (b) above) we can only argue that we must always be in some ground state. Thus, for example, in the case of $\nu = 1/3$ where there are three ground states, the cycle of inserting flux is

$$\text{insert } \phi_0 \rightarrow |GS_1\rangle \rightarrow \text{insert } \phi_0 \rightarrow |GS_2\rangle \rightarrow \text{insert } \phi_0 \rightarrow |GS_3\rangle \rightarrow \text{insert } \phi_0 \rightarrow |GS_1\rangle$$

where $GS$ here means ground state. Each insertion of flux pumps $e^* = e/3$ charge from one side to the other. After three fractionally charged particles move from one side to the other, this amounts to a single electron being moved from one side to the other, and we return to exactly the same ground state as we started with.

So now we need only figure out how it is that this unusual situation of fractionalized charges, and multiple ground states (indeed, this situation of a topological quantum field theory!) comes about.

Want an incompressible state: Ignore disorder for now

We need to understand how we have an incompressible state when a Landau level is partially filled. As with the integer case, disorder will be important in allowing us to have plateaus of finite width, but the fundamental physics of the fractional quantum Hall effect comes from the fact that we have a gapped incompressible systems at a particular filling fraction. We can thus choose to consider a system free from disorder with the understanding that localization of excitations will be crucial to actually observe a plateau.

Why This is a Hard Problem: Massive Degeneracy

We restrict our attention to a clean system with a partially filled (say, 1/3 filled) Landau level. If there are $N_e$ electrons in the system, there $3N_e$ available single electron orbitals in which to place these electrons. Thus in the absence of disorder, and in the absence of interaction, there are

$$\binom{3N_e}{N_e} \sim (27/4)^{N_e}$$

multiparticle states to choose from — and all of these states have the same energy! In the thermodynamic limit this is an insanely enormous degeneracy\(^4\). This enormous degeneracy is broken by the interaction between the electrons, which will pick out a very small ground state manifold (in this case being just 3 degenerate ground states), and will leave the rest of this enormous Hilbert space with higher energy.

\(^4\)For example, if our system of size 1 square cm has a typically $10^{11}$ electrons in it, the number of degenerate states at $\nu = 1/3$ is roughly 10 to the 100 billion power! Way way way more than the number of atoms in the universe.
37.0.1 Our Model Hamiltonian

Since we are to neglect disorder, we can write the Hamiltonian for our system of interacting electrons as

\[ H = \sum_i \left( \frac{(p_i + eA(r_i))^2}{2m} \right) + \sum_{i<j} V(r_i - r_j) \]

where the first term is just the kinetic energy of the electrons in the magnetic field, as discussed in Section 35.5, and the second term is the interaction between the electrons, which we might take to be of \(1/r\) Coulomb form, or perhaps a modified Coulomb form depending on the physical situation we are concerned with\(^5\).

Now we have already analyzed the first term in this Hamiltonian back in Eq. 35.5 resulting in the structure of Landau levels. If we further assume that the cyclotron energy \(\hbar \omega_c\) (the energy gap between Landau levels) is very large compared to the interaction energy scale \(V\), then we can assume that there is very little effect of higher Landau levels — the interaction simply breaks the massive degeneracy of the partially filled Landau level without mixing in the higher Landau levels (or putting holes in any completely filled Landau levels below the chemical potential). Another way to say this is that we are pursuing degenerate perturbation theory. The kinetic energy is completely determined (we just fill up Landau levels from the bottom up) and interaction only plays a role to break the degeneracy of the partially filled level.

The effective Hamiltonian is then just

\[ H = \sum_{i<j} V(r_i - r_j) \quad (37.1) \]

where the Hilbert state is now restricted to a single partially filled Landau level. But here it might look like we are completely stuck. We have an enormously degenerate Hilbert space — and we have no small parameter for any sort of expansion.

Laughlin’s insight was to simply guess the correct wavefunction for the system\(^6\). In order to describe this wavefunction we need to have a bit more elementary information about wavefunctions in a magnetic field (some of this is a homework problem!).

37.1 Landau Level Wavefunctions in Symmetric Gauge

We will now work in the symmetric gauge where the vector potential is written as

\[ A = \frac{1}{2} \mathbf{r} \times \mathbf{B} \]

where the magnetic field is perpendicular to the plane of the sample. (We can check that this gives \(\nabla \times A = B\).
In this gauge, lowest Landau level wavefunctions (as mentioned before in section 35.8) take the form\(^7\)

\[
\varphi_m(z) = C_m z^m e^{-|z|^2/(4\ell^2)} \tag{37.2}
\]

where

\[
z = x + iy = re^{i\theta}
\]

is the complex representation of the particle coordinate, \(\ell = \sqrt{\hbar/eB}\) is the magnetic length, \(C_m\) is a normalization constant and here \(m \geq 0\) is an integer. The most general lowest Landau level wavefunction for a single particle would be \(f(z)\) times the gaussian factor for any analytic function \(f\).

Note that the higher Landau level wavefunctions can all be obtained by application of a raising operator (which involve some prefactors of \(z^*\)) to the lowest Landau level wavefunctions. This algebra is discussed in a homework problem, so we will not belabor it here. A key point is that all Landau levels are effectively equivalent and any partially filled higher Landau level is equivalent to a partially filled lowest Landau level with an appropriately modified interaction. As such, we will focus exclusively on the lowest Landau level from here on.

Let us take a close look at the structure of the wavefunctions in Eq. 37.2. First we note that \(\varphi_m\) is an eigenstate of the angular momentum operator \(\hat{L}\) (centered around the point \(z = 0\))

\[
\hat{L} \varphi_m = \hbar m \varphi_m
\]

Secondly we should examine the spatial structure of \(\varphi_m\). Writing \(|\varphi_m|^2 \sim r^{2m} \exp(-r^2/(2\ell^2))\) and differentiating with respect to \(r\) we find that the maximum of this function is at radius

\[
r = \ell \sqrt{2m}
\]

Thus the function roughly forms a gaussian ring at this radius. The area enclosed by this ring is \(\pi r^2 = 2\pi m\ell^2 = m\phi_0/B\), which contains precisely \(m\) quanta of magnetic flux.

### 37.1.1 What We Want in a Trial Wavefunction

In building a trial wavefunction for fractional quantum Hall effect, several rules will be important to follow

1. **Analytic Wavefunction:** The wavefunction in the lowest Landau level should be comprised of single particle wavefunctions \(\varphi_m\) — that is, it must be a polynomial in \(z\) (with no \(z^*\)’s) times the gaussian factors. In other words we should have\(^8\)

\[
\Psi(r_1, \ldots, r_N) = \text{(Polynomial in } z_1, \ldots, z_N) \prod_{i=1}^{N} e^{-|z_i|^2/(4\ell^2)}
\]

---

\(^7\)We will ignore \(\partial z^*\) as before.

\(^8\)The polynomial can also be chosen so as to have all real coefficients. This is because the Hamiltonian, once projected to a single Landau level, i.e., Eq. 36.1, is time reversal symmetric.
(2) **Homogeneous in Degree:** Since the Hamiltonian is rotationally invariant, we can expect that the eigenstates will be angular momentum eigenstates. Since the $\hat{L}$ operator counts powers of $z$, this means that the (Polynomial in $z_1, \ldots, z_N$) part of the wavefunction must be homogeneous of degree.

(3) **Maximum Power of $z_i$ is $N_\phi = N/\nu$:** Since the radius of the wavefunction is set by the exponent of $z^m$, the full radius of the quantum Hall droplet is given by the largest power of any $z$ that occurs in the wavefunction. Since the area enclosed by the wavefunction should correspond to $N_\phi$ fluxes, this should be the maximum power.

(4) **Symmetry:** The wavefunction should be fully antisymmetric due to Fermi statistics, assuming we are considering fractional quantum Hall effect of electrons. It is actually very useful theoretically (and does not seem out of the question experimentally\(^9\)) to consider fractional quantum Hall effect of bosons as well — in which case the wavefunction should be fully symmetric.

Even given these conditions we still have an enormous freedom in what wavefunction we might write down. In principle this wavefunction should depend on the particular interaction $V(r)$ that we put in our Hamiltonian. The miracle here is that, in fact, the details of the interaction often do not matter that much!

### 37.2 Laughlin’s Ansatz

Laughlin simply guessed that a good wavefunction would be of the form\(^{10}\)

$$\Psi^{(m)}_{\text{Laughlin}} = \prod_{i<j} (z_i - z_j)^m \prod_{i=1}^N e^{-|z_i|^2/(4\ell^2)}$$

The proposed wavefunction is properly analytic and homogeneous in degree. The maximum power of the wavefunction is

$$N_\phi = m(N - 1)$$

thus corresponding to a filling fraction

$$\nu = N/N_\phi \to 1/m \quad \text{in large } N \text{ limit}$$

And the wavefunction is properly antisymmetric for $m$ odd, and is symmetric for $m$ even.

It is worth noting that for $m = 1$ the Laughlin wavefunction corresponds to a filled Landau level — that is, a single slater determinant filling all of the orbitals from $m = 0$ to $m = N_\phi = N - 1$. (This is a homework problem!)

\(^9\)While no one has yet produced fractional quantum Hall effect of bosons in the laboratory, proposals for how to do this with cold atoms or interacting photons are plentiful, and it seems very likely that this will be achieved in the next few years.

\(^{10}\)Note that this wavefunction is not normalized in any sense. The issue of normalization becomes important later in ***.
It is also worth noting that the density of the Laughlin wavefunction is completely constant in a disk up to its radius (and then the density falls quickly to zero). This constancy of density is proven by plasma analogy (which is another homework problem\textsuperscript{11}).

Why should we think this wavefunction is particularly good? As two particles approach each other, the wavefunction vanishes as $m$ powers. This means that the particles have low probability of coming close to each other — thus keeping the interaction energy low.

Being that the polynomial in each variable is of fixed degree $N_\phi$, the polynomial has a fixed number of analytic zeros. For the Laughlin wavefunction all of these zeros are on the positions of the other particles — thus the wavefunction arranges that the particles stay as far away from each other as possible in some sense.

### 37.2.1 Exact statements about Laughlin Wavefunction

It turns out that the Laughlin wavefunction is actually the exact ground state of a special inter-particle interaction\textsuperscript{12}.

**Bosons at $\nu = 1/2$**

Consider a system of bosons with the inter-particle interaction given by\textsuperscript{13}

$$V = V_0 \sum_{i<j} \delta(\mathbf{r}_i - \mathbf{r}_j)$$

with $V_0 > 0$. This is a non-negative definite interaction.

It is clear that the $\nu = 1/2$ Laughlin state of bosons $\Psi_{\text{Laughlin}}^{(m=2)}$ has zero energy for this interaction, since there is zero amplitude of any two particles coming to the same point. Further, however, the Laughlin state is the highest density wavefunction (lowest degree polynomial) that has this property\textsuperscript{14}. For example, the Laughlin state times any polynomial is also a zero energy state of this interaction, but since it has been multiplied by a polynomial, the total degree of the wavefunction is higher, meaning the wavefunction extends to higher radius, making the system

\textsuperscript{11}Roughly the story is as follows. The probability $|\Psi(z_1, \ldots, z_N)|$ of finding particles at position $z_1, \ldots, z_N$ can be phrased as a classical stat mech problem of a one-component 2d coulomb plasma in a background charge, by writing

$$|\Psi|^2 = e^{-\beta U(z_1, \ldots, z_N)}$$

with $\beta = 2/m$ and

$$U = -m^2 \sum_{i<j} \log(|z_i - z_j|) + \frac{m}{4} \sum_i |z_i|^2$$

where the first term is the coulomb interaction in 2d, and the second term is a background charge — which happens to be the charge associated with a uniform positive background (an easy thing to check using gauss’s law). Assuming this plasma screens the background charge, it will be of uniform density up to a constant radius.

\textsuperscript{12}This was discovered by Haldane in 1983, then again by Trugman and Kivelson and also Pokrovski and Tsalapov in 1985.

\textsuperscript{13}Actually this is a very realistic interaction for cold atom bosonic quantum Hall effect, should it be produced in the future.

\textsuperscript{14}Although with some thought this fact seems obvious, proving it rigorously is tricky.
less dense. A schematic of the ground state energy as a function of filling fraction for this case is shown in Fig. 37.1.

\[ \varepsilon \]

\[ \nu = \frac{N}{N_f} \]

**Fig. 37.1** Schematic of the ground state energy as a function of filling fraction for bosons with delta function interaction.

The key point is that the ground state energy has a cusp, which means there is a jump in the chemical potential

\[ \mu = \frac{\partial E}{\partial N} \]

This is precisely the same “incompressibility” as we have in the case of noninteracting electrons — where the chemical potential jumps between Landau levels! As in that case we presume that the presence of a cusp in the free energy, in the absence of disorder, will be enough to give us a plateau when disorder is added back in.

Now while we can easily show that there is a change of behavior at \( \nu = 1/2 \) in this plot, it is somewhat more difficult to be convincing that the slope coming from the right is finite — i.e., that the gap is actually finite. In order to do that, we would need to think about the elementary excitations, or resort to numerics.

**Fermions at \( \nu = 1/3 \)**

The arguments given for bosons at \( \nu = 1/2 \) can be easily generalized to the case of fermions (i.e., electrons) at \( \nu = 1/3 \) (and more generally to any \( \nu = 1/m \).) Obviously a \( \delta \)-function interaction will no longer do the job, since for fermions Pauli exclusion prevents any two fermions from coming to the same point already. However, consider an interaction of the form

\[ V = V_0 \sum_{i<j} \nabla^2 \delta(r_i - r_j) \]
Given a wavefunction $\Psi(r_1, \ldots, r_N)$ the interaction energy will be

$$E = \sum_{i<j} \int dr_1 \ldots dr_N |\Psi|^2 \nabla^2 \delta(r_i - r_j)$$

Writing

$$\Psi(dr_1 \ldots dr_N) = \phi(z_1 \ldots z_N) \prod_{i=1}^N e^{-|z_i|^2/(4\ell^2)}$$

with $\phi$ meaning the analytic polynomial part, for fermionic wavefunctions (that must vanish when $r_i = r_j$) the expression for the energy can be integrated by parts using $\nabla^2 = 4\partial_z \partial_{\bar{z}}$ to give

$$E = \sum_{i<j} \int dr_1 \ldots dr_N |\partial_z \phi|^2 \delta(r_i - r_j) \prod_{i=1}^N e^{-|z_i|^2/(2\ell^2)}$$

Thus we have a non-negative definite interaction. Further, if the wavefunction vanishes as a single power when two particles come together, then $\partial_z \phi$ will be nonzero and we will get a positive result (since $\partial_z(z_i - z_j)$ is nonzero). However, if the wavefunction vanishes as three powers $\partial_z \phi$ will remain zero (since $\partial_z(z_i - z_j)^3$ goes to zero when $z_i = z_j$).

Thus, entirely analogous to the above case of $\nu = 1/2$ with the $\delta$-function interaction, the Laughlin $m = 3$ ($\nu = 1/3$) wavefunction is the exact ground state (unique highest density zero energy wavefunction) of the $\nabla^2 \delta$-function interaction. With similar ideas, one can construct interactions for which any $\nu = 1/m$ Laughlin wavefunction is exact.

### 37.2.2 Real Interactions

Obviously electrons do not interact via a $\nabla^2 \delta$ interaction. They interact via a Coulomb interaction. What is perhaps surprising is that the Laughlin wavefunction is an almost perfect representation of the actual ground state. This statement comes from numerical tests. For example, for 9 electrons (on a spherical geometry to remove edge effects) the dimension of the fully symmetry reduced Hilbert space is 84, and yet the Laughlin trial wavefunction has an overlap squared of .988 with the exact ground state of the Coulomb interaction. This is absurdly accurate! The energy of the Laughlin wavefunction differs from the energy of the exact Coulomb ground state by less than a part in two thousand.

### 37.3 Quasiparticles

The Laughlin quantum hall ground state is a uniform density fluid (we will actually show this as a homework problem). Density perturbations are made in discrete units of charge known as quasiparticles. Positively

---

15 Generally one would expect derivatives of the gaussian part as well when we integrate by parts. However, because the polynomial is antisymmetric, the derivative must act on the polynomial part to prevent the wavefunction from vanishing when particle coordinates coincide.

16 Note that by antisymmetry the wavefunction must vanish as an odd number of powers as two particle positions approach each other.

17 In higher Landau levels, although the interaction is Coulomb, when the single Landau level problem is mapped to a single partly filled lowest Landau level (See the comments after Eq. 37.2), the interaction gets modified – this mainly effects the short range behavior.

18 The full Hilbert space is 45207 dimensional!

19 I need to recheck this number.
charged bumps of charge (opposite the charge of the electron) are known as quasiholes and negatively charged bumps of charge (same charge of the electron) are quasielectrons.

### 37.3.1 Quasiholes

For the quasiholes, it is fairly easy to guess their wavefunction (and indeed this was done by Laughlin). We start by considering adding a quasihole at position \(0\). This leaves the system rotationally invariant. We guess the solution

\[
\Psi_{qh}(0) = \left[ \prod_{i=1}^{N} z_i \right] \Psi_{Laughlin}
\]

where \(0\) indicates we have put the quasihole at position \(0\). Here the degree of the polynomial is increased by one for every variable, so each filled orbital gets pushed out to the next orbital. This leaves precisely one empty orbital open at position \(0\). Since our wavefunction has filling fraction \(\nu\), this means that on average a fraction \(\nu\) of the orbitals are filled. Thus leaving the orbital at the center completely empty corresponds to a positive charge of \(+\nu\), and our quasihole has a positive charge

\[
e^* = \nu e.
\]

Another way to think about the same wavefunction is to imagine adiabatically inserting a quantum of flux \(\phi_0\) at position \(0\). Analogous to the Laughlin argument for integer quantum Hall effect, this creates an azimuthal EMF. Since the system has quantized Hall conductance \(\sigma_{xy} = \nu e^2/h\), the total charge created is \(\nu e = \sigma_{xy} \phi_0\). Then once we have inserted the flux, the flux quantum can be gauged away leaving only the quasihole behind.

One can make quasiholes at any location \(w\) analogously,

\[
\Psi_{qh}(w) = \left[ \prod_{i=1}^{N} (z_i - w) \right] \Psi_{Laughlin}
\]

although this is no longer an angular momentum eigenstate. We can similarly consider multiple quasiholes the same way

\[
\Psi_{qhs}(w_1, \ldots, w_M) = \left[ \prod_{\alpha=1}^{M} \prod_{i=1}^{N} (z_i - w_\alpha) \right] \Psi_{Laughlin}
\]

Several interesting comments at this point:

1. While the \(z\)'s are physical electron coordinates, the \(w\) parameters are simply parameters of the wavefunction and can be chosen and fixed to any value we like. The wavefunction \(\Psi(w_1, \ldots, w_M; z_1, \ldots, z_N)\) is then the wavefunction of electrons \(z\) in the presence of quasiholes at fixed \(w\)
positions.

(2) Note that the phase of the wavefunction wraps by $2\pi$ when any electron moves around the position of a quasihole.

(3) For the special ultra-short-range wavefunctions for which the Laughlin ground state is an exact zero energy eigenstate, then this Laughlin quasihole is also an exact zero energy eigenstate (albeit one with lower density than the ground state since a hole has been inserted). Take for example the case of $\nu = 1/2$. With a $\delta$-function interaction, the energy is zero because no two particles come to the same point. Multiplying this wavefunction by any polynomial (as we have done to insert quasiholes) maintains this property and we still have a zero energy eigenstate. As is the case for the Laughlin ground state, the quasihole is not exact for the Coulomb interaction, but is extremely accurate numerically.

(4) At $\nu = 1/m$, if we insert $m$ quasiholes at the same point $w$, then the wavefunction is just the same as if we were to have an electron $e$ at the point $w$ (although the electron is not there). Thus we expect that “fusing” $m$ quasiholes together should precisely make an anti-electron (or a real hole).

### 37.3.2 Quasielectrons

The quasi-electron is a bump of negative charge (i.e., same charge as the electron). Unlike the case of quasiholes, there are no exact wavefunctions that we know of for quasi-electrons (not even for special short range interactions).

Whereas the quasi-hole increases the total degree of the polynomial wavefunction (thereby decreasing the density of the system) the quasi-electron should decrease the total degree of the wavefunction. Again, Laughlin made a very good guess of what the wavefunction for the quasi-electron should be. Considering a quasi-electron at the origin, we can write

$$\Psi_{qe}(0) = \left(\prod_{i=1}^{N} \frac{\partial}{\partial z_i}\right) \phi \prod_{i=1}^{N} e^{-|z_i|^2/(4\ell^2)}$$

where as in Eq. 37.3 we have written the Laughlin wavefunction as the polynomial part $\phi$ times the gaussian factors. Obviously the derivative correctly reduces the degree of the polynomial by one in each variable $z$, thus reducing the net angular momentum of each particle by one. Each particle moves to lower radius by one orbital, thus giving a pile-up of charge of $e^* = -e\nu$ at the origin.

In analogy to (but opposite that of) the quasihole, we might have looked for a quasi-electron where electrons accumulate a phase of $-2\pi$ when an electron moves around the quasiparticle. One might think of the operator $z^*$, but this operator does not live in the lowest Landau level. However, the projection of this operator to the lowest Landau level is given by

$$P_{LLL}z^* = 2\ell^2 \frac{\partial}{\partial z}$$
As mentioned above, the Laughlin quasi-electron is not exact for any known system. However, it is a fairly good trial wavefunction numerically for the Coulomb interaction. Note however, that other forms for the quasi-electron wavefunction have been found to be somewhat more accurate.

One can move the quasielectron to any position in a similar way as for quasiholes giving a wavefunction of the form

$$\Psi_{\text{qes}}(w) = \left( \prod_{i=1}^{N} \left( 2\ell^2 \frac{\partial}{\partial z_i} - w^* \right) \phi \right) \prod_{i=1}^{N} e^{-|z_i|^2/(4\ell^2)}$$

### 37.3.3 Fractional Charge and Statistics?

The quasiparticles of the Laughlin state thus have fractional charge. One should not lose sight of how surprising this is — that particles can emerge that are a fraction of the “elementary” particles of the system. If we lived at very low energy, we would experience these quasiparticles as the fundamental particles of the system and would not know of the existence of the underlying electron.

Once one accepts fractionalized charge, it is perhaps not surprising to discover that they also have fractional statistics. Proving this statement is nontrivial, and we will do it in several ways. Note that since the quasiparticles are charged, moving them around in a magnetic field incurs phases. We would like thus like to compare the phase of moving a particle in a loop versus moving a particle in a loop when another particle might be inside the loop, see fig. 37.2.

![Fig. 37.2](image.png)  

**Fig. 37.2** To find the statistical phase, we compare moving a particle in a loop versus moving it in the same loop when another particle is inside the loop.

We shall perform this comparison next after we introduce Berry’s phase, which is the effect which produces the statistical phase we are interested in.

### 37.4 Digression on Berry’s Phase

The Berry phase\(^{20}\) is one of the most fundamental ideas of modern physics. We recall the adiabatic theorem. If you start in an eigenstate and change a Hamiltonian sufficiently slowly, and there are no level crossings, then the system will just track the eigenstate as it slowly

---

\(^{20}\)Berry’s work on Berry Phase in 1984 had a number of precursors, most notably the work of Pancharatnam in 1956.
changes — i.e., it remains in the instantaneous eigenstate. However, during this process it takes a bit of thought to figure out what happens to the phase of the wavefunction.

To see how this correction arises, let us consider a Hamiltonian $H(R)$ which is a function of some general parameters which we will summarize as the vector $R$. In our case these parameters are going to represent the quasiparticle position — we will insert this information into the Hamiltonian by having some trapping potential which induces the quasiparticle at the point $R$ and we can then move around the trapping potential in order to move the particle. Let us write the instantaneous (here normalized!) eigenstate as $|\psi(R)\rangle$. So we have

$$H(R)|\psi(R)\rangle = E(R)|\psi(R)\rangle$$

Now let us write the full, time dependent wavefunction as

$$|\Psi(t)\rangle = e^{i\gamma(t)}|\psi(R(t))\rangle$$

so we are allowing for an additional phase out front of the instantaneous eigenstate. The time dependent Schroedinger equation is

$$i\hbar \frac{\partial}{\partial t}|\Psi(t)\rangle = H(R(t))|\Psi(t)\rangle$$

$$[-\hbar \dot{\gamma} + i\hbar \frac{\partial}{\partial t}]|\psi(R(t))\rangle = E(R(t))|\psi(R(t))\rangle$$

Projecting this equation onto the bra $\langle \psi(R) |$ we obtain

$$\dot{\gamma} = -E(R(t))/\hbar - i \left\langle \psi(R(t)) \left| \frac{\partial}{\partial t} \right| \psi(R(t)) \right\rangle$$

Integrating over some path $R(t)$ from some initial time $t_i$ to some final time $t_f$ gives

$$\gamma(t_f) - \gamma(t_i) = -\frac{1}{\hbar} \int_{t_i}^{t_f} E(R(t)) dt - i \int_{t_i}^{t_f} dR \cdot \langle \psi(R) | \nabla_R | \psi(R) \rangle$$

The first term is the expected dynamical phase — just accumulating a phase with time proportional to the energy. The second term on the right is the Berry phase contribution — a line integral along the particular path that $R(t)$ takes. Note that this term depends only on the geometry of the path and not on how long one takes to move through this path. In this sense it is a geometric phase.

### 37.5 Arovas-Schrieffer-Wilczek Calculation of Fractional Statistics

This section follows the approach of Arovas, Schrieffer and Wilczek\(^{21}\).
Let us consider a $\nu = 1/m$ wavefunction for a quasihole

$$\Psi(w) = N(|w|) \prod_{i=1}^{N} (z_i - w) \Psi_{\text{Laughlin}}^{(m)}$$

and we will imagine moving around the position $w$ in a circle of constant radius as shown in the right of Fig. 37.2. Here we have inserted a normalization constant out front, which can be shown to be a function of radius only. (This is argued by plasma analogy, which is part of the homework). We will then parameterize $w = |w|e^{i\theta}$.

The Berry phase from moving the particle in a loop will then be

$$\Delta \gamma = -i \int_{0}^{2\pi} d\theta \langle \Psi(\theta) | \partial_{\theta} | \Psi(\theta) \rangle$$

where we have written $|\Psi(\theta)\rangle$ to mean $|\Psi(|w|e^{i\theta})\rangle$. We then have

$$\partial_{\theta} |\Psi(\theta)\rangle = \frac{\partial w}{\partial \theta} \left( \sum_{i} \frac{-1}{z_i - w} \right) |\Psi(\theta)\rangle$$

Thus we have

$$\langle \Psi(\theta) | \partial_{\theta} | \Psi(\theta) \rangle = \frac{\partial w}{\partial \theta} \sum_{i} \left\langle \Psi(\theta) \left| \frac{-1}{z_i - w} \Psi(\theta) \right\rangle \right.$$ 

Thus from taking $w$ around in a circle we obtain the Berry phase

$$\Delta \gamma = -i \oint d\theta \langle \Psi(\theta) | \partial_{\theta} | \Psi(\theta) \rangle$$

$$= -i \oint dw \sum_{i} \left\langle \Psi(w) \left| \frac{-1}{z_i - w} \Psi(w) \right\rangle \right.$$ 

Now the integral around the loop of $1/(z - w)$ accumulates $2\pi i$ if and only if $z_i$ is inside the loop. Thus we obtain the phase

$$\Delta \gamma = 2\pi \langle \text{number of electrons in loop} \rangle$$

$$= 2\pi (1/m) \Phi/\Phi_0 = \gamma_{AB}$$

where $\Phi$ is the flux enclosed by the loop and $\Phi_0$ is the flux quantum (and here we have used $\nu = 1/m$). This is precisely the expected Aharonov-Bohm phase that we should expect for moving a charge $e/m$ around a flux $\Phi$.

Now we consider putting another quasiparticle in the center of the loop as shown in the left of Fig. 37.2 Using a normalization factor that is again a function of $|w|$ only, the same calculation holds, but now the

---

22 On can choose a more general path for the particle but we will then need the detailed form of $N(w)$. See the discussion below in section ***

23 The way this is written it is obviously a bit nonsense. Please fix it. I wrote this footnote, but now I don’t see what is wrong with what I have here! ***
number of electrons enclosed has changed by one quasiparticle charge $e/m$. Thus the phase is now

$$\Delta \gamma = \gamma_{AB} + \gamma_{\text{statistical}}$$

where the additional phase for having gone around another quasihole is given by

$$\gamma_{\text{statistical}} = \frac{2\pi}{m}$$

or in other words we have fractional statistics! For example, for the Laughlin state at $\nu = 1/2$, we have semionic statistics.

A more detailed version of this calculation (we will do this below) shows that the path of the particle does not matter --- the total phase is always the Aharonov-Bohm phase for taking a particle around flux, added to the statical phase of taking it around another quasiparticle.

Comment on the Fusion/Braiding Rules, and Chern-Simons theory

For the $\nu = 1/m$ Laughlin state thus we have a situation where the elementary quasi-holes have statistics $\theta = 2\pi/m$. We can assume that their antiparticles will have the same statistics (both opposite “charge” and “flux” in a charge-flux model). We also have that the fusion of $m$ elementary quasi-electrons or quasi-holes forms an electron or anti-electron.

In the case where $m$ is even, the underlying “electron” is a boson, in which case we can think of this electron as being identical to the vacuum — it has trivial braiding with all particles and it is essentially condensed into the ground state as some sort of background superfluid. Thus we have a simple anyon theory with $m$ particle types.

On the other hand, when $m$ is odd, we have the situation (discussed in our “charge-flux composite” section *** where the fusion of $m$ elementary anyons forms a fermion — and so there are actually $2m$ particle types — the fermion full-braids trivially with everything, but has fermionic statistics with itself. This situation is “non-modular” — it does not have as many ground states as it has particle types. There are only $m$ ground states, despite $2m$ particle types.

37.6 Gauge Choice and Monodromy

The Laughlin wavefunction with $M$ quasiholes takes the form

$$\Psi(w_1, \ldots, w_M; z_1, \ldots, z_N) =$$

$$\mathcal{N}(w_1, \ldots, w_N) \prod_{\alpha=1}^{M} \prod_{i=1}^{N} (z_i - w_{\alpha}) \psi^{(m)}_{\text{Laughlin}}(z_1, \ldots, z_N)$$

where $\mathcal{N}$ is a normalizing factor.

By using a plasma analogy (this is a homework assignment) we find
that the normalization must be of the form

$$|\mathcal{N}(w_1, \ldots, w_M)| = C \prod_{\alpha < \beta} |w_\alpha - w_\beta|^{1/m} \prod_{\alpha=1}^M e^{-|w_\alpha|^2/(4\ell^*^2)}$$

where $C$ is some constant and

$$\ell^* = \sqrt{\frac{\hbar}{e^* B}}$$

is the effective magnetic length for a particle of charge $e^* = e/m$. This choice of normalization assures that

$$\langle \Psi(w_1, \ldots, w_M) | \Psi(w_1, \ldots, w_M) \rangle$$

is independent of the position of the quasiholes.

Now, we can choose the phase of the factor $\mathcal{N}$ arbitrarily — this is essentially a gauge choice. In the above Arovas, Schrieffer, Wilczek calculation above, we chose the phase to be real. However, this is just a convention. An interesting different convention is to choose

$$\mathcal{N}(w_1, \ldots, w_N) = C \prod_{\alpha < \beta} (w_\alpha - w_\beta)^{1/m} \prod_{\alpha=1}^M e^{-|w_\alpha|^2/(4\ell^*^2)}$$

which is known as holomorphic or “fractional statistics” gauge — here the fractional statistics of the quasiparticles are put explicitly into the wavefunction! Note here that this function is not single valued in the $w$-coordinates. In this gauge, we see that the wavefunction has branch cuts and can be thought of as having Riemann sheets. This may look problematic, but it is not. While a wavefunction must be single-valued in the physical electron coordinates, the $w$’s are just parameters of the wavefunction, and we are allowed to choose wavefunctions’ phase conventions in any way we like — even in non-single-valued ways as we have done here.

What we would want to confirm is that the physical phase accumulated in moving one quasihole around another is independent of our gauge choice. To this end we note that the total phase accumulated can be decomposed into two pieces, the so-called monodromy and the Berry phase. The monodromy is the phase explicitly accumulated by the wavefunction when one coordinate is moved around another.

$$\text{Total Phase} = \text{Monodromy} + \text{Berry Phase}$$

In the above Arovas-Schrieffer-Wilczek calculation, we chose the phase of the normalization to be everywhere real. So there is no monodromy — no explicit phase as we move one particle around another. However, in fractional statistics gauge we see a phase of $2\pi/m$ for each particle which travels counterclockwise around another. In both gauges the total phase should be the same, so in the holomorphic gauge, the statistical
part of the phase should be absent. Let us see how this happens.

37.6.1 Fractional Statistics Calculation: Redux

Let us consider the case of two quasi-holes and repeat the argument of Arovas-Schrieffer-Wilczek but in holomorphic gauge. Putting one quasihole at position \( w \) and another at position \( w' \) the wavefunction is

\[
\Psi(w) = C \left( \frac{1}{m} e^{-|w|^2 + |w'|^2 / (4\ell^2)} \times \prod_i (z_i - w) \prod_i (z_i - w') \prod_{i<j} e^{-|z_i|^2 / (4\ell^2)} \right)
\]

with \( C \) chosen so that \( \Psi \) is normalized independent of the quasihole coordinates.

Let us parameterize the path of a quasiparticle as \( w(\tau) \).

\[\Delta \gamma = -i \oint d\tau \langle \Psi(\tau) | \partial \tau | \Psi(\tau) \rangle \]

We write

\[
\frac{\partial}{\partial \tau} = \frac{\partial w}{\partial \tau} \frac{\partial}{\partial w} + \frac{\partial w^*}{\partial \tau} \frac{\partial}{\partial w^*}
\] (37.6)

Now, because we are using holomorphic gauge of the wavefunction the \( \partial / \partial w^* \) only hits the gaussian factor, so we have

\[
\langle \Psi(w) | \partial_{w^*} | \Psi(w) \rangle = -\frac{w}{4\ell^2} \langle \Psi(w) | \Psi(w) \rangle = -\frac{w}{4\ell^2}
\]

To evaluate the derivative \( \partial / \partial w \) we integrate by parts so that it acts on the bra rather than the ket. Now since the bra is completely anti-holomorphic in \( w \) except the gaussian, the derivative acts only on the gaussian again to give

\[
\langle \Psi(w) | \partial_w | \Psi(w) \rangle = \partial_w \left[ \langle \Psi(w) | \Psi(w) \rangle \right] - [\partial_w \langle \Psi(w) \rangle] \langle \Psi(w) \rangle = \frac{w^*}{4\ell^2} \langle \Psi(w) | \Psi(w) \rangle = \frac{w^*}{4\ell^2}
\]

Note that the derivative on \( \langle \Psi | \Psi \rangle \) here is zero because the wavefunction is assumed normalized to unity for every value of \( w \).

We then have the Berry phase given by

\[
\Delta \gamma = -i \oint d\tau \langle \Psi(\tau) | \partial \tau | \Psi(\tau) \rangle = -i \frac{1}{4\ell^2} \oint (dw w^* - dw^* w)
\]

where we have used Eq. 37.6. We now use the complex version of Stokes theorem\(^{25}\) to obtain

\[
\Delta \gamma = \frac{\text{Area}}{\ell^2} = 2\pi (1/m) \Phi / \Phi_0
\]

which is the Aharonov-Bohm phase corresponding to the flux enclosed in the path – without giving the fractional statistical phase which has now been moved to the monodromy!
The key point here, which we emphasize, is that if we work with normalized holomorphic wavefunctions (i.e., holomorphic gauge), then the fractional statistics are fully explicit in the monodromy of the wavefunction — we can read the statistics off from the wavefunction without doing any work!

37.7 Appendix: Building an Effective (Chern-Simons) Field Theory

We can consider writing an effective field theory for this \( \nu = 1/m \) quantum Hall system. First let us think about how it responds to an externally applied electromagnetic field. It should have its density locked to the magnetic field, so we should have a change of electron density (In this section we set \( \hbar = e = 1 \) for simplicity)

\[
\delta n = j^0 = \frac{1}{2\pi m} \delta B
\]

Similarly we should expect a quantized Hall conductance, here with \( j \) being the current of electrons

\[
j^i = -\frac{1}{2\pi m} \epsilon^{ij} E_j
\]

Both of these can be summarized as the response to a perturbing vector potential

\[
j^\mu = -\frac{1}{2\pi m} \epsilon^{\mu\nu\lambda} \partial_\nu \delta A_\lambda
\]  

(37.7)

We must, of course have charge conservation as well. This is easy to enforce by writing the current in the form

\[
j^\mu = \frac{1}{2\pi} \epsilon^{\mu\nu\lambda} \partial_\nu a_\lambda
\]  

(37.8)

which then automatically satisfies

\[
\partial_\mu j^\mu = 0
\]

In this language, the effective Lagrangian that produces Eq. 37.7 as an equation of motion is then

\[
\mathcal{L} = -\frac{m}{4\pi} \epsilon^{\mu\nu\lambda} a_\mu \partial_\nu a_\lambda + \frac{1}{2\pi} \epsilon^{\mu\nu\lambda} A_\mu \partial_\nu a_\lambda + j_\eta^\mu a_\mu
\]

where \( j_\eta \) is the quasiparticle current. Note that without the \( A_\mu \) term, this is the same Chern-Simons theory we used for describing fractional statistics particles (now the quasiparticles).

To see the coupling to the external vector potential, note that the general (Noether) current associated with the local gauge symmetry will be

\[
j^\mu = \frac{\partial \mathcal{L}}{\partial A^\mu}
\]
which matches the expression from Eq. 37.8. By differentiating the Lagrangian with respect to $a_\mu$, we generate the equations of motion Eq. 37.7.

More here

37.8 Appendix: Quantum Hall Hierarchy

Good reference is https://arxiv.org/abs/1601.01697

Shortly after the discovery of the Laughlin $\nu = 1/3$ state additional fractional quantum Hall plateaus were discovered at filling fractions such as $\nu = 2/3, 2/5, 3/7$ and so forth. By now over 60 different plateaus have been observed in experiment!

The Laughlin theory only describes filling fractions $\nu = 1/m$ but it contains in it the right ideas to build possible theories for many of these fractions.

There are several approaches to building a hierarchy of quantum Hall states, however perhaps the most intuition comes from the original approaches by Haldane and Halperin in 1983.

The general idea is to begin with a Laughlin wavefunction for $N$ electrons with coordinates $z_i$ for $\nu = 1/m$ then change the magnetic field to add a large number $M$ of quasiparticles (say in the form of 37.4, in the case of quasiholes) at coordinates $w_\alpha$. Thus our wavefunction we write as $\Psi(w_1, \ldots w_M; z_1, \ldots z_N)$ as written in Eq. 37.4. We then write a pseudowavefunction to describe some dynamics of the quasiholes which we write as

$$\phi(w_1, \ldots, w_M)$$

An electron wavefunction is generated by integrating out the quasihole coordinates. Thus we have

$$\Psi(z_1, \ldots, z_N) = \int dw_1, \ldots dw_M \phi^*(w_1, \ldots, w_M) \Psi(w_1, \ldots w_M; z_1, \ldots z_N)$$

The general idea of this scheme is that the pseudo-wavefunction can itself be of the form of a Laughlin wavefunction. In the original Laughlin argument we wrote down wavefunctions for both boson and fermion particles. Here, the particles $w$ are anyons, so we need to write a slightly different form of a wavefunction. We expect

$$\phi(w_1, \ldots, w_M) = \prod_{\alpha<\beta} (w_\alpha - w_\beta)^{1/m + p}$$

with $p$ an even integer. The fractional power accounts for the fact that the anyon wavefunction must be multi-valued as one particle moves around another. The factor $p$ is to include a “Laughlin” factor repelling these anyons from each other without further changing the statistics.

The condensation of these quasi-particles into a Laughlin state gener-
ates a wavefunction for the filling fraction

\[ \nu = \frac{1}{m \pm 1/p} \]

with the ± corresponding to whether we are condensing quasiparticles or quasiholes. One can continue the argument starting with these new fractions and generating further daughter states and so forth. At the next level for example, we have

\[ \nu = \frac{1}{m \pm \frac{1}{p \pm \frac{1}{q}}} \]

By repeating the procedure, any odd denominator fraction \( \nu = p/q \) can be obtained.
Exercises

**Exercise 37.1 Filled Lowest Landau Level**

Show that the filled Lowest Landau level of non-interacting electrons (a single Slater determinant) can be written as

\[ \Psi^0_0 = N \prod_{1 \leq i < j \leq N} (z_i - z_j)^1 \prod_{1 \leq i \leq N} e^{-|z_i|^2/4\ell^2} \]  

(37.9)

with \( N \) some normalization constant. I.e., this is the Laughlin wavefunction with exponent \( m = 1 \).

**Exercise 37.2 Laughlin Plasma Analogy**

Consider the Laughlin wavefunction for \( N \) electrons at positions \( z_i \)

\[ \Psi^0_m = N \prod_{1 \leq i < j \leq N} (z_i - z_j)^m \prod_{1 \leq i \leq N} e^{-|z_i|^2/4\ell^2} \]  

(37.10)

with \( N \) a normalization constant. The probability of finding particles at positions \( \{z_1, \ldots, z_N\} \) is given by \( |\Psi_m(z_1, \ldots, z_N)|^2 \).

Consider now \( N \) classical particles at temperature \( \beta = \frac{1}{k_B T} \) in a plane interacting with logarithmic interactions \( v(\vec{r}_i - \vec{r}_j) \) such that

\[ \beta v(\vec{r}_i - \vec{r}_j) = -2m \log(|\vec{r}_i - \vec{r}_j|) \]  

(37.11)

in the presence of a background potential \( u \) such that

\[ \beta u(|\vec{r}|) = |\vec{r}|^2/(2\ell^2) \]  

(37.12)

Note that this log interaction is “Coulombic” in 2d (i.e., \( \nabla^2 v(\vec{r}) \propto \delta(\vec{r}) \)).

(a) Show that the probability that these classical particles will take positions \( \{\vec{r}_1, \ldots, \vec{r}_N\} \) is given by \( |\Psi_m(z_1, \ldots, z_N)|^2 \) where \( z_j = x_j + iy_j \) is the complex representation of position \( \vec{r}_j \). Argue that the mean particle density is constant up to a radius of roughly \( 2\sqrt{Nm} \). (Hint: Note that \( u \) is a neutralizing background. What configuration of charge would fully screen this background?)

(b) Now consider the same Laughlin wavefunction, but now with \( M \) quasiholes inserted at positions \( w_1, \ldots, w_M \).

\[ \Psi_m = N(w_1, \ldots, w_M) \left[ \prod_{1 \leq i \leq N} \prod_{1 \leq \alpha \leq M} (z_i - w_\alpha) \right] \Psi^0_m \]  

(37.13)

where \( N \) is a normalization constant which may now depend on the positions of the quasiholes. Using the plasma analogy, show that the \( w - z \) factor may be obtained by adding additional logarithmically interacting charges at positions \( w_\alpha \) with \( 1/m \) of the charge of each of the \( z \) particles

(c) Note that in this wavefunction the \( z \)'s are physical parameters (and the wavefunction must be single-valued in \( z \)'s), but the \( w \)'s are just parameters of the wavefunction — and so the function \( N \) could be fixed by normalization. Argue using the plasma analogy that in order for the wavefunction to remain normalized (with respect to integration over the \( z \)'s)
as the $w$’s are varied, we must have

$$|\mathcal{N}(w_1, \ldots, w_M)| = K \prod_{1 \leq \alpha < \gamma \leq M} |w_\alpha - w_\gamma|^{1/m} \prod_{1 \leq \alpha \leq M} e^{-|w_\alpha|^2/(4m^2)}$$  \hspace{1cm} (37.14)$$

with $K$ a constant so long as the $w$’s are not too close to each other. (Hint: a plasma will screen a charge).
38.1 Parabolic Confinement

For studying fractional quantum Hall edge states, it is perhaps most useful to consider a parabolic confinement potential. Considering the simple particle Hamiltonian, and adding this confining potential to the kinetic energy we have

\[ H_{\text{confined}} = H_0 + \gamma r^2 \]

where \(H_0\) is the single particle Hamiltonian in the absence of the confinement.

Since the confinement is rotationally symmetric, we can still classify all eigenstates by their angular momentum quantum numbers. Using symmetric gauge we can still write the single particle eigenstates as \(1\)

\[ \psi_m \sim z^m e^{-|z|^2/(4\ell^2)} \]

where \(m\) is the eigenvalue of the angular momentum squared operator \(\hat{L}\). Since the radius of these states is \(r \approx \ell \sqrt{2m}\) it is not surprising that the confinement energy \(\gamma r^2\) of each eigenstate is proportional to \(m\). We thus have

\[ H_{\text{confined}} = H_0 + \alpha \hat{L} \]

for some constant \(\alpha\).

For integer filling, the edge excitations are very much like the edge excitations we discussed above in Landau gauge. A round quantum Hall droplet fills \(m\) states up to a chemical potential along the edge. One can add a small amount of angular momentum to the edge by exciting a filled state from an \(m\) just below the chemical potential to an empty state just above the chemical potential.

38.2 Edges of The Laughlin State

We now consider adding an interaction term so as to produce a fractional quantum Hall state. It is convenient to think about the limit where the cyclotron energy is huge (so we are restricted to the lowest Landau level), the interaction energy is large, so we have a very well formed quantum Hall state, and finally, the edge confinement is weak.

In particular if we choose to consider the special ultra-short range interaction potentials (such as \(\delta\) function for bosons at \(\nu = 1/2\)) we still
have the ground state given exactly by the Laughlin state

$$\Psi^{(m)}_{\text{Laughlin}} = \prod_{i<j} (z_i - z_j)^m \prod_{i=1}^{N} e^{-|z_i|^2/(4\ell^2)}$$

such that it has zero interaction energy. The angular momentum of the Laughlin ground state is just the total degree of the polynomial

$$L_{\text{ground}} = m \frac{N(N - 1)}{2}$$

with confinement energy

$$E_{\text{ground}} = \alpha m \frac{N(N - 1)}{2}$$

While the Laughlin state has zero interaction energy it is also the case that any polynomial times the Laughlin state also has zero interaction energy since multiplying by a polynomial does not ruin the fact that the wavefunction vanishes as $m$ or more powers as two particles approach each other. Thus we can consider all possible wavefunctions of the form

$$\Psi = (\text{Any Symmetric Polynomial}) \Psi^{(m)}_{\text{Laughlin}}$$

where we insist that the polynomial is symmetric such that the symmetry of the wavefunction remains the same (i.e., antisymmetric for fermions and symmetric for bosons).

If the degree of the symmetric polynomial is $\Delta L$, then we have

$$L = L_{\text{ground}} + \Delta L$$
$$E = E_{\text{ground}} + \alpha \Delta L$$

We can organize the possible excitations by their value of $\Delta L$. We thus only need to enumerate all possible symmetric polynomials that we can write in $N$ variables of some given degree $\Delta L$.

We thus need some facts from the theory of symmetric polynomials. The symmetric polynomials on the $N$ variables $z_1, \ldots, z_N$ form a so-called “ring” (this means you can add and multiply them). A set of generators for this ring is given by the functions

$$p_m = \sum_{i=1}^{N} z_i^m$$

This means that any symmetric function on $N$ variables can be written as sums of products of these functions$^3$. Thus it is extremely easy to count symmetric functions. Of degree 1, we have only $p_1$. At degree 2, we have $p_1^2$ and also $p_2$. Thus the vector space of symmetric polynomials of degree two (with real coefficients) is two dimensional. We can build a corresponding table as shown in Table 38.1.

Thus the number of edge excitations at a given angular momentum
follows a pattern, 1, 2, 3, 5, 7, ... with energy increasing linearly with the added angular momentum. Note that this result holds also for the $\nu = 1$ Laughlin state (i.e., for the integer quantum Hall effect), and matches the counting for excitations of a chiral fermion (try this exercise\(^4\))

\subsection*{38.2.1 Edge Mode Field Theory: Chiral Boson}

An equivalent description of the edge modes is given by the Hamiltonian

$$ H = \sum_{m>0} (\alpha m) b_m^\dagger b_m $$

where the $b_m^\dagger$ are boson creation operators satisfying the usual commutations

$$ [b_m, b_n^\dagger] = \delta_{mn} $$

and we think of these boson creation operators $b_m^\dagger$ as creating an elementary excitation of angular momentum $m$ on the ground state which we will call $|0\rangle$ for now. We can build a table describing all of the states in Fock space of this Hamiltonian, ordered by their angular momentum as shown in Table 38.2. We see the Fock space is precisely equivalent to the above table of polynomials. In fact, the analogy is extremely precise. In the thermodynamic limit, up to a known normalization constant, application of $b_m^\dagger$ is precisely equivalent to multiplication of the wavefunction by $p_m$.

These operators describe a chiral boson – chiral because they only have

\(^4\)To get you started, consider filled states in a line filled up to the chemical potential. We can think of these as dots in a row. For example, let the ground state be

\[ \ldots \bullet \bullet \bullet \bullet \bullet \circ \circ \circ \circ \circ \circ \ldots \]

where $\bullet$ means a filled single particle eigenstate and $\circ$ means empty. Now if we add one unit of (angular) momentum, we have the unique state

\[ \ldots \bullet \bullet \bullet \bullet \bullet \circ \circ \circ \circ \circ \circ \ldots \]

adding two units can be done in two ways

\[ \ldots \bullet \bullet \bullet \bullet \bullet \bullet \circ \circ \circ \circ \circ \circ \circ \ldots \]

and

\[ \ldots \bullet \bullet \bullet \bullet \bullet \circ \circ \circ \circ \circ \circ \circ \circ \ldots \]

thus starting the series 1, 2, 3, 5, 7, ...
Fractional Quantum Hall Edges

<table>
<thead>
<tr>
<th>$L - L_{ground}$</th>
<th>dimension</th>
<th>basis fock states</th>
<th>Energy</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>$b_1^0$</td>
<td>$\alpha$</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>$b_2^0$, $b_2^1$</td>
<td>$2\alpha$</td>
</tr>
<tr>
<td>3</td>
<td>3</td>
<td>$b_3^0$, $b_3^1$, $b_3^2$</td>
<td>$3\alpha$</td>
</tr>
<tr>
<td>4</td>
<td>5</td>
<td>$b_4^0$, $b_4^1$, $b_4^2$, $b_4^3$, $b_4^4$</td>
<td>$4\alpha$</td>
</tr>
</tbody>
</table>

Table 38.2 Fock Space for Chiral Bosons

positive angular momentum $m > 0$ not negative angular momentum

38.3 Appendix: Edges and Chern-Simons theory

The existence of the edge theory could have been predicted from the effective Chern-Simons Lagrangian of the bulk. As mentioned previously, the Abelian Chern-Simons action is gauge invariant on a closed manifold. However, for a manifold with boundary, the action is not gauge invariant. This is what is known as an anomaly. The solution to this problem is that the action becomes gauge invariant only once it is added to an action for the low energy edge theory! We will not go through the detailed argument for this here.

---

$^5$An achiral boson field on a circle requires both positive and negative angular momentum modes).
Conformal Field Theory Approach to Fractional Quantum Hall Effect

In the last chapter we saw that we have an edge theory which is a chiral boson — a 1+1 dimensional dynamical theory. We can think of this theory as being a 2 dimensional cut out of a 3 dimensional space-time manifold. Now in a well-behaved topological theory, it should not matter too much how we cut our 3-dimensional space-time manifold. Thus we expect that the same chiral boson theory should somehow also be able to describe our 2+0 dimensional wavefunction. Since all chiral topological theories have gapless edges, this approach can be quite general.

1+1 dimensional gapless theories can all be described by conformal field theories (CFTs) possibly perturbed by irrelevant operators. And conformal field theories in 1+1 dimension are particularly powerful in that they are exactly solvable models, which can be used to describe either the dynamics of 1+1 dimensional systems or classical statistical mechanical models in 2 dimensions.

While we cannot provide a complete introduction to CFT here (see Ginsparg’s lectures, Fendley’s notes, or for a much more complete discussion, see the Big Yellow Book), it turns out that we need very little of the machinery to proceed. Furthermore, a large fraction of this machinery will look extremely familiar from our prior study of TQFTs. Indeed, there is an extremely intimate connection between CFTs and TQFTs — and much of what we know about TQFTs has grown out of the study of CFTs.

We will begin by seeing how this works for the chiral boson, which is perhaps the simplest of all 1+1d CFTs. Below we will show how the scheme works in more detail in the context of quantum Hall physics. This approach, first described by Moore and Read, has been extremely influential in the development of TQFTs and their relationship to the quantum Hall effect.

39.1 The Chiral Boson and The Laughlin State

An interesting feature of theories in 1+1d is that they can often be decomposed (mostly\(^1\)) cleanly into right moving and left moving pieces. So for example, if we take the simplest possible 1+1 d system, a free

\(^1\)There may be issues with the decomposition, for example, in the case of the boson, there is a complication associated with the so-called zero-mode, which we will ignore for simplicity.
boson, we can write an achiral Lagrangian density for a field \( \Phi(x, t) \) as
\[
\mathcal{L} \propto (\partial_{\mu} \Phi)(\partial^{\mu} \Phi)
\]
This can be decomposed into right and left moving pieces as
\[
\Phi(x, t) = \phi(x - vt) + \bar{\phi}(x + vt)
\]
where \( \phi \) is right-moving and \( \bar{\phi} \) is left-moving and these are two different fields. For simplicity we will set the velocity \( v = 1 \).

In the previous chapter we deduced that the edge theory of the Laughlin state could be described by a chiral boson Hamiltonian
\[
H = \sum_{m>0} (\alpha m) a_{m}^\dagger a_{m} + h.c.
\]
Quantizing the boson lagrangian we find that\(^2\)
\[
\phi(x) = \sum_{m>0} \frac{i}{\sqrt{m}} e^{2\pi i mx/L} a_{m}^\dagger + h.c. \quad \text{(39.1)}
\]
where \( L \) is the (periodic) length of the system.

We will often work in complex coordinates \( x + i\tau \), so we have
\[
\Phi(z, z^*) = \sum_{m>0} \frac{i}{\sqrt{m}} e^{2\pi i mx/L} a_{m}^\dagger + h.c.
\]
Note that we think of this correlation function as a correlation in a 1+1d theory even though we are working with complex \( z \).

From this chiral \( \phi \) operator we construct the so-called vertex operators
\[
V_{\alpha}(z) = \mathrm{e}^{i\alpha \phi(z)} \quad \text{where} \quad \mathrm{e}^{...}
\]
where \( \mathrm{e}^{...} \) means normal ordering\(^4\) A straightforward exercise (assigned as homework!) using Wick’s theorem then shows that\(^3\)
\[
\langle V_{\alpha_{1}}(z_{1})V_{\alpha_{2}}(z_{2}) \ldots V_{\alpha_{N}}(z_{N}) \rangle = e^{-\sum_{i<j} \alpha_{i}\alpha_{j}\langle \phi(z_{i})\phi(z_{j}) \rangle} = \prod_{i<j} (z_{i} - z_{j})^{\alpha_{i}\alpha_{j}} \quad \text{(39.2)}
\]
so long as
\[
\sum_{i} \alpha_{i} = 0 \quad \text{(39.3)}
\]
(otherwise the correlator vanishes).
39.1 Writing the Laughlin Wavefunction

We then define an “electron operator” to be

$$\psi_e(z) = V_\alpha(z)$$

where we will choose

$$\alpha = \sqrt{m}$$

This then enables us to write the holomorphic part of the Laughlin wavefunction as

$$\Psi_{\text{Laughlin}}^{(m)} = \langle \psi_e(z_1) \psi_e(z_2) \ldots \psi_e(z_N) \hat{Q} \rangle = \prod_{i<j} (z_i - z_j)^m$$

The index $\alpha$ must be chosen such that $\alpha^2$ is an integer such that the wavefunction is single valued in the electron coordinates. Note that here although the correlator means a 1+1d theory, we are constructing a wavefunction for a 2d system at fixed time!

Here, the operator $\hat{Q}$ can be chosen in two different ways. One possibility is to choose $\hat{Q} = V_{-N\alpha}$, i.e., a neutralizing charge at infinity such that Eq. 39.3 is satisfied and the correlator does not vanish. This approach is often used if one is only concerned with keeping track of the holomorphic part of the wavefunction (which we often do). A more physical (but somewhat more complicated) approach is to smear this charge uniformly over the system. In this case, the neutralizing charge, almost magically, reproduces precisely the gaussian factors that we want!\footnote{To see how this works, we divide the background charge into very small pieces (call them $\beta$) to obtain a correlator of the form

$$e^{-m \sum_{i<j} \log(|z_i - z_j|) - \epsilon \sqrt{m} \sum_{i,\beta} \log(|z_i - z_\beta|)}$$

the term with $\epsilon^2$ we throw away as we will take the limit of small $\epsilon$. Now here we realize that we are going to have a problem with branch cuts around these small charges — which we can handle if we work in a funny gauge. Changing gauge to get rid of the branch cuts we then get only the real part of the second term. The second term is then of the form

$$\sum_{i,\beta} \log(|z_i - z_\beta|) \to \int d^2r \log(|z - r|)$$

where we have taken the limit of increasing number of smaller and smaller charges. We define this integral to be $f(z)$. It is then easy to check that $f(z) \sim |z|^2$ which is most easily done by taking $\nabla^2 f(z)$ and noting that log is the coulomb potential in 2d so Gauss’s law just gives the total charge enclosed. Thus we obtain $e^{-|z|^2}$ as desired. A more careful calculation gives the constant correctly as well.}

39.1.2 Quasiholes

Let us now look for quasihole operators. We can define another vertex operator

$$\psi_{qh}(w) = V_\beta(w)$$

and now insert this into the correlator as well to obtain

$$\Psi_{qh}(w) = \langle \psi_{qh}(w) \psi_e(z_1) \psi_e(z_2) \ldots \psi_e(z_N) \hat{Q} \rangle$$

$$= \prod_i (z_i - w)^{\beta \sqrt{m}} \Psi_{\text{Laughlin}}^{(m)}$$

Since we must insist that the wavefunction is single valued in the $z$ coordinates, we must choose

$$\beta = \frac{p}{\sqrt{m}}$$

for some positive integer $p$, where the minimally charged quasiparticle is then obviously $p = 1$. (Negative $p$ is not allowed as it would create poles in the wavefunction).

Further, using this value of the charge $\beta$, along with the smeared out background charge, we correctly obtain the normalizing gaussian
factor for the quasiparticle
\[ e^{-|w|^2/(4m\ell^2)} \]

This is the correct gaussian factor, with an exponent 1/m times as big because the charge \( V_{1/\sqrt{m}} \) is 1/m times as big as that of the electron charge \( V_{\sqrt{m}} \).

If we are now to add multiple quasiholes, we obtain the wavefunction
\[
\Psi(w_1, \ldots, w_M) = \langle \psi_{qh}(w_1) \ldots \psi_{qh}(w_M) \psi_e(z_1) \ldots \psi_e(z_N) Q \rangle (39.5)
\]
\[
= C \prod_{\alpha<\beta} (w_\alpha - w_\beta)^{1/m} \prod_{\alpha=1}^M e^{-|w_\alpha|^2/(4\ell^2)} \left[ \prod_{\alpha=1}^M \prod_{i=1}^N (z_i - w_\alpha) \right] \Psi^{(m)}_{\text{Laughlin}}
\]

which is properly normalized
\[
\langle \Psi(w_1, \ldots, w_M) | \Psi(w_1, \ldots, w_M) \rangle = \text{Constant}
\]

and is in holomorphic gauge. As discussed previously in chapter *** with a normalized holomorphic wavefunction we can simply read off the fractional statistics as the explicit monodromy.

Note that we can consider fusion of several quasiparticles
\[
V_{1/\sqrt{m}} \times V_{1/\sqrt{m}} \to V_{2/\sqrt{m}}
\]

Fusion of \( m \) of these elementary quasiholes produces precisely one electron operator \( V_{2/\sqrt{m}} \). Since the electrons are “condensed” into the ground state, we view them as being essentially the identity operator, at least in the case of \( m \) even, which means we are considering a Laughlin state of bosons. Thus there are \( m \) species of particle in this theory. In the case of \( m \) odd, we run into the situation mentioned in chapter *** where the electron is a fermion, so really there are \( 2m \) species of particles in the theory.

The idea is that by using conformal field theory vertex operators we automatically obtain normalized holomorphic wavefunctions and we can determine the statistics of quasiparticles straightforwardly. This is a key feature of the Moore-Read approach. While there is no general proof that this will always be true (that the resulting wavefunctions will be properly normalized) it appears to hold up in many important cases.

We hope now to generalize this construction by using more complicated conformal field theories. This then generates more complicated fractional quantum Hall wavefunctions corresponding to more complicated TQFTs.
What We Need to Know About Conformal Field Theory

I can’t possibly explain CFT in a few pages. (See the big yellow book. Ginsparg’s lectures are nice for introduction. So are Fendley’s notes), but given what we already know about TQFTs many of the rules are going to seem very natural. Indeed, much of the math of TQFTs arose via CFTs.

CFTs are quantum theories in 1+1 dimension. They are generically highly interacting theories, and most often it is impossible to write an explicit Lagrangian for the theory, but due to the special properties of being in 1+1 and having conformal invariance (guaranteed by being gapless in 1+1 d) these models are exactly solvable.

A particular CFT is defined by certain information known as conformal data, which basically mimics the defining features of a TQFT:

1. There will be a finite set of so-called primary fields, which we might call \( \phi_i(z) \) (or we may use other notation). These are analogous to the particle types in a TQFT. Every CFT has an identity field often called \( I \) (which isn’t really a function of position). Correlators of these fields

\[
\langle \phi_{j_1}(z_1) \ldots \phi_{j_N}(z_N) \rangle
\]

are always holomorphic functions of the \( z \) arguments, although there may be branch cuts.

2. Each primary field has a scaling dimension or conformal weight or conformal spin, which we call \( h_i \). The scaling dimension of \( I \) is \( h_I = 0 \). We have see these quantities before when we discussed twists in world lines. Often we will only be interested in \( h \) modulo 1, since the twist factor is \( e^{2\pi i h} \). Each primary field has descendant fields which are like derivatives of the primary and they have scaling dimensions \( h_i \) plus an integer (we will typically not need these, but for example, \( \partial_z \phi_i \) has scaling dimension \( h_i + 1 \)).

3. Fusion relations exist for these fields, which are associative and commutative

\[
\phi_i \times \phi_j = \sum_k N_{ij}^k \phi_k
\]

where fusion with the identity is trivial

\[
I \times \phi_j = \phi_j
\]

In CFT we have the powerful relation that if we make a coordinate transform \( w(z) \) then any correlator of primary fields transforms as

\[
\langle \phi_{i_1}(w_1) \ldots \phi_{i_N}(w_N) \rangle = \left( \frac{\partial w_1}{\partial z_1} \right)^{-h_{i_1}} \ldots \left( \frac{\partial w_N}{\partial z_N} \right)^{-h_{i_N}} \langle \phi_{i_1}(z_1) \ldots \phi_{i_N}(z_N) \rangle
\]

However, we will not need this relationship anywhere for our discussion!
As with TQFTs, each particle type has a unique antiparticle. We will give a clearer meaning to these fusion relations in a moment when we discuss operator product expansion.

The expectation of any correlator in the theory is zero unless all the fields inside the correlator fuse to the identity. For example, if we have a \( \mathbb{Z}_3 \) theory where it requires three \( \psi \) particles fuse to the identity, then we would have \( \langle \psi(z) \psi(w) \rangle = 0 \). We saw this law previously in the neutrality condition for the chiral boson. The expectation of the identity \( I \) is unity.

The fundamental theorem we need, which is beyond the simple analogy with TQFT is the idea of an operator product expansion. The idea is that if you take two field operators in a conformal field theory and you put them close together, the product of the two fields can be expanded as sum of resulting fields

\[
\lim_{w \to z} \phi_i(w) \phi_j(z) = \sum_k C^k_{ij} (w - z)^{h_k - h_i - h_j} \phi_k(z) + \ldots
\]

Here the \( C^k_{ij} \) are coefficients which crucially are zero when \( N^k_{ij} \) is zero. In other words, when two fields are taken close together, the result looks like a sum of all the possible fusion products of these field. On the right hand side note that by looking at the scaling dimensions of the fields, we obtain explicit factors of \( (w - z) \). The \( \ldots \) terms are terms that are smaller (less singular) than the terms shown and are made of descendant fields and higher powers of \( (w - z) \). Crucially, no new types of branch cuts are introduced except those that differ by integers powers from (and are less singular than) those we write explicitly.

The convenient thing about the operator product expansion (or “OPE”) is that it can be used inside expectation values of a correlator. So for example

\[
\lim_{w \to z} \langle \psi_a(w) \psi_b(z) \psi_c(y_1) \psi_d(y_2) \ldots \psi_n(y_m) \rangle = \sum_k C^k_{ab} (w - z)^{h_k} \psi_c(y_1) \psi_d(y_2) \ldots \psi_n(y_m) \]

### 39.2.1 Example: Chiral Boson

The free boson vertex \( V_\alpha \) has scaling dimension

\[
h_\alpha = \frac{\alpha^2}{2}
\]

The fusion rules are

\[
V_\alpha V_\beta = V_{\alpha + \beta}
\]

corresponding to the simple addition of “charges”. The resulting operator product expansion is then

\[
V_\alpha(w)V_\beta(z) \sim (w - z)^{\alpha \beta} V_{\alpha + \beta}(z)
\]
where we have used the notation \( \sim \) to mean in the limit where \( w \) goes to \( z \), and where the exponent is here given as

\[
h_{\alpha+\beta} - h_{\alpha} - h_{\beta} = \frac{(\alpha + \beta)^2}{2} - \frac{\alpha^2}{2} = \alpha \beta
\]

Note that this fusion law for the chiral boson gives more precise meaning to the fusion law we wrote in Eq. 39.6.

### 39.2.2 Example: Ising CFT

The Ising CFT is actually the CFT corresponding to a 1+1 d free fermion, so it is particularly simple. The theory has three fields, \( I, \sigma, \psi \) with scaling dimensions

\[
h_I = 0 \\
h_\sigma = 1/16 \\
h_\psi = 1/2
\]

The fact that \( h_\psi = 1/2 \) is an indication that it is a fermion. The nontrivial fusion rules are (exactly as in the Ising TQFT previously)

\[
\psi \times \psi = I \\
\psi \times \sigma = \sigma \\
\sigma \times \sigma = I + \psi
\]

As in the case of TQFTs, it is the multiple terms on the right hand side that make a theory nonabelian.

We can write the operator product expansion

\[
\psi(w)\psi(z) \sim (w - z)^{h_{\psi} - h_\psi} I + \ldots \\
\sim \frac{I}{w - z} + \ldots
\]

The antisymmetry on the right hand side is precisely the behavior one should expect from fermions. It is crucial to note that within the ... all terms are similarly antisymmetric (and are less singular). Similarly, we have

\[
\psi(w)\sigma(z) \sim (w - z)^{h_{\sigma} - h_\psi} \sigma(z) + \ldots \\
\sim (w - z)^{-1/2} \sigma(z) + \ldots
\]

where again the ... indicates terms which have the same branch cut structure but are less singular. In other words, wrapping \( w \) around \( z \) should incur a minus sign for all terms on the right.

Finally we have the most interesting OPE

\[
\sigma(w)\sigma(z) \sim C^I_{\sigma \sigma}(w - z)^{-1/8} I + C^\psi_{\sigma \sigma}(w - z)^{3/8} \psi(z) + \ldots
\]

where all terms in the ... must have branch cuts that match one of the...
two leading terms.

Let us consider calculating a correlator,
\[ \lim_{w \to z} \langle \sigma(w) \sigma(z) \rangle \]

Since from rule (4) above, the two fields must fuse to the identity, we must choose the identity fusion channel only from the OPE. We then obtain
\[ \lim_{w \to z} \langle \sigma(w) \sigma(z) \rangle \sim (w - z)^{-1/8} \quad (39.8) \]

On the other hand, calculating
\[ \lim_{w \to z} \langle \sigma(w) \sigma(z) \psi(y) \rangle \]
in order to fuse to the identity, we must choose the \( \psi \) fusion of the two \( \sigma \) fields such that this \( \psi \) can fuse with \( \psi(y) \) to give the identity. We thus have
\[ \lim_{w \to z} \langle \sigma(w) \sigma(z) \psi(y) \rangle \sim (w - z)^{3/8} \quad (39.9) \]

Similarly one can see that fusion of two \( \sigma \)'s in the presence of any even number of \( \psi \) fields will be similar to Eq. 39.8 whereas in the presence of any odd number of \( \psi \) fields it will be like Eq. 39.9.

Since the Ising CFT is actually a free fermion theory, we can use Wick's (fermionic) theorem for correlators of the \( \psi \) fermi fields with the added information that
\[ \langle \psi(z) \psi(w) \rangle = \frac{1}{z - w} \]

which is exactly true, not only in the OPE sense. However, we cannot use Wick's theorem on correlators of the \( \sigma \) fields which are sometimes known as “twist” fields — we can think of these as altering the boundary conditions.

39.3 Quantum Hall Wavefunction Based on Ising CFT: The Moore-Read State

Let us try to build a quantum Hall wavefunction based on the Ising CFT. We must first choose a field which will represent our electron. One might guess that we should use the fermion field. However, when two \( \psi \) fields come together the correlator (and hence our wavefunction) diverges, so this cannot be acceptable. Instead, let us construct an electron field which is a combination of the Ising \( \psi \) field and a chiralbose vertex \( V_\alpha \)
\[ \psi_e(z) = \psi(z)V_\alpha(z) \]

These two fields are from completely different 1+1d theories and are simply multiplied together.

We then look at the operator product expansion to see what happens.
when two electrons approach each other
\[ \psi_e(z)\psi_e(w) \sim \left[ \frac{1}{z - w} \right] (z - w)\alpha^2 V Z \]
where the first bracket is from the Ising part of the theory and the second bracket is from the boson part of the theory. In order for this to not be singular, we must have \( \alpha^2 \) be a positive integer. If we choose
\[ \alpha^2 = m \]
with \( m \) odd we have an overall bosonic operator \( \psi_e(z)\psi_e(w) = \psi_e(w)\psi_e(z) \)
wheras if we choose \( m \) even we have an overall fermionic operator. However, we cannot choose \( m = 0 \) since that leaves a singularity. Thus we have the electron operator of the form
\[ \psi_e(z) = \psi(z)V_{\sqrt{m}}(z) \]
with \( m \geq 1 \). Using this proposed electron operator we build the multi-particle wavefunction
\[ \Psi = \langle \psi(z_1)\psi(z_2)\ldots\psi(z_N)Q \rangle \]
where \( Q \) is the background charge for the boson field. Since the Ising and boson fields are completely separate theories we can take the expectation for the boson field to give
\[ \Psi = \langle \psi(z_1)\psi(z_2)\ldots\psi(z_N) \rangle \prod_{i<j}(z_i - z_j)^m \prod_{i=1}^N e^{-|z_i|^2/(4\ell^2)} \]
where the correlator is now in the Ising theory alone.

Now the Ising correlator must be zero unless there are an even number of \( \psi \) fields (since we need them to fuse to the identity). If the number of fermi fields is indeed even, then we can use the fact that \( \psi \) is a free fermi field and we can invoke Wick’s theorem to obtain
\[ \langle \psi(z_1)\psi(z_2)\ldots\psi(z_N) \rangle = \mathcal{A} \left[ \frac{1}{z_1 - z_2} \frac{1}{z_3 - z_4} \ldots \frac{1}{z_{N-1} - z_N} \right] \]
\[ \equiv \text{Pf} \left( \frac{1}{z_i - z_j} \right) \quad (39.10) \]
Here \( \mathcal{A} \) means antisymmetrize over all reordering of the \( z \)'s. Here we have written the usual notation for this antisymmetrized sum Pf which stands for “Pfaffian”\(^\text{12}\). Thus we obtain the trial wavefunction based on the Ising CFT
\[ \Psi = \text{Pf} \left( \frac{1}{z_i - z_j} \right) \prod_{i<j}(z_i - z_j)^m \prod_{i=1}^N e^{-|z_i|^2/(4\ell^2)} \]
which is known as the Moore-Read wavefunction. For \( m \) odd this is a

\(^\text{12}\) Several interesting facts about the Pfaffian: A BCS wavefunction for a spinless superconductor can be written as \( \text{Pf}[g(r_i - r_j)] \) where \( g \) is the wavefunction for a pair of particles. Any antisymmetric matrix \( M_{ij} \) has a Pfaffian
\[ \text{Pf}[M] = \mathcal{A}[M_{12}M_{34}\ldots]. \]
Also it is useful to know that \( (\text{Pf}[M])^2 = \text{det}M \).
wavefunction for bosons and for \( m \) even it is a wavefunction for fermions. To figure out the filling fraction, we note that the Pfaffian prefactor only removes a single power in each variable. Thus the filling fraction is determined entirely by the power \( m \), and is given (like Laughlin) by 
\[
\nu = \frac{1}{m}.
\]

### 39.3.1 Some Exact Statements About the Moore-Read Wavefunction

For simplicity, let us consider the \( m = 1 \) case \( \nu = 1 \) for bosons, which is the easiest to think about analytically. The wavefunction does not vanish when two particles come to the same point, since the zero of \((z_1 - z_2)\) can be canceled by the pole of the Pfaffian. However, it is easy to see that the wavefunction must vanish (quadratically) when three particles come to the same point (three factors from \((z - z)^1\) but then one factor in the denominator of the Pfaffian).

Note that, even were we to not have an explicit expression for the Moore-Read wavefunction we would still be able to use the operator product expansion to demonstrate that the wavefunction (for \( m = 1 \)) must vanish quadratically when three particles come to the same point\(^{13}\).

To see this, note that taking the first two particles to the same point gives
\[
\lim_{z_2 \to z_1} \psi_e(z_1) \psi_e(z_2) \sim IV_2(z_1)
\]

Then fusing the third particle
\[
\lim_{z_3 \to z_1} \psi_e(z_3) V_2(z_1) \sim (z_3 - z_1)^2 \psi V_3(z_1)
\]

Analogous to the case of the Laughlin wavefunction, it turns out that the Moore-Read wavefunction (for \( m = 1 \)) is the exact (highest density) zero energy ground state of a three-body delta function interaction
\[
V = V_0 \sum_{i<j<k} \delta(r_i - r_j) \delta(r_i - r_k)
\]

Similarly one can construct a potential for fermions such that the \( \nu = 1/2 \) Moore-Read state (\( m = 2 \)) is the highest density zero energy state. This is quite analogous to what we did for the Laughlin state:
\[
V = V_0 \sum_{i<j<k} [\nabla^2 \delta(r_i - r_j)] \delta(r_i - r_k)
\]

### Non-Exact Statements

Although the Coulomb interaction looks nothing like the three body interaction for which the Moore-Read Pfaffian is exact, it turns out that \( \nu = 1/2 \) Moore-Read Pfaffian \( m = 2 \) is an extremely good trial state\(^{14}\) for electrons at \( \nu = 5/2 \) interacting with the usual Coulomb interaction. This is very suggestive that the \( \nu = 5/2 \) is topologically equivalent to the Moore-Read Pfaffian wavefunction (i.e., they are in the same phase of matter)\(^{15}\). Further, the most natural interaction for bosons, the simple two-body delta function interaction has a ground state at \( \nu = 1 \) which is extremely close to the Moore-Read \( m = 2 \) Pfaffian.

\(^{13}\)To see this, note that taking the first two particles to the same point gives
\[
\lim_{z_2 \to z_1} \psi_e(z_1) \psi_e(z_2) \sim IV_2(z_1)
\]

Then fusing the third particle
\[
\lim_{z_3 \to z_1} \psi_e(z_3) V_2(z_1) \sim (z_3 - z_1)^2 \psi V_3(z_1)
\]

\(^{14}\)Here we have used a mapping between Landau levels, that any partially filled higher Landau level can be mapped to a partially filled lowest Landau level at the price of modifying the inter-electron interaction. This mapping is exact to the extent that there is no Landau level mixing. I.e., that the spacing between Landau levels is very large.

\(^{15}\)There is one slight glitch here. It turns out that with a half-filled Landau level, the wavefunction and its charge-conjugate (replace electrons by holes in the Landau level) are inequivalent! The breaking of the particle-hole symmetry is very weak and involves Landau-level mixing. From numerics it appears that the \( \nu = 5/2 \) state is actually in the phase of matter defined by the conjugate of the Moore-Read state. **add refs**
39.4 Quasiholes of the Moore-Read state

We now try to construct quasiholes for the Moore-Read Pfaffian wavefunction. As we did in Eq. 39.4, we want to write
\[ \Psi_{qh}(w) = \langle \psi_{qh}(w) \psi_e(z_1) \psi_e(z_2) \ldots \psi_e(z_N) \hat{Q} \rangle \]
but we need to figure out what the proper quasihole operator \( \psi_{qh} \) is.

Laughlin Quasihole

One obvious thing to try would be to write a simple vertex operator
\[ \psi_{qh}^L(w) = V_\beta(w) \]
Looking at the OPE we have (**include fields on the right? **)
\[ \psi_{qh}^L(w) \psi_e(z) \sim (w-z)^{\beta \sqrt{m}} \psi(z) \]
In order to have the correlator be single valued in \( z \) (i.e., no branch cuts) we must choose \( \beta = p/\sqrt{m} \) for some integer \( p \) (the smallest quasihole of this type corresponding to \( p = 1 \) then). This generates the wavefunction
\[ \Psi_{qh}^L(w) = \langle \psi_{qh}^L(w) \psi_e(z_1) \psi_e(z_2) \ldots \psi_e(z_N) \hat{Q} \rangle \]
(39.11)
\[ = \prod_{i=1}^{N} (z_i - w) \psi^{(m)}_{Moore-Read} \]
which is just a regular Laughlin quasihole factor. By the same arguments, the charge of this quasihole is \( e^* = e\nu \).

Minimal quasihole

However, the Laughlin quasihole is not the minimal quasihole that can be made. Let us try using an operator from the Ising theory as part of the quasihole operator. Suppose
\[ \psi_{qh}(w) = \sigma(w)V_\beta(w) \]
We then have the operator product expansion
\[ \psi_{qh}(w) \psi_e(z) \sim [\sigma(w)\psi(z)] \left[ V_\beta(w)V_{\sqrt{m}}(z) \right] \sim (w-z)^{-1/2}(w-z)^{\beta \sqrt{m}} \]
In order for the wavefunction not to have any branch cuts for the physical electron \( z \) coordinates, we must choose \( \beta = (p + 1/2)/\sqrt{m} \) for \( p \geq 0 \), with the minimal quasihole corresponding to \( p = 0 \). Thus we have the minimal quasihole operator of the form
\[ \psi_{qh}(w) = \sigma(w)V_{\frac{1}{\sqrt{m}}}(w) \]
Note that when we consider correlators, by the general rule (4) from section 39.2, the operators must fuse to the identity in order to give a nonzero result. Thus, we must always have an even number of $\sigma$ fields.  

We thus consider the wavefunction of the form

$$\Psi_{qh}(w, w') = \langle \psi_{qh}(w) \psi_{qh}(w') \psi_e(z_1) \psi_e(z_2) \ldots \psi_e(z_N) \hat{Q} \rangle$$

(39.12)

$$= (w - w') \frac{1}{4\pi} e^{-(|w|^2+|w'|^2)/4\ell^*} \prod_{i=1}^{N} (w - z_i)^{1/2} (w' - z_i)^{1/2}$$

(39.13)

$$\times \langle \sigma(w)\sigma(w') \psi(z_1) \psi(z_2) \ldots \psi(z_N) \rangle \prod_{i<j} (z_i - z_j)^{\nu} \prod_{i=1}^{N} e^{-|z_i|^2/(4\ell^2)}$$

Several comments are in order here. First of all, from the first line of Eq. 39.13 it looks like there are branch cuts with respect to the $z$ coordinates. However, these fractional powers are precisely canceled by branch cuts in the correlator on the second line. Secondly the charge of the quasihole is determined entirely by the power of the $(z - w)$ factor, since it tells us how much the electrons are pushed away from the hole. (The correlator does not give an extensive number of zeros, similar to the Pfaffian of Eq. 39.10). If the exponent of $(z - w)$ were one, this would be a regular Laughlin quasihole with charge $e\nu$, thus here we have a quasihole charge of

$$e^* = e\nu/2.$$ 

I.e., the Laughlin quasihole has fractionalized into two pieces! This charge is reflected in the effective magnetic length $\ell^* = \sqrt{\hbar/e^* B}$.

Note that this wavefunction is still an exact zero energy state of the special interaction discussed above for which the Moore-Read wavefunction is the exact highest density zero energy state (the wavefunction here is higher degree and thus less dense, as we would expect given that we have added quasiholes). We can demonstrate the current wavefunction is still zero energy by bringing together three electrons to the same point and examining how the wavefunction vanishes. Since this can be fully determined by the operator product expansion, it does not matter if we add quasiholes to the wavefunction, the vanishing property of the wavefunction remains the same, and thus this is an exact zero energy state of the special interaction.

A crucial assumption

The wavefunction here is single valued in all electron coordinates (as it should be) and is holomorphic in all coordinates (all $z$’s and $w$’s) except for the gaussian exponential factors. In this holomorphic gauge, as discussed above, we can read off the fractional statistics of the quasiparticles given the assumption that the wavefunction is properly normalized. This is a crucial assumption and it is not a simple result of CFT, but always requires an assumption about some sort of plasma being in a screening phase — and often the mapping to a plasma is highly non-
trivial. Nonetheless, from extensive numerical work, it appears that physics is kind to us and that these wavefunctions do indeed come out to be properly normalized!

Fusion and Braiding of Two Quasiholes in Identity Channel (even number of electrons)

Let us assume that the number of electrons is even. In this case the two $\sigma$’s of the quasiholes fuse to the identity as in Eq. 39.8. As the two quasiholes approach each other we then have \(^{(1)}\) (** insert also h-h-h derivation of R? **) \[
\psi_{qh}(w)\psi_{qh}(w') \sim (w - w')^{1/4m - 1/8m}
\]
where the $1/4m$ is written explicitly in the first line of Eq. 39.13 and the $-1/8m$ is from the operator product expansion Eq. 39.8. Invoking now the crucial assumption that the wavefunctions are normalized, since they are obviously holomorphic, we simply read off the statistical phase (the monodromy) we get for wrapping one quasihole around another!

One might object that the operator product expansion only tells us the behavior of the correlator as $w$ and $w'$ come close to each other. However, we are guaranteed that there are no other branch cuts in the system — the only branch cut in the wavefunction for $w$ is when it approaches $w'$. Thus, no matter how far $w$ is from $w'$, when $w$ circles $w'$ it must always accumulate the same monodromy! In the notation we defined in earlier chapters we have \(\psi\) (move I downstairs here to fit with our conventions?, change notation “I” to 2qh-I?) \[
[R^I_{qh-qh}]^2 = e^{2\pi i(1/16 - 1/8m)}
\]
Recall that if $a \times b \to c$ we should have $[R^c_{ab}]^2 = e^{2\pi i(h_c - h_a - h_b)}$. Here, the total scaling dimension of the quasihole is $h_{qh} = 1/16 + 1/(8m)$ with the second piece from the bose vertex operator $V_1/\sqrt{m}$. The fusion product $"I" = V_1/\sqrt{m}$ has quantum dimension $h":"I" = 1/2m.$

Fusion and Braiding of Two Quasiholes in $\psi$ Channel (odd number of electrons)

Let us now assume that the number of electrons is odd. In this case the two $\sigma$’s of the quasiholes fuse to $\psi$ as in Eq. 39.9. As the two quasiholes approach each other we then have \(\psi\) \[
\psi_{qh}(w)\psi_{qh}(w') \sim (w - w')^{1/4m + 1/8m}
\]
where the $1/4m$ is written explicitly in the first line of Eq. 39.13 and the $1/8m$ is from the operator product expansion Eq. 39.9. Again we just read off the monodromy from this OPE. Thus, one obtains a different phase depending on the fusion channel of the two quasiholes. In the notation

---

17See work by Bonderson et al **.

18Strictly speaking on the right hand side we should also write the identity operator $I$ for the Ising theory and $V_1/\sqrt{m}$ for the boson sector.

19Strictly speaking on the right hand side we should also write the operator $\psi$ for the Ising theory and $V_1/\sqrt{m}$ for the boson sector.
we defined in earlier chapters we have

\[ R_{q_h-q_h}^{\psi \psi} = e^{2\pi i \left( \frac{q_h}{q} + \frac{q}{q} \right)} \]

### 39.5 Multiple Fusion Channels and Conformal Blocks

We will next address the issue of what happens when we have more than two quasiholes. It is clear what will happen here, we will obtain a correlator (like that in Eq. 39.13) but now it will have more \( \sigma \) fields. We will thus have to figure out how to make sense of correlators with many (nonabelian) \( \sigma \) fields. As an example to show how this works, let us get rid of the \( \psi \) fields for a moment and consider a correlator

\[ G(w_1, w_2, w_3, w_4) = \langle \sigma(w_1)\sigma(w_2)\sigma(w_3)\sigma(w_4) \rangle \]

Let us imagine that we will bring \( w_1 \) close to \( w_2 \) and \( w_3 \) close to \( w_4 \). Now in order for the correlator to give a nonzero value, the four fields have to fuse to unity (rule (4) from section 39.2). There are two different ways in which this can happen

\[
\sigma(w_1)\sigma(w_2) \rightarrow I \\
\sigma(w_3)\sigma(w_4) \rightarrow I
\]

OR we could have

\[
\sigma(w_1)\sigma(w_2) \rightarrow \psi \\
\sigma(w_3)\sigma(w_4) \rightarrow \psi
\]

and the two \( \psi \) fields could then fuse to the identity.

So which one is right? In fact both happen at the same time! To understand this we should think back to what we know about a 2d systems with nonabelian quasiparticles in them — they are described by a vector space. In order to know which particular wavefunction we have in a vector space we need some sort of initial condition or space-time history. Nowhere in the correlator have we specified any space-time history, so we should be getting a vector space rather than a single wavefunction. The multiple wavefunctions in the vector space arise from choosing different roots of the branch cuts of the holomorphic functions. To see a detailed example of this let us write out the explicit form of the correlator in Eq. 39.14. We note that the calculation that leads to this requires some substantial knowledge of conformal field theory and will not be presented here. However many of these sorts of results have simply been tabulated in books and can be looked up when necessary.

\[ \text{In fact due to conformal invariance, knowing the correlator for any fixed three points and one point } z \text{ free, we can determine the correlator for any other four points, but this is beyond the scope of the current discussion!} \]
\[
\lim_{w \to \infty} \langle \sigma(0) \sigma(z) \sigma(1) \sigma(w) \rangle = a_+ G_+(z) + a_- G_-(z) \quad (39.15)
\]

where
\[
G_{\pm} = (wz(1-z))^{-1/8} \sqrt{1 \pm \sqrt{1-z}} \quad (39.16)
\]

are known as conformal blocks and here \(a_+\) and \(a_-\) are arbitrary complex coefficients (usually with some normalization condition implied). I.e, the correlator itself represents not a function, but a vector space (with basis vectors being conformal blocks) with arbitrary coefficients yet to be determined by the history of the system!

Let us analyze some limits to see which fusion channels we have here. Taking the limit of \(z \to 0\) we find that
\[
\begin{align*}
\lim_{z \to 0} G_+ &\sim z^{-1/8} \quad (\sigma(0) \sigma(z) \to I) \\
\lim_{z \to 0} G_- &\sim z^{3/8} \quad (\sigma(0) \sigma(z) \to \psi)
\end{align*}
\]

Thus (comparing to Eqs. 39.8 and 39.9) we see that \(G_+\) has \(\sigma(0)\) and \(\sigma(z)\) fusing to \(I\) whereas \(G_-\) has them fusing to \(\psi\). Since the four \(\sigma\)'s must fuse to the identity, this tells us also the fusion channel for \(\sigma(1)\) and \(\sigma(w)\).

The most general wavefunction is some linear combination \((a_+ + a_-)\) of the two possible fusion channels. This is what we expect, the state of a system can be any superposition within this degenerate space.

Now consider what happens as we adiabatically take the coordinate \(z\) in a circle around the coordinate 1. Looking at Eq. 39.16 we see that we accumulate a phase of \(e^{-2\pi i/8}\) from the factor of \((1-z)^{-1/8}\) outside the square-root. In addition, however, the \(\sqrt{1-z}\) inside the square root comes back to minus itself when \(z\) wraps around 1, thus turning \(G_+\) to \(G_-\) and vice versa! The effect of monodromy (taking \(z\) around 1) is then
\[
\begin{pmatrix} a_+ \\ a_- \end{pmatrix} \longrightarrow e^{-2\pi i/8} \begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix} \begin{pmatrix} a_+ \\ a_- \end{pmatrix}
\]

(This result should be somewhat familiar from the homework exercise on Ising anyons!)

We thus see that in this language, the multiple fusion channels are just different choices of which Riemann sheet we are considering, and the fact that braiding (monodromy) changes the fusion channel is simply the fact that moving coordinates around on a Riemann surface, you can move from one Riemann sheet to another!

So long as we can assume that the conformal blocks are orthonormal (see comment above on “crucial assumption” about normalization of wavefunctions. Orthonormality, is now adding a further assumption\(^{21}\)) then we can continue to read off the result of physically braiding the particles around each other by simply looking at the branch cuts in the wavefunction.

\(^{21}\) As with the discussion above, this assumption appears to be true, but “proofs” of it always boil down to some statement about some exotic plasma being in a screening phase, which is hard to prove. *** myabe move bon-derson ref here!
We have seen how to describe the fusion of \( \sigma(0) \) and \( \sigma(z) \). What if now we instead take \( z \) close to 1 such that we can perform an operator product expansion of \( \sigma(z)\sigma(1) \). Taking this limit of Eq 39.16 it naively looks like both

\[
\lim_{z \to 1} G_+ \sim (1 - z)^{-1/8} \\
\lim_{z \to 1} G_- \sim (1 - z)^{-1/8}
\]

But examining this a bit more closely we realize we can construct the linear combinations

\[
\tilde{G}_+ = \frac{1}{\sqrt{2}} (G_+ + G_-) \\
\tilde{G}_- = \frac{1}{\sqrt{2}} (G_+ - G_-)
\]

where here we have inserted the prefactor of \( 1/\sqrt{2} \) such that the new basis \( \tilde{G}_\pm \) is orthonormal given that the old basis \( G_\pm \) was. With this new basis we now have the limits

\[
\lim_{z \to 1} \tilde{G}_+ \sim (1 - z)^{-1/8} \\
\lim_{z \to 1} \tilde{G}_- \sim (1 - z)^{-1/8} \left[ \sqrt{1 + \sqrt{1 - z}} - \sqrt{1 - \sqrt{1 - z}} \right] \\
\sim (1 - z)^{-1/8} (1 - z)^{1/2} \sim (1 - z)^{3/8}
\]

Thus we see that in this twiddle basis \( \tilde{G}_\pm \) we have in this limit that \( \tilde{G}_+ \) is the fusion of \( \sigma(z) \) and \( \sigma(1) \) to identity and \( \tilde{G}_- \) is the fusion to \( \psi \).

The transformation between the two bases \( G_\pm \) and \( \tilde{G}_\pm \) is precisely the \( F \)-matrix transformation.

\[
\begin{pmatrix}
\tilde{G}_+ \\
\tilde{G}_-
\end{pmatrix}
= \frac{1}{\sqrt{2}} \begin{pmatrix} 1 & 1 \\ 1 & -1 \end{pmatrix}
\begin{pmatrix}
G_+ \\
G_-
\end{pmatrix}
\]

which should look familiar to anyone who did the homework! (We also got the same result from writing the ising theory in terms of cabled Kauffman strings). Diagrammatically this transform is shown in Fig. 39.11.
39.6 More Comments on Moore-Read State with Many Quasiholes

Although we have presented this discussion about multiple fusion channels and braiding in terms of \( \sigma \) operators, the situation is extremely similar once we use quasihole operators \((\sigma(z)\mathcal{V}_\beta(z))\) and we put them in a wavefunction as in Eq. 39.13 but possibly with more quasihole operators. As we might expect just from looking at the fusion rules, the number of fusion channels (the number of Riemann sheets!) is \(2^{M/2-1}\) where \(M\) is the number of quasiholes, and the -1 arises because the overall fusion channel must be the identity. Further, the \(F\)-matrices and braiding properties all follow very much in a similar manner. The only slightly problematic piece is that we must continue to assume that the conformal blocks form an orthonormal basis — which is hard to prove, but appears to be true.

39.7 Generalizing to Other CFTs

The principles we used for building a quantum Hall state from the Ising CFT can be generalized to build quantum Hall states from other CFTs as well. The general principles are as follows:

1. Construct an electron field which gives a ground state which is single valued in the electron coordinates. This is done by starting with an abelian field from the CFT (one that does not have multiple fusion channels) and combining it with a chiral boson vertex operator. The filling fraction is determined entirely by the charge on the vertex operator.

2. Identify all of the possible quasiholes by looking at all the fields in the CFT and fusing them with a chiral boson vertex operator and enforcing the condition that the electron coordinates must not have branch cuts. The charge of the quasihole is determined by the charge on the vertex operator (and the charge on the electron vertex operator).

3. Some of the braiding properties can be determined immediately.
from the operator product expansion while others require more detailed information about the form of the CFT.

39.7.1 $\mathbb{Z}_3$ Parafermions (briefly)

As an example, let us consider the $\mathbb{Z}_3$ Parafermion CFT. Its primary fields and fusion rules are given by

<table>
<thead>
<tr>
<th></th>
<th>$h$</th>
<th>$\times$</th>
<th>$\psi_1$</th>
<th>$\psi_2$</th>
<th>$\sigma_1$</th>
<th>$\sigma_2$</th>
<th>$\epsilon$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\psi_1$</td>
<td>$2/3$</td>
<td>$\psi_1$</td>
<td>$\psi_2$</td>
<td>$\sigma_2 + \psi_1$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\psi_2$</td>
<td>$2/3$</td>
<td>$\psi_2$</td>
<td>$\epsilon$</td>
<td>$\sigma_2 + \psi_1$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\sigma_1$</td>
<td>$1/15$</td>
<td>$\sigma_1$</td>
<td>$\sigma_1$</td>
<td>$I + \epsilon$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\sigma_2$</td>
<td>$1/15$</td>
<td>$\sigma_2$</td>
<td>$\epsilon$</td>
<td>$\sigma_1 + \psi_2$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\epsilon$</td>
<td>$2/5$</td>
<td>$\epsilon$</td>
<td>$\sigma_2$</td>
<td>$\sigma_1 + \psi_2$</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

These fusion rules might look very complicated, but in fact they can be thought of as an abelian $\mathbb{Z}_3$ theory (with fields $I, \psi_1, \psi_2 = \bar{\psi}_1$) fused with a Fibonacci theory (with fields $I$ and $\tau$). We then have

$$\sigma_1 = \psi_2 \tau$$

$$\sigma_2 = \psi_1 \tau$$

$$\epsilon = \tau$$

and using the Fibonacci fusions $\tau \times \tau = I + \tau$ and the $\mathbb{Z}_3$ fusions $\psi_i \times \psi_j = \psi_{(i+j) \mod 3}$ with $\psi_0$ being the identity, we recover the full fusion table. Let us propose an electron field

$$\psi_e(z) = \psi_1(z)V_{\sqrt{m + \frac{2}{3}}}^m(z)$$

where $m$ is a nonnegative integer (even for bosons, odd for fermions). It is easy to check from the OPE that

$$\psi_e(z)\psi_e(w) \sim (z-w)^m \psi_2(z)V_{\sqrt{m + \frac{2}{3}}}^m(z)$$

The resulting wavefunction is then

$$\Psi = \langle \psi_e(z_1)\psi_e(z_2)\ldots\psi_e(z_N)\rangle Q$$

which is known as the Read-Rezayi $\mathbb{Z}_3$ parafermion wavefunction.

The filling fraction of the wavefunction is determined by the vertex operator and is given by

$$\nu = \frac{1}{m + \frac{2}{3}}$$

For the $m = 0$ case this is $\nu = 3/2$ bosons, while for the $m = 1$ case this is $\nu = 3/5$ fermions.

For the case of $m = 0$ it is easy to check that the wavefunction does not vanish when two particles come to the same point, nor does it vanish when three particles come to the same point, but it does vanish when...
four particles come to the same point. Thus the wavefunction is an exact (densest) zero energy ground state of a four particle delta function.

While there are 4-particle interactions for these systems for which wavefunctions are the exact ground state, it turns out that there are physically relevant cases where the Read-Rezayi $Z_3$ parafermion wavefunction is an extremely good trial wavefunction. For bosons interacting with a simple two body $\delta$-function potential at filling fraction $\nu = 3/2$, the $Z_3$ parafermion wavefunction is extremely good. For electrons interacting with simple coulomb interaction (in realistic quantum well samples), it turns out that the wavefunction is extremely good for $\nu = 2 + 2/5$, which we need to particle-hole conjugate in the partly filled Landau level to get a $\nu = 3/5$ wavefunction. (** add cites **)

To construct a quasihole we can try building a quasihole from any of the primary field operators. It turns out the one with the lowest charge is constructed from $\sigma_1$

$$\psi_{qh}(z) = \sigma_1(z)V_\beta(z)$$

Using the OPE we have

$$\sigma_1(w)\psi_1(z) \sim (z - w)^{-1/3}\epsilon(z)$$

We thus choose

$$\beta = \frac{p}{3\sqrt{m + \frac{2}{3}}}$$

with the smallest charge quasihole then being $p = 1$. With this choice, for a quasihole at position $w$ we generate a factor of

$$\prod_i (z - w)^{1/3}$$

meaning the charge of the quasihole is

$$e^* = e\nu/3$$
Exercise 39.1 Bose Vertex Operators

In lecture we needed the following identity
\[ \langle V_{\alpha_1}(z_1)V_{\alpha_2}(z_2)\ldots V_{\alpha_N}(z_N) \rangle = \prod_{i<j}(z_i - z_j)^{\alpha_i\alpha_j} \]  
(39.17)

where
\[ \sum_i \alpha_i = 0 \]  
(39.18)

where the vertex operators are defined by
\[ V_\alpha(z) = :e^{i\alpha \varphi(z)}: \]  
(39.19)

with \( \varphi \) a chiral bose field and colons meaning normal ordering.

(a) To get to this result, let us first show that for a bose operator \( a \), such that \([a, a^\dagger] = 1\), we have
\[ e^{\alpha a} e^{\beta a^\dagger} = e^{\beta a^\dagger} e^{\alpha a} e^{\alpha \beta} \]  
(39.20)

(b) Thus derive
\[ \langle V_{A_1}V_{A_2}\ldots V_{A_N} \rangle = e^{\sum_{i<j} \langle A_i A_j \rangle} \]  
(39.21)

where
\[ A_i = u_i a^\dagger + v_i a \]  
(39.22)

and
\[ V_{A_i} = :e^{A_i} := e^{u_i a^\dagger} e^{v_i a} \]  
(39.23)

with the colons meaning normal ordering (all daggers moved to the left).

(c) Show that Eq. (39.21) remains true for any operators \( A_i \) that are sums of different bose modes \( a_k \), i.e., if
\[ A_i = \sum_k [u_i(k) a_k^\dagger + v_i(k) a_k] \]  
(39.24)

Set \( A_i = i \alpha_i \varphi(z_i) \) such that \( V_{A_i} = V_{\alpha_i}(z_i) \). If \( \varphi \) is a free massless chiral bose field which can be written as the sum of fourier modes of bose operators such that
\[ \langle \varphi(z)\varphi(w) \rangle = -\ln(z - w) \]  
(39.25)

conclude that Eq. (39.17) holds.

Note: This result is not quite correct, as it fails to find the constraint Eq. (39.18) properly. The reason it fails is a subtlety which involves how one separates a bose field into two chiral components. (More detailed calculations that get this part right are given in the Big Yellow CFT book (P. Di Francesco, P. Mathieu, and D. Senechal) and in a different language in A. Tsvelik’s book.)

There is, however, a quick way to see that the constraint must be true. Note that the lagrangian of a massless chiral bose field is
\[ \mathcal{L} = \frac{1}{2\pi} \partial_x \varphi (\partial_x + v \partial_t) \varphi \]  
(39.26)
which clearly must be invariant under the global transformation $\phi \rightarrow \phi + b$.

(d) Show that the correlator Eq. 39.17 (with Eq. 39.19) cannot be invariant under this transformation unless Eq. 39.18 is satisfied, or unless the value of the correlator is zero.

**Exercise 39.2 Z$_4$ Quantum Hall State**

In this problem we intend to construct a quantum Hall state from the the Z$_4$ parafermion conformal field theory (Details of the CFT can be found in A. B. Zamolodchikov and V. A. Fateev, Soviet Physics JETP 62, 216 (1985), but we will not need too many of the details here).

The wavefunction we construct is known as the Z$_4$ Read-Rezayi wavefunction (N. Read and E. Rezayi, Phys. Rev. B 59, 8084 (1999)).

The Z$_4$ parafermion conformal field theory has 10 fields with corresponding conformal weights (scaling dimension)

<table>
<thead>
<tr>
<th>field</th>
<th>$\psi_1$</th>
<th>$\psi_2$</th>
<th>$\psi_3$</th>
<th>$\sigma_+$</th>
<th>$\sigma_-$</th>
<th>$\epsilon$</th>
<th>$\rho$</th>
<th>$\chi_+$</th>
<th>$\chi_-$</th>
</tr>
</thead>
<tbody>
<tr>
<td>weight $h$</td>
<td>0</td>
<td>$\frac{7}{4}$</td>
<td>1</td>
<td>$\frac{3}{4}$</td>
<td>$\frac{1}{10}$</td>
<td>$\frac{1}{10}$</td>
<td>$\frac{1}{12}$</td>
<td>$\frac{9}{10}$</td>
<td>$\frac{9}{10}$</td>
</tr>
</tbody>
</table>

and the fusion table is given by

<table>
<thead>
<tr>
<th>$\times$</th>
<th>1</th>
<th>$\psi_1$</th>
<th>$\psi_2$</th>
<th>$\psi_3$</th>
<th>$\sigma_+$</th>
<th>$\sigma_-$</th>
<th>$\epsilon$</th>
<th>$\rho$</th>
<th>$\chi_+$</th>
<th>$\chi_-$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>$\psi_1$</td>
<td>$\psi_2$</td>
<td>$\psi_3$</td>
<td>$\sigma_+$</td>
<td>$\sigma_-$</td>
<td>$\epsilon$</td>
<td>$\rho$</td>
<td>$\chi_+$</td>
<td>$\chi_-$</td>
</tr>
<tr>
<td>$\psi_1$</td>
<td>$\psi_1$</td>
<td>$\psi_2$</td>
<td>$\psi_3$</td>
<td>$\sigma_+$</td>
<td>$\sigma_-$</td>
<td>$\epsilon$</td>
<td>$\rho$</td>
<td>$\chi_+$</td>
<td>$\chi_-$</td>
<td></td>
</tr>
<tr>
<td>$\psi_2$</td>
<td>$\psi_1$</td>
<td>$\psi_2$</td>
<td>$\psi_3$</td>
<td>1</td>
<td>$\psi_1$</td>
<td>$\sigma_+$</td>
<td>$\sigma_-$</td>
<td>$\epsilon$</td>
<td>$\rho$</td>
<td>$\chi_+$</td>
</tr>
<tr>
<td>$\psi_3$</td>
<td>1</td>
<td>$\psi_1$</td>
<td>$\psi_2$</td>
<td>$\psi_3$</td>
<td>$\sigma_+$</td>
<td>$\sigma_-$</td>
<td>$\epsilon$</td>
<td>$\rho$</td>
<td>$\chi_+$</td>
<td>$\chi_-$</td>
</tr>
<tr>
<td>$\sigma_+$</td>
<td>$\sigma_+$</td>
<td>$\sigma_+$</td>
<td>$\sigma_+$</td>
<td>1</td>
<td>$\psi_1 + \rho$</td>
<td>$\psi_3 + \rho$</td>
<td>$\psi_1 + \psi_3 + \rho$</td>
<td>$\psi_2 + \psi_3 + \rho$</td>
<td>$\psi_1 + \rho$</td>
<td></td>
</tr>
<tr>
<td>$\sigma_-$</td>
<td>$\sigma_-$</td>
<td>$\sigma_-$</td>
<td>$\sigma_-$</td>
<td>$\psi_1 + \rho$</td>
<td>$\psi_3 + \rho$</td>
<td>$\psi_1 + \psi_3 + \rho$</td>
<td>$\psi_2 + \psi_3 + \rho$</td>
<td>$\psi_1 + \rho$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\epsilon$</td>
<td>$\epsilon$</td>
<td>$\epsilon$</td>
<td>$\epsilon$</td>
<td>$\epsilon$</td>
<td>$\psi_1 + \psi_3 + \rho$</td>
<td>$\psi_2 + \psi_3 + \rho$</td>
<td>$\psi_1 + \psi_3 + \rho$</td>
<td>$\psi_2 + \psi_3 + \rho$</td>
<td>$\psi_1 + \rho$</td>
<td></td>
</tr>
<tr>
<td>$\rho$</td>
<td>$\rho$</td>
<td>$\rho$</td>
<td>$\rho$</td>
<td>$\rho$</td>
<td>$\psi_1 + \psi_3 + \rho$</td>
<td>$\psi_2 + \psi_3 + \rho$</td>
<td>$\psi_1 + \psi_3 + \rho$</td>
<td>$\psi_2 + \psi_3 + \rho$</td>
<td>$\psi_1 + \rho$</td>
<td></td>
</tr>
<tr>
<td>$\chi_+$</td>
<td>$\chi_+$</td>
<td>$\chi_+$</td>
<td>$\chi_+$</td>
<td>$\chi_+$</td>
<td>$\chi_+$</td>
<td>$\chi_+$</td>
<td>$\chi_+$</td>
<td>$\chi_+$</td>
<td>$\chi_+$</td>
<td></td>
</tr>
<tr>
<td>$\chi_-$</td>
<td>$\chi_+$</td>
<td>$\chi_+$</td>
<td>$\chi_+$</td>
<td>$\chi_+$</td>
<td>$\chi_+$</td>
<td>$\chi_+$</td>
<td>$\chi_+$</td>
<td>$\chi_+$</td>
<td>$\chi_+$</td>
<td></td>
</tr>
</tbody>
</table>

If I have not made any mistake in typing this table, the fusion rules should be associative

\[(a \times b) \times c = a \times (b \times c)\]  

(39.27)

Note of interest: These fusion rules may look mysterious, but in fact they are very closely related to the fusion rules of SU(2) appropriately truncated (i.e., this is the SU(2)$_2$ WZW model). We can write each field as a young tableau with no more than 2 (for SU(2)) columns and no more than 4 − 1 = 3 rows

<table>
<thead>
<tr>
<th>field</th>
<th>$\psi_1$</th>
<th>$\psi_2$</th>
<th>$\psi_3$</th>
<th>$\sigma_+$</th>
<th>$\sigma_-$</th>
<th>$\epsilon$</th>
<th>$\rho$</th>
<th>$\chi_+$</th>
<th>$\chi_-$</th>
</tr>
</thead>
<tbody>
<tr>
<td>tableau</td>
<td>empty</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The fusion rules are just a slight modification of the usual young tableau
Exercises

manipulations for $SU(2)$ where columns are removed if they have 4 boxes. (See the big yellow book for details).

Using the techniques discussed in lecture:

(a) Use the operator product expansion (dimension counting) to find the singularity as two $\psi_1$ fields come close together. I.e., find the exponent $\alpha$ in the relation

$$\lim_{z' \to z} \psi_1(z')\psi_1(z) \sim (z' - z)\alpha \psi_1(z) \quad (39.28)$$

(b) Construct all possible “electron” fields by making a product of the $\psi_1$ field and a chiral boson vertex operator of the form

$$\psi_e(z) = \psi_1(z)e^{i\beta\phi(z)} \quad (39.29)$$

that give a single-valued and nonsingular wavefunction for the electron. (See Eq. 39.17, but ignore the sum condition Eq. 39.18). I.e., find all acceptable values of $\beta$. Consider both the case where the “electron” is a boson or a fermion. What filling fractions do these correspond to? (There are multiple allowable solutions for both bosons and fermions). Consider among the bosonic solution, the one solution of the highest density. The ground state wavefunction in this case is the highest density zero energy state of a 5-point delta function interaction. Show that the wavefunction does not vanish when 4 particles come to the same point, but does indeed vanish as 5 particles come to the same point.

(c) Given a choice of the electron field, construct all possible quasihole operators from all fields $\varphi$ in the above table

$$\varphi_{qh}(w) = \varphi(w)e^{i\kappa\phi(w)} \quad (39.30)$$

For each case, fix the values of $\kappa$ by insisting that the wavefunction remain single-valued in the electron coordinates. Determine the quasihole with the lowest possible (nonzero) electric charge. What is this charge?

(d) Two such quasiholes can fuse together in two possible fusion channels. What is the monodromy in each of these channels. I.e., what phase is accumulated when the two quasiholes are transported around each other (assuming the Berry matrix is zero – which is a statement about wavefunctions being properly orthonormal – which we usually assume is true).

(e) Draw a Bratteli diagram (a tree) describing the possible fusion channels for many of these elementary particles. Label the number of paths in the diagram for up to 10 quasiholes. If there are 8 quasiparticles and the number of electrons is divisible by 4, what is the degeneracy of the ground state? If there are 4 quasiparticles and the number of electrons is $4m + 2$ what is the degeneracy of the ground state?

(f) Construct a 5 by 5 transfer matrix and show how to calculate the ground state degeneracy in the presence of any number of quasiholes. Finding the largest eigenvalue of this matrix allows you to calculate the “quantum dimension” $d$ which is the scaling

$$\text{Degeneracy} \sim d^{|\text{Number of Quasiholes}|} \quad (39.31)$$

in the limit of large number of quasiholes. While diagonalizing a 5 by 5 matrix seems horrid, this one can be solved in several easy ways (look for a trick or a nice factorization of the characteristic polynomial).
(g) Consider instead constructing a wavefunction from the $\psi_2$ field

$$\psi_4(z) = \psi_2(z)e^{i\beta\phi(z)} \quad (39.32)$$

What filling fraction does this correspond to (for bosons or fermions)? In the highest density case, what are the properties of this wavefunction (how does it vanish as how many many electrons come to the same point).
Appendix: Resources for TQFTs

Working out the details of a TQFT is an often tedious task and except in the simplest cases, one does not want to go through the pain of doing this. At the end of this chapter we list a number of resources for finding details of many TQFTs.

Perhaps the most useful single resource I have found for obtaining data about TQFTs is a computer program called Kac written by A. N. Schellekens. The complicated part of the algorithm is described by Fuchs et al. [1996]. More details are given on the project webpage.

The program can be downloaded from the webpage

https://www.nikhef.nl/~t58/Site/Kac.html

While the program has many capabilities (and I encourage you to RTFM\(^1\)), it is probably useful to give here an annotated example of how it works. Note that the program uses Dynkin diagram (Cartan) notation for describing Lie algebras. The correspondence is given by

\[
\begin{align*}
A_r &= \mathfrak{su}(r+1) \\
B_r &= \mathfrak{so}(2r+1) \\
C_r &= \mathfrak{sp}(2r) \\
D_r &= \mathfrak{so}(2r)
\end{align*}
\] (40.1)

One can also use the \(E_6, E_7, E_8, F_4\) and \(G_2\) Lie algebras.

Here we present some annotated sessions with Kac.

**MY LINUX BOX**

Kac (on MY LINUX BOX), version 8.05468, compiled on Sep 1 2016, at 16:27:29

Started Sun 30 Aug 05:19:53 BST 2020

Non-interactive mode; Assuming default answer: OK

> tensor

The tensor command tells the program that we might be tensoring together multiple theories.

> g a 1 2

This inputs the group (\(g\) for group) with the Cartan notation \(a\ 1\), or \(A_1\) which is \(\mathfrak{su}(2)\) as given by the correspondence Eq. [40.1] above, and the 2 indicates level 2. So we are asking it to compute information about \(SU(2)_2\).
> display
CFT \{A1:2\}; 3 primaries (2 simple currents)

<table>
<thead>
<tr>
<th>Lbl</th>
<th>Comb.</th>
<th>Weights</th>
<th>Wts.</th>
<th>F.l.</th>
<th>F.m.</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>{0}</td>
<td>0.0000000</td>
<td>0</td>
<td>-</td>
<td>1</td>
</tr>
<tr>
<td>1</td>
<td>{1}</td>
<td>0.5000000</td>
<td>1/2</td>
<td>-</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>{2}</td>
<td>0.1875000</td>
<td>3/16</td>
<td>-</td>
<td>1</td>
</tr>
</tbody>
</table>

The fields are numbered 0, 1, 2, and we see their corresponding weights $h = 0, 1/2, 3/16$. The simple currents are always listed first. Recall that twist factors are given by $\theta = e^{2\pi i h}$. Note also that the weights are only correct modulo one. We can then ask for quantities like the fusion rules, or the $S$-matrix, the Frobenius-Schur indicator, or the central charge

> fusion

\[
\begin{align*}
(0) \times (0) &= (0) \\
(0) \times (1) &= (1) \\
(0) \times (2) &= (2) \\
(1) \times (1) &= (0) \\
(1) \times (2) &= (2) \\
(2) \times (2) &= (0) + (1)
\end{align*}
\]

> S

\[
\begin{align*}
S(0,0) &= 0.50000000 \\
S(0,1) &= 0.50000000 \\
S(0,2) &= 0.70710678 \\
S(1,1) &= 0.50000000 \\
S(1,2) &= -0.70710678 \\
S(2,2) &= 0.00000000
\end{align*}
\]

> Get Schur 2

-1

> Browse Central

Central charge 1.500000000000000

If we had wanted to look at the opposite chirality theory, we use $h$ rather than $g$. To wipe the memory of the program and return to tensor mode we use `reset tensor`. So for example, we have
> reset tensor
> h a 1 2
> display
CFT {A1:2}; 3 primaries (2 simple currents)

<table>
<thead>
<tr>
<th>Lbl</th>
<th>Comb.</th>
<th>Weights</th>
<th>Wts.</th>
<th>F.l.</th>
<th>F.m.</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>{}</td>
<td>0.0000000 (mod 1)</td>
<td>0</td>
<td>-</td>
<td>1</td>
</tr>
<tr>
<td>1</td>
<td>{1}</td>
<td>0.5000000 (mod 1)</td>
<td>1/2 (mod 1)</td>
<td>-</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>{2}</td>
<td>0.8125000 (mod 1)</td>
<td>13/16 (mod 1)</td>
<td>-</td>
<td>1</td>
</tr>
</tbody>
</table>

Note that the weight of the 2 field is 13/16 = −3/16 mod 1 so this is the opposite chirality version of $SU(2)_2$ which we write as $SU(2)_2$.

The program can also handle $U(1)$ Chern-Simons theory, and accepts a parameter for so-called “radius” of the boson (which substitutes for the level of the Chern-Simons theory). Since there is some disagreement in the literature as to how you label the level of a $U(1)$ Chern-Simons theory, and as to how you label the radius, it is worth stating explicitly that in the convention used by this program, the theory with radius $R$ has $R$ different fields. In the convention we use in section 20.4.2 we have $U(1)_{N/2}$ corresponding to radius $N$ for $N$ even. We produce these theories using the code g u followed by the radius as follows.

> reset tensor
> g u 4
> display
CFT {U4:0}; 4 primaries (4 simple currents)

<table>
<thead>
<tr>
<th>Lbl</th>
<th>Comb.</th>
<th>Weights</th>
<th>Wts.</th>
<th>F.l.</th>
<th>F.m.</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>{}</td>
<td>0.0000000</td>
<td>0</td>
<td>-</td>
<td>1</td>
</tr>
<tr>
<td>1</td>
<td>{1}</td>
<td>0.1250000</td>
<td>1/8</td>
<td>-</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>{2}</td>
<td>0.5000000</td>
<td>1/2</td>
<td>-</td>
<td>1</td>
</tr>
<tr>
<td>3</td>
<td>{3}</td>
<td>0.1250000</td>
<td>1/8</td>
<td>-</td>
<td>1</td>
</tr>
</tbody>
</table>

The program can handle condensation, as well as splitting. Let us consider the example used in section 23.4 of $SU(2)_4$. We first produce the $SU(2)_4$ theory

> reset tensor
> g a 1 4
> display
CFT {A1:4}; 5 primaries (2 simple currents)

<table>
<thead>
<tr>
<th>Lbl</th>
<th>Comb.</th>
<th>Weights</th>
<th>Wts.</th>
<th>F.l.</th>
<th>F.m.</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>{}</td>
<td>0.0000000</td>
<td>0</td>
<td>-</td>
<td>1</td>
</tr>
<tr>
<td>1</td>
<td>{1}</td>
<td>1.0000000</td>
<td>1</td>
<td>-</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>{2}</td>
<td>0.1250000</td>
<td>1/8</td>
<td>-</td>
<td>1</td>
</tr>
<tr>
<td>3</td>
<td>{3}</td>
<td>0.6250000</td>
<td>5/8</td>
<td>-</td>
<td>1</td>
</tr>
<tr>
<td>4</td>
<td>{4}</td>
<td>0.3333333</td>
<td>1/3</td>
<td>-</td>
<td>1</td>
</tr>
</tbody>
</table>

Note that one of the simple currents is a boson (integer weight). To condense it we issue the command current and the name of the field
we want to condense.

> current 1
> display

CFT \{A1:4\}; 3 primaries

<table>
<thead>
<tr>
<th>Lbl</th>
<th>Comb.</th>
<th>Weights</th>
<th>Wts.</th>
<th>F.l.</th>
<th>F.m.</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>{0}</td>
<td>0.000000</td>
<td>0</td>
<td>-</td>
<td>1</td>
</tr>
<tr>
<td>1</td>
<td>{4}</td>
<td>0.333333</td>
<td>1/3</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>{4}</td>
<td>0.333333</td>
<td>1/3</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

> fusion

\(0 \times 0 = (0)\)
\(0 \times 1 = (1)\)
\(0 \times 2 = (2)\)
\(1 \times 1 = (2)\)
\(1 \times 2 = (0)\)
\(2 \times 2 = (1)\)

Which correctly splits the 4-particle as we discussed in section \[23.4\].

To generate product theories, we just input several theories in a row. For example, to look at a product theory, \(SU(2)_2 \times SU(2)_1 \times SU(2)_1\), we write

> reset tensor
> g a 1 2
> h a 1 1
> h a 1 1
> display

CFT \{A1:2_A1:1_A1:1\}; 12 primaries (8 simple currents)

<table>
<thead>
<tr>
<th>Lbl</th>
<th>Comb.</th>
<th>Weights</th>
<th>Wts.</th>
<th>F.l.</th>
<th>F.m.</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>{0,0,0}</td>
<td>0.0000000</td>
<td>(mod 1)</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>1</td>
<td>{0,0,1}</td>
<td>0.7500000</td>
<td>(mod 1)</td>
<td>3/4</td>
<td>(mod 1)</td>
</tr>
<tr>
<td>2</td>
<td>{0,1,0}</td>
<td>0.7500000</td>
<td>(mod 1)</td>
<td>3/4</td>
<td>(mod 1)</td>
</tr>
<tr>
<td>3</td>
<td>{0,1,1}</td>
<td>0.5000000</td>
<td>(mod 1)</td>
<td>1/2</td>
<td>(mod 1)</td>
</tr>
<tr>
<td>4</td>
<td>{1,0,0}</td>
<td>0.5000000</td>
<td>(mod 1)</td>
<td>1/2</td>
<td>(mod 1)</td>
</tr>
<tr>
<td>5</td>
<td>{1,0,1}</td>
<td>0.2500000</td>
<td>(mod 1)</td>
<td>1/4</td>
<td>(mod 1)</td>
</tr>
<tr>
<td>6</td>
<td>{1,1,0}</td>
<td>0.2500000</td>
<td>(mod 1)</td>
<td>1/4</td>
<td>(mod 1)</td>
</tr>
<tr>
<td>7</td>
<td>{1,1,1}</td>
<td>0.0000000</td>
<td>(mod 1)</td>
<td>0</td>
<td>(mod 1)</td>
</tr>
<tr>
<td>8</td>
<td>{2,0,0}</td>
<td>0.1875000</td>
<td>(mod 1)</td>
<td>3/16</td>
<td>(mod 1)</td>
</tr>
<tr>
<td>9</td>
<td>{2,0,1}</td>
<td>0.9375000</td>
<td>(mod 1)</td>
<td>15/16</td>
<td>(mod 1)</td>
</tr>
<tr>
<td>10</td>
<td>{2,1,0}</td>
<td>0.9375000</td>
<td>(mod 1)</td>
<td>15/16</td>
<td>(mod 1)</td>
</tr>
<tr>
<td>11</td>
<td>{2,1,1}</td>
<td>0.6875000</td>
<td>(mod 1)</td>
<td>11/16</td>
<td>(mod 1)</td>
</tr>
</tbody>
</table>

Since \(SU(2)_2\) has 3 fields, and each \(SU(2)_1\) has 2 fields, the product
of these three theories has 12 fields. The second column of the output shows how each field is constructed from the constituent factors. For example, the output field labeled 9 in the far left column comes from the 2 field of \( SU(2) \), the 0 field from the first \( SU(2)_1 \) and the 1 field from the second \( SU(2)_1 \).

Let us now construct the coset \( SU(2)_2/(SU(2)_1 \times SU(2)_1) \). Recall from section 23.6 that one can construct this coset by starting with \( SU(2)_2 \times SU(2)_1 \times SU(2)_1 \) and condensing all possible simple current bosons. Notice in the above output that there are 8 simple currents, and the one labeled 7 or \( \{1,1,1\} \) is a boson. We thus issue the command

\[
\text{current 1 1 1} \\
\text{display}
\]

\[
\text{CFT \{A1:2,A1:1,A1:1\}; 3 primaries (2 simple currents)}
\]

\begin{tabular}{llll}
Lbl & Comb. & Weights & F.L. & F.m. \\
0 & \{0,0,0\} & 0.0000000 & (mod 1) & 0 - 1 \\
1 & \{0,1,1\} & 0.5000000 & (mod 1) & 1/2 (mod 1) - 1 \\
2 & \{2,0,1\} & 0.9375000 & (mod 1) & 15/16 (mod 1) - 1 \\
\end{tabular}

Giving us the result that this coset is actually Ising.

**Further Reading**

Note that many of the following references give only the so-called “modular data” for TQFTs — meaning the \( S \)-matrices (which imply the fusion rules via the Verlinde formula, Eq. 17.13) and the twist factors \( \theta_a \). However, it has recently been established that there can be cases where more than one modular TQFT can share the same modular data (Mignard and Schauenburg [2017]). However the simplest such case known where the modular data does not uniquely define the TQFT has 49 different particle types and for all simple TQFTs the modular data is, at least in principle, full information.

- A useful reference on conformal field theory, including WZW theories (which give you the content of the corresponding Chern-Simons theory) is given by Di Francesco et al. [1997].
- Many details of the simplest few modular tensor categories on the periodic table are given by Rowell et al. [2009]; A discussion for fermionic models is given by Bruillard et al. [2017, 2020].
- Some nice data for some simple categories is given by Bonderson [2007]. This includes, for example, the \( F \)-matrices for \( SU(2)_2 \) and a number of other simple theories.
- \( F \)-matrices for many more complicated theories are given by Ardonne and Slingerland [2010].
- Online databases of vertex algebras, modular categories, fusion rings etc are given at
  
  - https://www.math.ksu.edu/~gerald/voas/
  - http://www.thphys.nuim.ie/AnyonWiki/index.php/Main_Page

\[\text{2 See also Bonderson et al. 2019 and Delaney and Tran 2018 for discussion of what additional data might be added to make the TQFT unique}\]
Some Mathematical Basics

Many undergraduates (and even many graduates) do not get any proper education in advanced mathematics. As such I am including a very short exposition of most of what you need to know in order to read this book. For much of the book, you won’t even need to know this much! If you have even a little background in mathematics you will probably know most of this already.

41.1 Manifolds

We sometimes write $\mathbb{R}$ to denote the real line, i.e., it is a space where a point is indexed by a real number $x$. We can write $\mathbb{R}^n$ to denote $n$-dimensional (real) space — a space where a point is indexed by $n$-real numbers $(x_1, \ldots, x_n)$. Sometimes people call these spaces “Euclidean” space.

Definition 41.1 A Manifold is a space that locally looks like a Euclidean space.

If a manifold is bounded, contains all its limit points, and has no boundary we call it closed.

41.1.1 Some Simple Examples: Euclidean Spaces and Spheres

- $\mathbb{R}^n$ is obviously a manifold (it is not bounded, so therefore not closed).
- The circle $S^1$, also known as a 1-sphere (hence the notation, the index 1 meaning it is a 1-dimensional object) is defined as as all points in a plane equidistant from a central point. Locally this looks like a line since position is indexed by a single variable (the “curvature” of the circle is not important locally). Globally, one discovers that the circle is not the same as a real line, as position is periodic (if you walk far enough in one direction you come back to where you start). We sometimes define a circle as a real number from 0 to $2\pi$ which specifies the angle around the circle.
- The 2-sphere $S^2$ is what we usually call (the surface of) a sphere in our regular life. We can define this similarly as all points in $\mathbb{R}^3$ equidistant from a central point.
- One can generally define the $n$-sphere, $S^n$, as points equidistant from a central point in $\mathbb{R}^{n+1}$.
Often when we discuss a manifold, we will be interested in its topological properties only. In other words, we will not care if a circle is dent ed as shown in Fig. 41.1; it is still topologically $S^1$. Mathematicians say that two objects that can be smoothly deformed into each other are homeomorphic, although we will not use this language often.

It is sometimes convenient to view the circle $S^1$ as being just the real line $\mathbb{R}$ with a single point added “at infinity” — think about joining up $+\infty$ with $-\infty$ to make a circle. We can do the same thing with the sphere $S^2$ and $\mathbb{R}^2$ — this is like taking a big flat sheet and pulling the boundary together to a point to make it into a bag and closing up the top (which gives a sphere $S^2$). Obviously the idea generalizes: $S^3$ is the same as $\mathbb{R}^3$ “compactified” with a point at infinity, and so forth.

### Orientability

We say a manifold is orientable if we can consistently define a vector normal to the manifold at all points. Another way of defining orientability (that does not rely on embedding the manifold in a higher dimension) is that we should be able to consistently define an orientation of the coordinate axes at all points on the manifold. Throughout this book we will almost always assume that all manifolds are orientable.

An example of a nonorientable manifold is the Möbius strip shown in Fig. 41.2. If we smoothly move the coordinate axes around the strip, when we come back to the same point, the upward pointing normal will have transformed into a downward facing normal.

There is a very simple classification of orientable closed (bounded and without boundary) two dimensional manifolds by the number of “holes” which is known as its “genus”, $g$. A sphere ($g = 0$) has no holes, a torus ($g = 1$) has one hole, a two handled torus ($g = 2$) has two holes, and so forth. See Fig. 41.3.

### 41.1.2 Unions of Manifolds $\mathcal{M}_1 \cup \mathcal{M}_2$

We can take a “disjoint” union of manifolds, using the notation $\cup$. For example, $S^1 \cup S^2$ is two circles (not connected in any way). If we think of this as being a single manifold, it is a manifold made of two disjoint pieces (or a disconnected manifold). Locally it still looks like a Euclidean space.

### 41.1.3 Products of Manifolds: $\mathcal{M}_3 = \mathcal{M}_1 \times \mathcal{M}_2$

One can take the product of two manifolds, or “cross” them together, using the notation $\times$. We write $\mathcal{M}_3 = \mathcal{M}_1 \times \mathcal{M}_2$. This means that a point in $\mathcal{M}_3$ is given by one point in $\mathcal{M}_1$ and one point in $\mathcal{M}_2$. This multiplication is often called the direct or Cartesian product.

- $\mathbb{R}^2 = \mathbb{R}^1 \times \mathbb{R}^1$. Here, a point in $\mathbb{R}^1$ is specified by a single real number. Crossing two of these together, a point in $\mathbb{R}^2$ is specified...
by two real numbers (one in the first \( \mathbb{R}^1 \) and one in the second \( \mathbb{R}^1 \)).

- \( T^2 = S^1 \times S^1 \). The 2-torus \( T^2 \), or surface of a doughnut\(^1\) is the product of two circles. To see this note that a point on a torus is specified by two angles, and the torus is periodic in both directions. Similarly we can build higher dimensional tori (tori is the plural of torus) by crossing \( S^1 \)'s together any number of times.

1. Alternatively spelled “donut” if you are from the states and you like coffee.

### 41.1.4 Manifolds with Boundary:

One can also have manifolds with boundary. A boundary of a manifold locally looks like an \( n \)-dimensional half-Euclidean space. The interior of a manifold with boundary looks like a Euclidean space, and near the boundary it looks like a half-space, or space with boundary. For example, a half-plane is a 2-manifold with boundary. An example is useful:

- The \( n \)-dimensional ball, denoted \( B^n \) is defined as the set of points in \( n \) dimensional space such that the distance to a central point is less than or equal to some fixed radius \( r \). Note: Often the ball is called a disk and is denoted by \( D^n \) (so \( D^n = B^n \)). The nomenclature makes good sense in two dimensions, where what we usually call a disk is \( D^2 \). The one-dimensional ball is just an interval (one-dimensional segment) which is sometimes denoted \( I = D^1 = B^1 \).

Note that a boundary of a manifold may have disconnected parts. For example, the boundary of an interval (segment) in 1-dimension \( I = B^1 \) is two disconnected points at its two ends\(^2\).

One can take cartesian products of manifolds with boundaries too. For example, consider the interval (or 1-ball) \( I = B^1 \) which we can think of as all the points on a line with \( |x| \leq 1 \). The cartesian product \( I \times I \) is described by two coordinates \((x, y)\) where \( |x| \leq 1 \) and \( |y| \leq 1 \). This is a square including its interior. However, in topology we are only ever concerned with topological properties, and a square-with-interior can be continuously deformed into a circle-with-interior, or a 2-ball (2-disc), \( B^2 \).

\(^2\)In the notation of Section 41.1.3 below, \( \partial I = \text{pt} \cup \text{pt} \) where \( \text{pt} \) means a point and here \( \cup \) means the union of the two objects as described above in 41.1.2.

- The same reasoning gives us the general topological law \( B^n \times B^m = B^{n+m} \).
- The cylinder (hollow tube) is expressed as \( S^1 \times I \) (two coordinates, one periodic, one bounded on both sides).
- The solid donut is expressed as \( D^2 \times S^1 \) (\( = B^2 \times S^1 \)), a 2-disk crossed with a circle.
41.1.5 **Boundaries of Manifolds: \( \mathcal{M}_1 = \partial \mathcal{M}_2 \).**

The notation for boundary is \( \partial \), so if \( \mathcal{M}_1 \) is the boundary of \( \mathcal{M}_2 \) we write \( \mathcal{M}_1 = \partial \mathcal{M}_2 \). The boundary \( \partial \mathcal{M} \) has dimension one less than that of \( \mathcal{M} \).

- The boundary of \( D^2 \), the 2-dimensional disk is the one dimensional circle \( S^1 \).
- More generally, the boundary of \( B^n \) (also written as \( D^n \)) is \( S^{n-1} \).

It is an interesting topological principle that the boundary of a manifold is always a manifold without boundary. Or equivalently, the boundary of a boundary is the empty set. We sometimes write \( \partial^2 = 0 \) or \( \partial (\partial \mathcal{M}) = \emptyset \) where \( \emptyset \) means the empty set.

- The boundary of the 3-dimensional ball \( B^3 \) is the sphere \( S^2 \). The sphere \( S^2 \) is a 2-manifold without boundary.

The operation of taking a boundary obeys the Leibnitz rule analogous to taking derivatives

\[
\partial(\mathcal{M}_1 \times \mathcal{M}_2) = (\partial \mathcal{M}_1) \times \mathcal{M}_2 \cup \mathcal{M}_1 \times (\partial \mathcal{M}_2)
\]

Let’s see some examples of this:

- Consider the cylinder \( S^1 \times I \). Using the above formula with find its boundary

\[
\partial(S^1 \times I) = (\partial S^1) \times I \cup S^1 \times \partial I = S^1 \cup S^1
\]

To see how we get the final result here, start by examining the first term, \((\partial S^1) \times I\). Here, \( S^1 \) has no boundary so \( \partial S^1 = \emptyset \) and therefore everything before the \( \cup \) symbol is just the empty set. In the second term the boundary of the interval is just two points \( \partial I = \text{pt} \cup \text{pt} \). Thus the second term gives the final result \( S^1 \cup S^1 \), the union of two circles.

- Consider writing the disk (topologically) as the product of two intervals \( B^2 = I \times I \). It is best to think of this cartesian product as forming a filled-in square. Using the above formula we get

\[
\partial B^2 = \partial(I \times I) = (\text{pt} \cup \text{pt}) \times I \cup I \times (\text{pt} \cup \text{pt}) = (I \cup I) \cup (I \cup I) = \text{top} \cup \text{bottom} \cup \text{left} \cup \text{right} = \text{square (edges only)} = S^1
\]

The formula gives the union of four segments denoting the edges of the square.
41.2 Groups

A group $G$ is a set of elements $g \in G$ along with an operation that we think of as multiplication. The set must be closed under this multiplication. So if $g_1, g_2 \in G$ then $g_3 \in G$ where

$$g_3 = g_1 g_2$$

where by writing $g_1 g_2$ we mean multiply $g_1$ by $g_2$. Note: $g_1 g_2$ is not necessarily the same as $g_2 g_1$. If the group is always commutative (i.e., if $g_1 g_2 = g_2 g_1$ for all $g_1, g_2 \in G$), then we call the group abelian\(^3\). If there are at least some elements in the group where $g_1 g_2 \neq g_2 g_1$ then the group is called nonabelian\(^4\).

A group must always be associative

$$g_1 (g_2 g_3) = (g_1 g_2) g_3 = g_1 g_2 g_3$$

Within the group there must exist an identity element which is sometimes\(^5\) called $e$ or $I$ or 0 or 1. The identity element satisfies

$$ge = eg = g$$

for all elements $g \in G$. Each element of the group must also have an inverse which we write as $g^{-1}$ with the property that

$$gg^{-1} = g^{-1}g = e$$

We often write $|G|$ to mean the number of elements in the group $G$.

41.2.1 Some Examples of Groups

- The group of integers $\mathbb{Z}$ with the operation being addition. The identity element is 0. This group is abelian.
- The group $\{1, -1\}$ with the operation being the usual multiplication. This is also called the group $\mathbb{Z}_2$. The identity element is 1. We could have also written this group as $\{0, 1\}$ with the operation being the usual addition modulo 2, where here the identity is 0. This group is abelian.
- The group $\mathbb{Z}_N$ which is the set of complex numbers $e^{2\pi ip/N}$ with $p$ an integer (which can be chosen between 1 and $N$ inclusive) and the operation being multiplication. This is equivalent to the set of integers modulo $N$ with the operation being addition. This group is abelian.
- The group of permutations of $N$ elements, which we write as $S_N$ (known as the permutation group, or symmetric group). This group is nonabelian. There are $N!$ elements in the group. Think of the elements of the group as being a one-to-one mapping from the set of the first $N$ integers into itself.

\(^3\)Named after Abel, the Norwegian mathematician who studied such groups in the early 1800s despite living in poverty and perishing at the young age of 26 from tuberculosis. The word “abelian” is usually not capitalized due to its ubiquitous use. There are a few similar words in English which are not capitalized despite being named after people, such as “galvanic”.

\(^4\)Apparently named after someone named Nonabel.

\(^5\)It may seem inconvenient that the identity has several names. However, it is sometimes convenient. If we are thinking of the group of integers and the operation of addition, we want to use 0 as the identity. If we are thinking about the group $\{1, -1\}$ with the operation of usual multiplication, then it is convenient to write the identity as 1. For more abstract groups, $e$ or $I$ is often most natural.
• The simplest nonabelian group is $S_3$. In $S_3$, one of the elements is

$$X = \begin{cases} 
1 & \rightarrow 2 \\
2 & \rightarrow 1 \\
3 & \rightarrow 3 
\end{cases}$$

Another element is

$$R = \begin{cases} 
1 & \rightarrow 2 \\
2 & \rightarrow 3 \\
3 & \rightarrow 1 
\end{cases}$$

where $X$ stands for exchange (exchanges 1 and 2) and $R$ stands for rotate. The multiplication operation $XR$ is meant to mean, do $R$ first, then do $X$ (you should be careful to make sure your convention of ordering is correct. Here we choose a convention that we do the operation written furthest right first. You can choose either convention, but then you must stick to it! You will see both orderings in the literature!) So, if we start with the element 1, when we do $R$ the element 1 gets moved to 2. Then when we do $X$ the element 2 gets moved to 1. So in the product $XR$ we have 1 getting moved back to position 1. In the end we have

$$XR = \begin{cases} 
1 & \rightarrow 1 \\
2 & \rightarrow 3 \\
2 & \rightarrow 1 
\end{cases}$$

Note that if we multiply the elements in the opposite order we get a different result (hence this group is nonabelian)

$$RX = \begin{cases} 
1 & \rightarrow 3 \\
2 & \rightarrow 2 \\
1 & \rightarrow 2 
\end{cases}$$

It is easy to check that

$$X^2 = R^3 = e \quad (41.1)$$

and further we have

$$XR = R^2 X \quad (41.2)$$

There are a total of 6=3! elements in the group which we can list as $e, R, R^2, X, XR, XR^2$. All other products can be reduced to one of these 6 possibilities using Eqs. (41.1) and (41.2).

**Product of Groups**

Two groups $G$ and $H$ can be multiplied to form $G \times H$ in an obvious way. An element of $G \times H$ is a pair $(g, h)$ with $g \in G$ and $h \in H$. The multiplication in $G \times H$ is given by $(g_1, h_1)(g_2, h_2) = (g_1g_2, h_1h_2)$. 
All Abelian Groups

The so-called **fundamental theorem of finite abelian groups**\(^6\) tells us that any abelian groups can be written as

\[ \mathbb{Z}_{N_1} \times \mathbb{Z}_{N_2} \times \ldots \times \mathbb{Z}_{N_p} \]

for some number of factors \(p\).

41.2.2 More Features of Groups

A **subgroup** is a subset of elements of a group which themselves form a group. For example, the integers under addition form a group. The even integers under addition are a subgroup of the integers under addition.

The **centralizer** of an element \(g \in G\) often written as \(Z(g)\) is the set of all elements of the group \(G\) that commute with \(g\). I.e., \(h \in Z(g)\) iff \(hg = gh\). Note that this set forms a subgroup (proof is easy!). For an abelian group \(G\) the centralizer of any element is the entire group \(G\).

A **conjugacy class** of an element \(g \in G\) is defined as the set of elements \(g' \in G\) such that \(g' = hgh^{-1}\) for some element \(h \in G\). We sometimes write \(C_g\) for the conjugacy class of \(g\), and \(|C_g|\) is the number of elements in the conjugacy class.

A useful result of group theory is that the number of elements in the conjugacy class of an element times the centralizer of the element gives the total number of elements in the group\(^7\)

\[ |Z(g)| \cdot |C_g| = |G| \quad (41.3) \]

**Example:** \(S_3\)  Above we listed some of the properties of the group \(S_3\). \(S_3\) has several subgroups:

- The group containing the identity element \(e\) alone
- The group containing \(\{e, X\}\)
- The group containing \(\{e, R, R^2\}\)
- The group \(S_3\) itself (which is not a so-called “proper” subgroup)

The centralizer is just the identity element \(Z(S_3) = e\), since it is the only element of the group \(S_3\) that commutes with all elements of the group. The group has three conjugacy classes

- The identity element \(e\)
- The rotations \(\{R, R^2\}\)
- The reflections \(\{X, XR, XR^2\}\)

We can check that conjugating any element in any class gives another element within the same class. For example, consider the element \(X\) and conjugate it with the element \(R\). We have \(RXR^{-1} = XR\) which is in the same conjugacy class as \(X\).
Some Mathematical Basics

41.2.3 Lie Groups and Lie Algebras

A **Lie group** is a group which is also a manifold. Roughly, a group with a continuous (rather than discrete) set of elements. Examples include:

- The group of invertible $n \times n$ complex matrices. We call this group $GL(n, \mathbb{C})$. Here $GL$ stands for “general linear”. The identity is the usual identity matrix. By definition all elements of the group are invertible.
- The group of invertible $n \times n$ real matrices. We call this group $GL(n, \mathbb{R})$.
- The group, $SU(2)$, the set of 2 by 2 unitary matrices with unit determinant. In this case the fact that this is also a manifold can be made particularly obvious. We can write all $SU(2)$ matrices as
  \[
  \left( \begin{array}{cc}
  x_1 + ix_2 & -x_3 + ix_4 \\
  x_3 + ix_4 & x_1 - ix_2
  \end{array} \right)
  \]
  with all $x_j$ any real numbers with the constraint that $x_1^2 + x_2^2 + x_3^2 + x_4^2 = 1$. Obviously the set of four coordinates $(x_1, x_2, x_3, x_4)$ with the unit magnitude constraint describes the manifold $S^3$.
- $SU(N)$, the group of unitary $N$ by $N$ matrices of determinant one is a Lie group
- $SO(N)$, the group of real rotation matrices in $N$ dimensions is a Lie group.
- The vector space $\mathbb{R}^n$ with the operation being addition of vectors, is a Lie group.

Note that certain Lie groups are known as “simple” because as manifolds they have no boundaries and no nontrivial limit points (For example, $GL(n)$ is not simple because there is a nontrivial limit — you can continuously approach matrices which have determinant zero (or are not invertable) and are therefore not part of the group. The set of simple Lie groups (including, $SU(N)$ and $SO(N)$ and just a few others) is extremely highly studied.

A **Lie Algebra** is the algebra generated by elements infinitesimally close to the identity in a Lie group. For matrix valued Lie groups $G$, we can write any element $g \in G$ as
\[
g = e^X = 1 + X + (X)^2/2 + \ldots
\]
where $X$ is an element of the corresponding Lie algebra (make it have small amplitude such that $g$ is infinitesimally close to the identity). Conventionally if a Lie group is denoted as $G$ the corresponding Lie algebra is denoted $\mathfrak{g}$.

- For the Lie group $SU(2)$, we know that a general element can be written as $g = \exp(i \mathbf{n} \cdot \sigma)$ where $\mathbf{n}$ is a real three-dimensional vector and $\sigma$ are the Pauli matrices. In this case $i\sigma_x$, $i\sigma_y$ and $i\sigma_z$. 

---

8 Pronounced “Lee”, named after Sophus Lie, also a Norwegian Mathematician of the 1800s. Like Ski-Jumping, Norway seems to punch above its weight in the theory of groups.

9 A slightly more rigorous definition is that a Lie algebra is an algebra of elements $u,v,w...$ which can be added with coefficients $a,b,c$ to give $X = au + bv + cw + \ldots$ where we have a commutator $[\cdot,\cdot]$ which satisfies $[X,X] = 0$ for all $X$ as well as bilinearity $[au + bv, X] = a[u,X] + b[v,X]$ and similarly $[X, au + bv] = a[X,u] + b[X,v]$ for all $X,a,b,u,v$, and finally we must have the Jacobi identity $[[X,Y],Z] + [[Y,Z],X] + [[Z,X],Y] = 0$. 

---

548 Some Mathematical Basics
are the three generators of the Lie algebra $\mathfrak{su}(2)$ (in the, so-called, fundamental representation).

- For the Lie group $GL(n, \mathbb{R})$ the corresponding Lie algebra $\mathfrak{gl}(n, \mathbb{R})$ is just the algebra of $n \times n$ real matrices.

Add something about Lie Algebra?

### 41.2.4 Representations of Groups:

A representation is a group homomorphism. This means it is a mapping from one group to another which preserves multiplication. We will be concerned with the most common type of representation, which is a homomorphism into the general linear group, i.e., the group of matrices. Almost always we will work with complex matrices. Thus an $n$-dimensional representation is a mapping $\rho$ to $n$-dimensional complex matrices

$$\rho : G \rightarrow GL(n, \mathbb{C})$$

preserving multiplication. I.e.,

$$\rho(g_1)\rho(g_2) = \rho(g_1g_2)$$

for all $g_1, g_2 \in G$.

Typically in quantum mechanics we are concerned with representations which are unitary, i.e., $\rho(g)$ is a complex unitary matrix of some dimension. (In case you don’t remember, a unitary matrix $U$ has the property that $UU^\dagger = U^\dagger U = 1$).

A representation is reducible if the representing matrices decomposes into block diagonal form. I.e., $\rho$ is reducible if $\rho = \rho_1 \oplus \rho_2$ for two representations $\rho_1$ and $\rho_2$. An irreducible representation is one that cannot be reduced.

The dimensions of the irreducible representations follow the law

$$\sum_R \dim(R)^2 = |G|$$

(41.4)

where $|G|$ is the number of elements in the group and the sum is over irreducible representations.

An amazing fact from representation theory of discrete groups is that the number of irreducible representations of a group is equal to the number of distinct conjugacy classes.

**Schur’s Second Lemma** is a very useful result stating that if a matrix $A$ commutes with every $\rho^R(g)$ for all elements $g$ in the group for an irreducible representation $R$, then $A$ is proportional the unit matrix. In particular this means that any element $h$ that commutes with all elements of the group has $\rho^R(h)$ a complex phase times the identity. A corollary of this is that all of the irreducible representations of abelian groups are one dimensional.

Following from Shur’s lemma we can, for example, write all the irreducible representations of the group $\mathbb{Z}_N$ in the following way. Let the
group $\mathbb{Z}_N$ be written as $g = 0, \ldots, (N - 1)$ with the operation of addition. Let $\omega = e^{2\pi i/N}$. There are exactly $N$ irreps of this group which we label as $\omega^p$ for $p = 0, \ldots, (N - 1)$. These irreps are given by

$$\rho^p : g \rightarrow (\omega^p)^g$$

(41.5)

Orthogonality and Characters

Irreducible unitary representations matrices satisfy a beautiful orthogonality relationship known as the grand orthogonality theorem (or Schur orthogonality)

$$\sum_{g \in G} [\rho^R(g^{-1})]_{mn} [\rho^{R'}(g)]_{pq} =$$

$$\sum_{g \in G} [\rho^R(g)]^*_{nm} [\rho^{R'}(g)]_{pq} = \frac{\delta_{np}\delta_{mq}\delta_{RR'}}{|G|} \dim(R)$$

(41.6)

where the superscript $R$ indicates a particular irreducible representation, the subscript are the matrix elements of the $\rho$ matrix, $\dim(R)$ is the dimension of the representation $R$, and $|G|$ is the total number of elements in the group.

A character is the trace of a representation matrix.

$$\chi_R(g) = \text{Tr}[\rho^R(g)]$$

(41.7)

where the superscript $R$ indicates we are considering a particular representation $R$. Because of the cyclic property of the trace $\text{Tr}[ab] = \text{Tr}[ba]$ the character is the same for all elements of a conjugacy class. One can find tables of characters for different groups in any book on group theory or on the web.

Representation theory of groups is a huge subject, but we won't discuss it further here!

41.3 Fundamental Group $\Pi_1(\mathcal{M})$

A powerful tool of topology is the idea of the fundamental group of a manifold $\mathcal{M}$ which is often called the first homotopy group, or $\Pi_1(\mathcal{M})$. This is essentially the group of topologically different paths through the manifold starting and ending at the same point.

First, we choose a point in the manifold. Then we consider a path through the manifold that starts and ends at the same point. Any other path that can be continuously deformed into this path (without changing the starting point or ending point) is deemed to be topologically equivalent (or homeomorphic, or in the same equivalence class). We only want to keep one representative of each class of topologically distinct paths.

These topologically distinct paths form a group. As one might expect, the inverse of a path (always starting and ending at the same point) is given by following the same path in a backward direction. Multiplication
41.3 Fundamental Group $\pi_1(M)$

of two paths is achieved by following one path and then following the other to make a longer path.

41.3.1 Examples of Fundamental Groups

- If the manifold is a circle $S^1$ the topologically distinct paths (starting and ending at the same point) can be described by the number $n$ of clockwise wrappings the path makes around the circle before coming back to its starting point (note $n$ can be 0 or negative as well). Thus the elements of the fundamental group are indexed by a single integer. We write $\pi_1(S^1) = \mathbb{Z}$.

- If the manifold is a torus $S^1$ the topologically distinct paths can be described by two integers indicating the number of times the path winds around each handle. We write $\pi_1(S^1 \times S^1) = \mathbb{Z} \times \mathbb{Z}$.

It is in fact, easy to prove that $\pi_1(M_1 \times M_2) = \pi_1(M_1) \times \pi_1(M_2)$.

- A fact known to most physicists is that the group of rotations of three dimensional space $SO(3)$ is not simply connected — a $2\pi$ rotation (which seems trivial) cannot be continuously deformed to the trivial rotation, whereas a $4\pi$ rotation can be continuously deformed to the trivial rotation.\(^{10}\) Correspondingly the fundamental group is the group with two elements $\pi_1(SO(3)) = \mathbb{Z}_2$.

\(^{10}\)This is the origin of half-odd integer angular momenta.

Chapter summary

Some mathematical ideas introduced in this chapter:

- **Manifolds** are locally like Euclidean space: Examples include sphere $S^2$, circle $S^1$, torus surface $T^2 = S^1 \times S^1$, etc. Manifolds can also have boundaries, like a two dimensional disk $B^2$ (or $D^2$) bounded by a circle.

- **Groups** are mathematical sets with an operation, and identity and an inverse: Important examples include, $\mathbb{Z}$ the integers under addition, $\mathbb{Z}_N$ the integers mod $N$ under addition, the symmetric (or permutation group) on $N$ elements $S_N$, and Lie groups such as $SU(2)$ which are also manifolds at the same time as being groups.

- **The Fundamental Group** of a manifold is the group of topologically different paths through the manifold starting and ending at the same point.

- **Isotopy** is the topological equivalence of knot diagrams (what can be deformed to what without cutting).

- **Writhe and Linking Number** characterize pictures of oriented knots and links.
For background on more advanced mathematics used by physicists, including some topological ideas, see:


For further information on mathematics of knots, isotopy, and Riedemeister moves, writhing and linking, see


Something on group theory
(1) A general reference which should be useful for much of the book is the review article by Nayak, Simon, et al. [Nayak et al. 2008].

(2) A wonderful little book which is really fun to read that introduces the Kauffman invariant and many other ideas of knot theory is the book Knots and Physics by Kauffman [Kauffman 2001], now in its 3rd edition. This book really inspired me when I was a grad student. It appears to be available online in several places (not certain which, if any, are legal). Although the whole book is fun; and much of it is written at a very introductory level, mainly the end of part 1 is the most relevant part where he explains the connection of Kauffman invariant to Chern-Simons theory (and pieces get to be well beyond introductory). There is a lot in here; the deep parts are easy to gloss over.

(3) A very nice introduction to non abelian anyons and topological quantum computation is given in John Preskill’s lecture notes, available online [Preskill 2004].

(4) Frank Wilczek has two books which both discuss Berry phase and abelian anyons [Wilczek 1990]. Both have mainly reprints in them with some commentary by Wilczek. Often it is enough to read the commentary!

(5) If you need a refresher on path integrals, consider the first 15 pages of Fabian Essler’s notes [Essler 2004]. Also consider the nice article by Richard MacKenzie [MacKenzie 2004]. MacKenzie includes some useful applications such as Aharanov-Bohm effect. Look mainly at the first 22 pages.

(6) The classic paper by Ed Witten which launched the field is Witten [Witten 1989]. This is a tremendously deep paper which introduces a lot of brilliant ideas. I find something new every time I read it. I find it to be tough reading in some places and easy in others.

(7) From a more mathematical viewpoint several articles by Sir Michael Atiya are very useful [Atiya 1988, 1997]. These are both introductions to topological quantum field theories. There is also a more detailed book by the same author [Atiya 1990]. The full book might be hard to read unless you have a very strong maths background.

(8) There are several nice references on the structure of topological quantum field theories and diagrammatic calculus, Parsa Bonderson’s thesis: http://thesis.library.caltech.edu/2447/2/thesis.pdf. This is a more detailed version of the long article by Kitaev (“Anyons in exactly solvable models”) which I mention below. Note there is
some slight change of convention between the two articles.
Also a good reference is the book on Topological Quantum Computation by Zhenghan Wang
If you are more mathematical, you might like the thesis of Bruce Bartlett available online here
https://arxiv.org/abs/math/0512103

https://arxiv.org/abs/cond-mat/0506438
This brings the ideas of topological quantum field theory into the condensed matter arena. This is not easy reading, but a ton of great ideas are buried in this paper.
introduces the famous toric code, discusses quantum error correction, and generalizes the toric code model to arbitrary non-abelian groups.
Kitaev’s work on the quantum wire (which we might get to at the end of the course) is here.
https://arxiv.org/abs/cond-mat/0010440
A brief digest of some of the many ideas introduced in these three papers is given by notes taken by Laumann of Kitaev’s lectures, available here.
https://arxiv.org/abs/0904.2771
Loop gases are introduced in this paper by Freedman et al. It has a lot of sections which are hard to parse.
http://stationq.cnsi.ucsb.edu/ freedman/Publications/83.pdf
The double-fibonacci string-net is discussed in some detail in this work by Fidkowski et al,
https://arxiv.org/abs/cond-mat/0610583
The classic paper on string-nets very generally is this by Levin and Wen.
https://arxiv.org/abs/cond-mat/0404617
The standard reference on introductory quantum hall effect is the classic book, ”The Quantum Hall Effect”, edited by Prange and Girvin, published by Springer. The first chapter, and the chapters by Laughlin and Haldane are probably the best. The experimental chapters are good for context too.
A short review article by Macdonald is pretty nice and is available here.
The article that introduced the ideas of conformal field theory into the field of quantum Hall effect is by Moore and Read, available online here.
http://www.physics.rutgers.edu/gmoore/MooreReadNonabelions.pdf
A recent review article on Fractional quantum Hall hierarchies (and also discusses nonabelian quantum Hall and conformal field theory) is online here.
https://arxiv.org/abs/1601.01697

A few random digressions:

(10) If you are interested in 2+1 D quantum gravity, see this article .
I can’t vouch for it, but the introduction is interesting;
This is the article by Witten explaining how 2+1 D gravity is ”ex-actly solvable.” More from Witten here. There is reconsideration many years later, again by Witten, see here .

(11) I’ve been told the book by Jiannis Pachos on topological quantum computation is a good resource.

(12) If you are interested in the topology of manifolds in 3 and 4 dimensions, there are several good books. One by Kirby is online here.
https://math.berkeley.edu/kirby/papers/Kirby
There is a book by Gompf and Stipicz ”4-manifolds and Kirby Calculus” which is nice. Note that parts of this book are online free if you google them.
https://www.amazon.co.uk/4-Manifolds-Calculus-Graduate-Studies-Mathematics/dp/0821809946

(13) For more information on conformal field theory. The standard reference is the Big yellow book (Conformal Field Theory Authors: Philippe Di Francesco, Pierre Mathieu, David Sénéchal) . The first part of this book (up to chapter 12) is excellent, but even that much is a lot of reading. There is a short set of lectures from les Houches by Ginsparg.
I also like the short set of notes by Fendley .
http://galileo.phys.virginia.edu/pf7a/msmCFT.pdf
For even shorter introduction of what you need to apply CFT to quantum Hall, see the appendix of Ref. 1 above, or the appendix of ***.
The book by Kauffman and Lins gives more details of constructing a full anyon theory from the kauffman invariant.
http://press.princeton.edu/titles/5528.html
Neilsen and Chuang for quantum computation in general, although there are plenty of other refs.
References


017. URL http://www.sciencedirect.com/science/article/pii/S0022404918303098


O. Buerschaper and M. Aguado. Mapping kitaev’s quantum double lattice models to levin and wen’s string-net models. *Phys. Rev. B*, 80:


REFERENCES


Y. Hu, S. D. Stirling, and Y.-S. Wu. Ground-state degeneracy in the

Y. Hu, Y. Wan, and Y.-S. Wu. Twisted quantum double model of

Y. Hu, Y. Wan, and Y.-S. Wu. Twisted quantum double model of


C. Jones and D. Penneys. Operator algebras in rigid $c^*$-tensor cate-
gories. *Communications in Mathematical Physics*, 355(3):1121–1188,
Nov 2017. ISSN 1432-0916. doi: 10.1007/s00220-017-2964-0. URL
https://doi.org/10.1007/s00220-017-2964-0

V. F. R. Jones. A polynomial invariant for knots via von neumann
1985.

L. Kauffman. State models and the Jones polynomial. *Topology*, 26:

L. Kauffman and S. Lins. *Temperley Lieb Recoupling theory and invari-


R. Kirby. A calculus for framed links in $s^3$. *Inventiones Mathematicae*,

R. Kirby and P. Melvin. Canonical framings for 3-manifolds. *Turkish J.
Math; Proceedings of 6th Gokova Geometry-Topology conference*, 23

R. C. Kirby. *The Topology of 4-Manifolds*, volume 6 of *Nankai Institute

publication date 2003.


F. Ladisch. Finite groups in which every character has real values grading the representation. *Math Overflow*, 2011. URL [https://mathoverflow.net/questions/53126/finite-groups-in-which-every-character-has-real-values-grading-the-representati](https://mathoverflow.net/questions/53126/finite-groups-in-which-every-character-has-real-values-grading-the-representati).


S. MacLane. Categories for the working mathematician. Springer-Verlag, 1971.


REFERENCES


