New exercises for Entropy, Order Parameters, and Complexity: Statistical Mechanics

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2 Advanced Statistical Mechanics

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Chapter 1

Entropy, Order Parameters, and Complexity: New Exercises


N1.1 Entropy and currents. ☞

Consider a locally conserved density $\rho(x,t)$ in an isolated one-dimensional system with a current $J(x,t)$. Imagine that the system has a complicated diffusion constant, with strong density dependence and also dependent on position, so $J(x,t)$ may not be simply related to $\rho$. Imagine, however, that the internal energy is not important in the system, so entropy $S = -k_B \int \rho \log \rho \, dx$ must increase with time. Are there conditions on $J(x,t)$ which would guarantee this?

Show that the entropy will increase with time if $J(x,t)$ is in the direction of decreasing $\rho$. (This is closely related to Exercise 5.10; again, you will need to integrate by parts. You may assume $J(x)$ and $\rho(x)$ go to zero as $x \to \pm \infty$.)
N1.2 **First-digit law and priors.** (Statistics) \(\square\)

Bayesian statistics, like statistical mechanics, incorporates known experimental results into a probabilistic prediction for the behavior of the system in the future (see Exercise 6.14). In statistical mechanics, if we only know the energy of a system then Liouville’s theorem tells us that all points in the energy shell are equally likely *a priori*. In Bayesian statistics, they have no theorem like Liouville’s, so they need to assume a *prior*. For example, if you want to estimate a time constant \(\tau\) for a chemical reaction (which can range from nanoseconds to years), you might want a prior \(P_\tau(\tau)\) that gives equal weight to each decade: finding \(\tau\) in the range \((10^{-9}, 10^{-8})\) seconds is equally plausible as finding \(\tau\) in the range \((10^5, 10^6)\) seconds.

Show that \(P_\tau(\tau) \propto 1/\tau\) has this reasonable property. Show that this choice also makes the decay rates \(\Gamma = 1/\tau\) have this same nice property: \(P_\Gamma(\Gamma) \propto 1/\Gamma\). (Hint: If \(\tau\) lies in a small range \(\Delta\tau\), then \(\Gamma\) will lie in a corresponding small range \(\Delta\Gamma\), so \(P_\Gamma(\Gamma)|\Delta\Gamma| = P_\tau(\tau)|\Delta\tau|\).) Show that this distribution predicts that the first non-zero decimal digit \(d\) of \(\tau\) will have probability \(\log_{10}(1 + 1/d)\) (Fig. N1.1). (Hint: Do it assuming \(\tau\) lies in one decade first.) Show your steps. (Note: Feel free to consult the extensive discussions on the Web.)

Simon Newcomb, using a book of logarithms in 1881\(^1\) discovered this by noticing that the pages in the beginning (1.000001, 1.000002, \ldots) were dirtier than the ones at the end (9.000001, 9.000002, \ldots). Frank Benford fleshed this out in 1938, showing that areas of rivers, molecular weights of compounds, and physical constants like the proton mass, Planck length, and Avogadro’s constant (Fig. N1.1) also obey this law.

N1.3 **Accelerators vs. ergodicity.**\(^2\) (Accelerator, Mathematics, Computation, Dynamical

---

\(^1\)Before calculators, people used printed books of logarithms, which allow one to multiply and divide quickly.

\(^2\)Hints for the computations can be found at the book website [31].
Fig. N1.2 Orbit of particle around synchrotron. Dashed line shows the intended center of the beam; solid line the trajectory of a particular particle. The particle oscillates around the center with a phase \( \theta \) and an amplitude proportional to \( \sqrt{p} \), with anharmonicity determined by \( K \). The standard map is a Poincaré section; it tells one what \( \theta \) and \( p \) will be as the particle next crosses the horizontal axis along the red line.

Synchrotrons push charged particles around circular orbits at near the speed of light, stably over billions of orbits (Fig. N1.2). Why haven’t the nonlinear fields used to steer the particles kicked them out of the beam? Like the cat map in Exercise 5.8 and the three-body problem of Exercise 4.4, physicists distill the dynamics using a Poincaré section into a two-dimensional area-preserving nonlinear map. The classic example, due to Chirikov, is the so-called standard map

\[
\begin{align*}
\theta_{n+1} &= \theta_n + p_n + K \sin(\theta_n) \\
p_{n+1} &= p_n + K \sin(\theta_n).
\end{align*}
\]  

(N1.1)

Here \( p \) is proportional to the squared amplitude of the deviation from center of beam and \( \theta \) represents the phase of the oscillation around that center (Figure N1.2). Large \( K \) represents a strong nonlinearity in the fields. For plotting, we shall take \( \theta \mod 2\pi \), and occasionally also \( p \mod 2\pi \).

We remember from Exercise 5.8 that area-preserving maps mimic the volume-preservation in phase space guaranteed by Liouville’s theorem.

(a) Show that the standard map (eqn N1.1) preserves area, by showing that its Jacobian has determinant one.

(b) Iterate the standard map a few thousand times for \( K = 4 \), starting with \( (\theta_0, p_0) = (0.1, 0.11) \) (chosen arbitrarily), and plot \( p_n \) versus \( n \) for a thousand points \( n \). (For this part, do not take \( p \mod 2\pi \)). Does the amplitude of the particle oscillations around the beam path (as measured by \( p \)) remain bounded in amplitude as \( n \) gets large? Would designing an accelerator with this large an anharmonicity be a good idea?

The situation changes for smaller anharmonicity.
CHAPTER 1. EOPC: NEW EXERCISES

Fig. N1.3 Trajectories at $K = 0.2$ for a variety of initial conditions.

(c) Implement the standard map graphically, so that you can interactively select a variety of initial conditions and run each for a few thousand steps to explore the trajectories, using mod($2\pi$) for $\theta$ but not for $p$. Check that you get trajectories for $K = 0.2$ that are compatible with those shown in Fig. N1.3. Plot the behavior for $K = 0.7$, selecting a variety of initial conditions. Be sure to illustrate “oval” KAM$^3$ tori around stable periodic orbits, “horizontal” KAM tori that span from $\theta = 0$ to $2\pi$, and chaotic regions. (Remember that your trajectory is a cross section once per orbit around the accelerator. So a closed curve becomes a tube, which mathematically is called a torus.) Explore initial conditions spanning $p \in (0, 2\pi)$. Do the trajectories stay bounded in $p$? Which keeps $p$ from growing indefinitely – the oval or horizontal tori? Keeping the magnets close to the beam is best – until the beam starts hitting the magnets. The coordinate $p$ is related to the squared maximum deviation of the particle from the designed beam center. Based on your plot, what range $\Delta p$ would you use to set the magnet positions for an accelerator at $K = 0.7$?

The equilibrium state predicted from statistical mechanics would fill all of phase space uniformly. The map at $K = 0.7$ is not ergodic.

In Section 4.2 we defined a map (or time evolution) to be ergodic if and only if all the ergodic components of the space either have zero volume or have a volume equal to the space. Here ergodic components were sets $R$ that remain invariant under the map (so if $(\theta_0, p_0)$ is in $R$ then $(\theta_n, p_n)$ is also in $R$).

(d) Identify a few ergodic components in your plot of $K = 0.7$ in part (c). Are the KAM tori ergodic components? Do they have non-zero volume? Do they surround ergodic components with non-zero volume? Does the chaotic regions surrounding $\theta \approx p \approx 0$ appear to be an ergodic component with non-zero volume?

(e) Alter your standard map to also take $p \mod 2\pi$, to keep the motion bounded.$^4$ Plot

---

$^3$KAM stands for Kolmogorov, Arnol’d, and Moser, who proved that tori survive for certain irrational winding numbers; see part (e).

$^4$Note that your map of $K = 0.7$ in part (c) is periodic in $p$ with period $2\pi$, so this makes sense.
and examine carefully the time evolution at $K = 4$. Do all initial conditions fill out the entire volume? If not, describe ergodic components that neither have zero volume nor fill the square.

We define the **winding number** of a particle trajectory as the average number of oscillations $\theta/2\pi$ per iteration of the map. The KAM theorem tells us that the tori whose winding numbers are difficult to approximate by rationals are the most stable.

(f) What is the winding number at $K = 0$, for a trajectory starting at initial condition $(\theta_0, p_0)$? Write a routine to approximate the winding number for $K = 0.7$ and $\theta = \pi$, and plot it for $0 < p < 2\pi$. By examining your plot of the different trajectories for $K = 0.7$ from part (c), explain the plateaus you observe at rational winding numbers.

The last “horizontal” KAM torus for the standard map is destroyed by the nonlinearity at $K_c = 0.971635 \ldots$. Its winding number is the inverse Golden Mean, $(\sqrt{5} - 1)/2 = 0.618033 \ldots$, which is the most irrational number – the number hardest to approximate by ratios of small integers. This transition is associated with self-similar behavior and scaling, and has been studied using the renormalization group methods we study in Chapter 12.

### N1.4 Bosons in two states. (Quantum) 3

(a) Consider three noninteracting identical bosons in a system with two single-particle energy eigenstates at energies $E_0 = 0$ and $E_1 = \epsilon$ at temperature $T$. Give a formula for the partition function $Z$ and the expected number of bosons $\langle n \rangle$ in the excited state $E_1$, as a sum over states.

(b) What would your answer be for $\langle n \rangle$ if there were an infinite number of bosons distributed between the two levels? What is the expected number of bosons in the upper state at temperature $k_B T = 10\epsilon$? (Hint: How does your answer relate to the thermal occupation probabilities for harmonic oscillator eigenstates?)

We treat the excitations of harmonic oscillators as particles (phonons and photons) because they act like noninteracting bosons (see Exercise 7.2). Each harmonic oscillator acts like a different single-particle eigenstate, for a grand canonical ensemble of bosonic photons with chemical potential zero. The $n^{\text{th}}$ harmonic excitation is $n$ photons occupying the state. Our lower level here can be viewed as a particle analogy of the heat bath; the heat bath sets the temperature, our lower level sets the chemical potential.

(c) Now consider a system with 1000 distinguishable, noninteracting particles that can each occupy the same two states, $E_0 = 0$ and $E_1 = \epsilon$. What is the expected number of distinguishable particles in the upper state at temperature $k_B T = 10\epsilon$? Discuss how and why the behavior of bosons and distinguishable particles are different in this exercise.

(d) Are the bosons in part (b) forming a Bose–Einstein condensate?
CHAPTER 1. EOPC: NEW EXERCISES

N1.5 Beer and bubble nucleation.

A glass of beer illustrates many aspects of statistical mechanics. The head of foam at the top of the beer exhibits rigidity like a solid (Exercise N1.7), and the foam coarsens with time (Exercise N1.6). In this exercise, we shall study the phase transitions triggered when one opens the bottle. You may wish to experiment with a bottle or can of beer or soda.

We shall model beer and soda, for the purposes of this exercise, as a phase consisting of carbon dioxide and water. When one opens the bottle, the absolute pressure drops from around $P_{\text{beer}} \sim 200 \text{kPa}$ (depending on the beer), to atmospheric pressure $P_{\text{atm}} \approx 100 \text{kPa}$. The solubility of CO$_2$ at 0 C in water at atmospheric pressure is about 1.8 cm$^3$ CO$_2$ per gram of water [15]. Note that there is a small amount of gas above the beer. Henry’s law tells us that the number of dissolved CO$_2$ molecules is proportional to its partial pressure in the gas.\(^5\) H$_2$O at 0 C has a partial pressure of 0.6 kPa, so we assume that the pressure in the gas atop the beer bottle is almost completely due to CO$_2$.

When you pour a glass of beer, much of the volume is filled with foam. Bartenders will pour off the foam to give you a full glass of liquid. Assume the carbonated liquid starts at $P_{\text{beer}}$, and ignore the volume change in the liquid as the pressure drops and the gas separates. Assume the two form a closed system.

(a) As the pressure drops and the liquid separates from the vapor, what fraction of the volume in the system at the new equilibrium is gas? Does that correspond qualitatively to your observations? (Note that cold beer may need to be shaken considerably to extract all of the gas.)

Let $x_{\text{CO}_2}$ be the fraction of the molecules in the liquid that are CO$_2$ molecules. Henry’s law for our system can be written

$$P_{\text{CO}_2} = K x_{\text{CO}_2}, \quad (\text{N1.2})$$

with Henry’s constant $K \approx 70 \text{MPa}$ at 0 C (ice-cold beer) [6].

(b) Compare the solubility determined from Henry’s constant to the quoted solubility 1.8 cm$^3$/gram.

We shall use $X$ to denote the fraction of CO$_2$ molecules in the entire system: the total number of CO$_2$ molecules (both dissolved and in vapor), divided by the total number of molecules including the water (both liquid and vapor). Hence $x = X$ if the CO$_2$ is entirely dissolved in the liquid.

The vapor pressure of water at $T = 0 \text{C}$ is $\sim 0.6 \text{kPa}$, independent of the pressure and roughly independent of the amount of CO$_2$ dissolved. CO$_2$ is a gas at all pressures of interest, and both gases are well-approximated as ideal gases of molecules. Let us draw the P–X phase diagram.

---

\(^5\)The pressure of a gas mixture in the ideal gas approximation is the sum of the partial pressures contributed by each constituent.
(c) Use Henry’s law to determine the total pressure (vapor pressure of water plus CO$_2$ partial pressure) at which the liquid beer begins to coexist with the vapor. Find the pressure at low pressures and high CO$_2$ molar fraction $X$ at which the liquid disappears (the gas begins to coexist with the liquid). Draw the $P$–$X$ phase diagram for the H$_2$O/CO$_2$ system at $T = 0$ C, for $P \in (0, 400)$ kPa and $X \in (0, 0.01)$, and include a line connecting the beer initial and final pressures.\(^6\) Draw the phase diagram again for $P \in (0, 10)$ kPa and $X \in (0, 1)$ to illustrate the other side of the coexistence curve. In both diagrams, indicate the liquid phase, the gas phase, and the two-phase coexistence region.

It is apparent that the phase separation in beer is nucleated through the formation of bubbles of gas. Let us calculate the nucleation rate of a spherical bubble. The surface tension of water in contact with air is $\sigma \sim 7.56$ N/m at 0 C. The prefactor $\Gamma_0$ for the Arrhenius rate (called “prefactors” in eqn 11.11) may be roughly estimated \([5, 3]\) as $\Gamma_0 = 4 \times 10^{41}$ m$^{-3}$ s$^{-1}$, corresponding to a physically sensible prefactor $\approx 10^{13}$/s of the number of attempts to form a critical droplet centered at each water molecule in the cubic meter.

(d) What is the pressure energy gained by forming a bubble of radius $R$? What is the surface tension energy cost? What is the radius of the critical bubble? What is the nucleation barrier for generating a bubble in the bulk of the beer (homogeneous nucleation)? Would you plan to wait until a bubble forms when you open a half-liter bottle of beer?

Water has an unusually large surface tension, and our calculation clearly shows that homogeneous nucleation cannot occur. Even for extremely clean liquids with low surface tension (like liquid helium), homogeneous nucleation almost always is unimportant compared to nucleation on surfaces, dust particles, dislocation tangles, scratches on surfaces, etc. Many of your beer bubbles probably form on small flaws on the glass surface. If your glass of beer has not gone flat yet, see if you notice a chain of bubbles all emitting from the same spot on the glass.

N1.6 Beer foam and coarsening.\(^7\)

Pour a beer. Note that the bubbles rise to the top under gravity, and form a foam—the “head” of a beer. (If you do not have a beer handy, soap suds or shampoo bubbles will work as well.) As time goes on, the bubbles in the foam will grow larger, as the big bubbles absorb the little ones. This is loosely reminiscent of the coarsening seen after abrupt phase transitions (Section 11.4.1).

There are two major mechanisms for this coarsening: the popping of the walls between bubbles as they grow thin, and the diffusion of gas between bubble walls. Which is more important can be controlled through changing the composition of the fluid. In

\(^6\)Let $X$ be determined by the liquid portion of the beer; assume the small amount of gas at the top escaped when we open the bottle.

\(^7\)Thanks to Douglas Durian for helpful consultations and Fig. N1.4.
this exercise, we shall study a theory of bubble coarsening via diffusion, following a classic analysis in two dimension by von Neumann [33] and Mullins [23] (see [18]).

The bubbles near the top of the beer are dry; they have thin walls separating relatively large bubbles. In a “two-dimensional” experiment with thickness much smaller than the bubbles, these walls form a network of curves connecting nodes where three bubbles touch (Fig. N1.4). These walls move to balance the forces on them, on a time scale much faster than the diffusion of gases between bubbles that drive the coarsening we study.

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{foam_scaling.png}
\caption{Foam scaling. Two snapshots of a simulated 2D foam, coarsening by diffusion, with elapsed time $t$ in minutes, and a rescaled version of the early time snapshot (graciously provided by Anthony Chieco and Douglas Durian). Note that the patterns look statistically similar, apart from an overall growth in the size of the bubbles.}
\end{figure}

Let us first examine the structure of the force-balanced configurations. Let the line tension (energy per unit length) of the walls be $\lambda$, and assume the energy per node is negligible.

(a) Draw a free body diagram for the forces on a node at a three-wall intersection. (Nodes where more than three bubbles meet are non-generic. That is, any tiny perturbation will split them into three-fold intersections.) What force will each wall exert on the node? Show that the three walls must intersect at $120^\circ$ angles.

Each bubble will be filled with a gas of a certain pressure. The pressure difference $\Delta P$ across a wall determines its radius of curvature $R = \Lambda/\Delta P$.

(b) Determine the constant $\Lambda$ in terms of the wall tension $\lambda$. 
Fig. N1.5 2D Bubble geometry. A thin-walled bubble with five neighbors, whose walls are arcs of circles of angle $\theta_i$, radii $R_i$, and arc length $c_i$. The bubble walls meet at nodes, forming angles of $2\pi/3 = 120^\circ$ (red dashed lines). The local tangent vector to the bubble perimeter winds by $2\pi$ as one circles the bubble (black arrows), but jumps by $\pi/3 = 60^\circ$ every time it passes a node. It rotates by the opening angle $\theta$ during the arc $c_i$ for each bubble.

Hence a thin-walled bubble with $n$ neighbors with pressure changes $\Delta P_i$ will be formed by $n$ circular arcs of radii $R_i \propto 1/\Delta P_i$ connecting the nodes where walls meet. As one might imagine, the rate of gas diffusion through the wall per unit length is given by a constant $\mu$ times the pressure drop $\Delta P$. So

$$\frac{dA_n}{dt} = \mu \sum_{i=1}^{n} \Delta P_i c_i$$

where $c_i$ is the arc length of the corresponding wall (Fig. N1.5).

John von Neumann, known for his work in computer science and game theory, deduced a law for the size evolution of two-dimensional domains [33] like our bubbles. The area $A_n$ of a domain with $n$ neighbors will vary as

$$\frac{dA_n}{dt} = K_0(n - 6)$$

The bubbles with only a few neighbors vanish. But the average number of neighbors should stay around six (which you can show using Euler’s theorem about faces, edges, and nodes).

(c) Find a video of this coarsening process, such as that in Ref. [16]. Identify a bubble with six neighbors. Does it stay roughly the same size? When it adds / subtracts a neighbor, does it start growing / shrinking as expected? Identify a bubble with more than six neighbors at early times, and follow its evolution. Does it die? (If not, start again with a second many-neighbored bubble. Most bubbles should eventually disappear, since so few survive.) Does the number of neighbors of your bubble fluctuate both up and down?

What was von Neumann’s argument? It was purely geometrical.
(d) Show that \( \Delta P_i c_i = \Lambda \theta_i \), where \( \theta_i \) is the angle between the tangents to the \( i^{th} \) boundary at the nodes on either end as shown in Fig. N1.5 (negative for convex boundaries ‘bulging out’, positive for concave boundaries). Noticing that the tangent vector jumps by \( \pi/3 = 60^\circ \) at each node (Fig. N1.5) and that the tangent vector must rotate by \( 2\pi \) as it winds around the bubble, derive von Neumann’s law (eqn N1.4). What is \( K_0 \) in terms of \( \mu \) and \( \Lambda \)?

For non-conserved order parameters, we saw in Section 11.4.1 that the typical length scale \( L(t) \sim t^\beta \) with \( \beta = \frac{1}{2} \). For conserved order parameters, where the particles or spins must diffuse over a length scale \( L(t) \) to rearrange the structure, we found \( \beta = \frac{1}{3} \).

What is \( \beta \) for 2D dry foams? In Section 11.4.1, we argued qualitatively that if droplets or features of size \( L \) disappear in a time \( t \propto L^{1/\beta} \), then the typical size of the surviving features will grow as \( L \sim t^\beta \). In the next part, you can give an analogous argument for foams.

(e) In part (c), you found that the number of neighbors of a droplet changed in a complicated way with time. For simplicity, let us consider a droplet with a constant number of neighbors \( n \), which at \( t = 0 \) has area \( A_0 \). Write a formula for \( A(t) \). At what time \( t_0 \) does \( A(t) \) vanish? What happens to \( t_0 \) for \( n > 6 \)? If we define \( L(t) = \sqrt{A(t)} \), write \( L(t) = C(n)|t - t_0|^{\beta} \). What is \( C(n) \)? Does \( \beta \) depend on the number of sides \( n \)?

See also Exercise N1.5, which studies the bubble formation in beer, and Exercise N1.7, which studies the transition from liquid beer to rigid foam . . .

N1.7 Beer and rigidity: Jamming.

Beer is a liquid: the bubbles in the beer do not prevent it from pouring. But the head of a beer (the frothy foam on the top of the beer where the bubbles accumulate) is rigid. The foam is a random packing of bubbles that can support shear, just as the glass containing the beer is a random arrangements of atoms bonded together that does not flow under stress.

Fig. N1.6 A head of beer. Beer is liquid on the bottom and forms a rigid network of bubbles in the foam at the top.

---

8This exercise was developed in collaboration with Stephen Thornton.
As in Exercise N1.5, we can view the beer as exploring a slice of the vapor/fluid phase diagram, with gravity pulling the fluid downward leading to polyhedral bubbles separated by thin walls forming the head, gradually becoming rounder and wetter with depth, until they separate into individual spherical bubbles in the bulk of the glass.

Somewhere, as we move upward in the glass, there must be a first point where we go from a fluid phase to a rigid phase. What is the nature of this phase transition?

For large objects like bubbles, powders, sand, and colloids, this transition is a problem in mechanics. When two bubbles touch, they resist moving closer together, which we can think of as an overlap energy. The bubbles can therefore pack together without costing overlap energy only until there are enough contacts that each bubble is held in place by its neighbors.

This kind of rigidity was first studied by Maxwell in his study of the stability of mechanical trusses. A contact between two spherical bubbles constrains their separation to be the sum of their radii; a truss is a rigid rod that constrains its endpoints to be a fixed distance apart. The Maxwell criterion involves subtracting the number of constraints in a system (bubble contacts) from the number of motional degrees of freedom of the system (three translations per bubble). Roughly speaking, when the two are equal the system becomes rigid. If there are fewer constraints, the system will be floppy, with each missing constraint replaced by a zero-energy mode of deformation.

**Fig. N1.7 Rigidity and Maxwell constraint counting.** An array of nodes (blue dots), connecting rotors (black lines) to springs (red dashed lines). Is the system floppy or stiff? Each rotor and each spring can be viewed as a constraint. The two coordinates of the nodes are each degrees of freedom. The green arrows indicate that there is at least one floppy mode. With permission from Chen et al. [8].

Let us consider an example, a mechanical network inspired by recent work on topological insulators. Figure N1.7 shows the right-hand side of a chain of rotors whose ends are nodes connected by springs.

---

9One can envision a tall beer in weak gravity, which could act to gradually change the chemical potential of the fluid and the gas. On Earth the gradient of chemical potential for a bubble is rather steep.

10Yes, it is the same Maxwell from electromagnetism and the Maxwell relations.

11The Maxwell criterion has various refinements. Truss networks can have states of self stress, where some regions are overconstrained. A network of fibers can become rigid with fewer constraints if they are pulled straight; this can be important in cells. Ellipsoids can become rigid even though they can still rotate. None of these subtleties are significant in bubble networks.
(a) Check, for an infinite system, that the number of constraints (springs and rotors) per node in this mechanical network would be equal to the number of directions the node can move. The figure shows that when the chain ends, the missing final spring leads to a floppy mode. The other end of the chain also ends with a rotor with only one spring attached. Use the Maxwell criterion to argue that there is only one floppy mode: the other dangling end of the chain must be rigid. (See Ref. [8] for many more fascinating aspects of the problem.)

Fig. N1.8 Force densities and tractions. (a) The shear elastic constant of a solid and the viscosity of a liquid are both defined as the response to a traction $T$ (a force per unit area) applied to two ends 0 and $L$ of a sample. (b) The transverse susceptibility $\tilde{\chi}_T(q,\omega)$ is the response to a force per unit volume $f(x,y,z,t) \propto \exp(iqx) \exp(-i\omega t) \hat{y}$. We can view the traction in (a) as a force per unit volume $f(x,y,z) = T(\delta(x-L) - \delta(x))$.

Let us prepare to study this rigidity transition by considering the response of normal solids and liquids to external shear. At long lengths and low frequencies, both liquids and isotropic solids respond to a one-dimensional, time-dependent, transverse force density $f(x,y,z,t) \equiv f(x,t) \hat{y}$ by obeying the equation

$$\rho \frac{\partial^2 u}{\partial t^2} = G \frac{\partial^2 u}{\partial x^2} + \eta \frac{\partial^3 u}{\partial x^2 \partial t} + f$$

where $u(x,t)$ is the motion in the $\hat{y}$ direction, $\rho$ is the mass density, $G$ is the shear modulus, and $\eta$ is the dynamic viscosity (see Fig. N1.8(b)).

We can test this equation by considering the case of uniform shear, given by a traction on the top and bottom surfaces of the cube $f(x,t) = T(\delta(x-L) - \delta(x))$.

(b) If the material is rigid ($G > 0$), show that this force can result in a static deformation satisfying $G \frac{\partial u}{\partial x} = T$, agreeing with the standard definition of the shear modulus.

(c) If the material is a liquid ($G = 0$), write an equation for $\dot{u}$, and show that this force produces a steady-state velocity gradient $\eta \frac{\partial \dot{u}}{\partial x} = T$, agreeing with the standard definition of the viscosity.

\[12\] Note that the viscosity is in the form of Kelvin damping, the lowest-order form allowed by Galilean invariance (see Exercises 9.6, 9.14, and 10.9).
(d) Use the equation of motion N1.5 to write the transverse susceptibility at long wavelengths \( \tilde{\chi}_T(q, \omega) = \frac{u}{\tilde{f}} \). Show that

\[
G^{-1} = \lim_{q \to 0} \lim_{\omega \to 0} q^2 \tilde{\chi}_T(q, \omega). \tag{N1.6}
\]

Show that

\[
\eta = \lim_{\omega \to 0} \lim_{q \to 0} (\rho^2 \omega^3 / q^2) \chi''_T(q, \omega), \tag{N1.7}
\]

where as usual \( \chi'' = \text{Im}(\tilde{\chi}) \).

Since our transverse susceptibility describes the typical long-wavelength, low frequency behavior of any isotropic fluid or solid, these last two formulas can be considered the definitions of shear modulus and dynamic viscosity for other systems (like foams near jamming) where the behavior is more interesting at short times and lengths.

We now turn our focus back to beer. Maxwell’s constraint on the rigidity of systems also holds for general disordered packings of spheres (like the bubbles in our beer). As we move upwards through the foam, there is a point where the foam can first support shear deformation. This is known as the jamming transition. The tunable parameter that induces this transition is \( \delta z \equiv z - z_c \), where \( z \) is the coordination number counting the average number of nearest neighbor contacts each bubble has. The critical coordination number \( z_c = 2d \) in \( d \) dimensions by Maxwell’s constraint-counting argument.

Recent work [17] provides us with an explicit formula for the universal scaling function for the transverse susceptibility near the jamming transition. Near jamming in three spatial dimensions in a viscous medium, the transverse part of the susceptibility has a particular universal scaling form in terms of our distance to jamming \( \delta z \) and intrinsic damping parameter \( \gamma \):

\[
\tilde{\chi}_T(q, \omega) = \left[ a q^2 |\delta z| \left( \sqrt{1 - c \frac{i \gamma \omega + d \rho \omega^2}{|\delta z|^2}} \pm 1 \right) \right.
\]

\[
- i \gamma q^2 \omega - \rho \omega^2 \left]^{-1} \right. \tag{N1.8}
\]

where the \( \pm \) refers to the solid “rigid foam” and liquid “wet foam” sides of the transition, respectively.

(e) Calculate shear the modulus \( G \) of the material (eqn N1.6) in terms of constants and the distance to jamming \( |\delta z| \), both on the floppy and the rigid side of the transition. Is the transition from rigid to floppy continuous, in terms of the shear elastic constant? Calculate the dynamic viscosity \( \eta \) on the floppy side of the transition (eqn N1.7) in terms of constants and the distance to jamming \( |\delta z| \). (Hint: Expand the square root to first order, and use \( 1/(a + bi) = (a - bi)/(a^2 + b^2) \). The algebra is a bit messy.) How does the viscosity depend on the distance to jamming? Explain how we might view this as a continuous transition.

\[13\]One might consider this a requirement for a material to be described as a fluid or isotropic solid.
So, unlike the abrupt way that liquids becomes rigid when they freeze into crystals, jammed solids have a continuous transition in their response to external shear. Their elastic response to shear \((1/G)\) diverges from the rigid side, and their viscous response to shear \((1/\eta)\) from the floppy side smoothly goes to zero. Equation N1.8 explains many other unusual properties exhibited by jammed solids like beer foam.

**N1.8 Where is the antimatter?**

The Universe experimentally is composed primarily of matter. This is a surprise, since in all other ways matter and antimatter appear the same. Sakharov in 1967 proposed a set of three necessary conditions for the period during which the Universe generated this baryon asymmetry.

1. Reactions that do not conserve net baryon number \(B\) (the number of baryons minus antibaryons) must be present.

2. The quantum theory of the Universe must violate both the symmetries \(C\) and \(CP\) (more details below).

3. The Universe must be out of equilibrium.

In this exercise, we shall give a cartoon discussion of “electroweak baryogenesis” [10], which generates this asymmetry as bubble walls sweep through the Universe during an abrupt phase transition.

First, we need to discuss important symmetries in quantum field theory. There are three symmetries of Schrödinger’s equation that we often take for granted. \(P\) (parity) is the symmetry inverting all positions through a point, \(x \rightarrow -x\). \(T\) (time reversal) is the symmetry inverting the direction of time (e.g., taking momenta \(p \rightarrow -p\)). And \(C\) (charge conjugation) replaces each particle with its antiparticle. It seems clear that breaking charge conjugation symmetry in Sakharov’s second condition is necessary to get more matter than antimatter. Parity is also important to Sakharov because of spin. We shall ignore the spin of the baryons in our model of the bubble wall (see note 19 on page 22).

It was a great surprise when it was discovered in 1956 by Chien–Shiung Wu that the weak interaction violates parity symmetry \(P\). (Her discovery was prompted by the theoretical suggestion that it had never been tested.) Thus Sakharov’s second condition is satisfied, even in our current Universe. Later, weak interactions were also shown to violate \(CP\). But there is a famous CPT theorem arguing that any Lorentz-invariant quantum field theory must have \(CPT\) invariance. That is, if one simultaneously inverts time, space, and swaps particles with their antiparticles, the laws of the Universe would be unchanged.

---

\(^{14}\)This exercise was developed in collaboration with Mitrajyoti Ghosh.

\(^{15}\)Protons and neutrons are baryons, as are all particles made up of three quarks. Three antiquarks make up antiprotons, antineutrons and other antibaryons.
We can treat $C$, $P$, $T$, and baryon number $B$ as quantum operators, like the Hamiltonian of the Universe $\mathcal{H}$. The CPT theorem tells us that $[CPT, \mathcal{H}] = 0$.

(a) How do $C$, $P$, and $T$ individually affect the baryon number operator? In particular, use $C^2 = T^2 = P^2 = 1$ to compute $CBC^{-1}$, $PBP^{-1}$, and $TBT^{-1}$ in terms of $B$. (Hint: One approach is to check their behavior applied to an eigenstate of baryon number $B|b\rangle = b|b\rangle$.)

We can now derive Sakharov’s third condition. You might think that the energy of a baryon is given by its mass, $mc^2$, so that in equilibrium the densities of baryons and antibaryons would be the same (since their masses are the same). But baryons interact in complex ways with the particles around them (think nuclear physics and the polarization of local vacuum fluctuations), and these are governed by rules that (for the weak interaction) violate $C$. Can we get a general proof that the equilibrium densities are the same?

An equilibrium Universe has a density matrix $\rho = \exp(-\beta \mathcal{H})$. The net baryon number is thus given by $\langle B \rangle = \text{Tr}(\rho B)$.

(b) Use CPT symmetry to show that $\langle B \rangle = 0$. (Hint: Use $(CPT)(CPT)^{-1} = 1$. Insert 1 into the trace, use the fact that $CPT$ commutes with the Hamiltonian, and use the cyclic invariance of the trace.)

Experiments in our current cold Universe have never observed baryon number violations. (Even a very small decay rate of protons to, say, a positron, photon, and antineutrino would cause a huge flood of $\gamma$ rays emanating from the Earth.)

In the early Universe, we believe there was an electroweak epoch when particles were massless (like the photon is now), and the four bosons $W^{\pm}$, $Z$, and photon were symmetric partners of one another. During this epoch when the electroweak symmetry is unbroken, the theory allows for a sphaleron reaction that transforms nine quarks (three baryons) into three antileptons (changing $B$, as Sakharov required). At the electroweak transition, the Higgs boson was created and many particles became massive, and the sphaleron reaction developed a high energy barrier (explaining why protons do not decay now).

The transition in electroweak baryogenesis is abrupt, and happens via the nucleation of rare bubbles that expand at nearly the speed of light—transforming the equilibrium unbroken phase with $\langle B \rangle = 0$ into a high-temperature version of the current Universe. Calculations show that the sphalerons are turned off almost completely in our current Universe (with breaking of the electroweak symmetry).

---

16 Do not confuse the density matrix $\rho$ with the baryon density $\rho$ in later parts of this exercise.

17 Sphalerons are non-perturbative solutions to the equations of motion of the quantum Hamiltonian of our universe. In particular, sphalerons are a minimum action solution to the field equations with a non-trivial topology. The order parameter space is $SU(2)$, and the field configuration takes the Universe (with topology $S^3$) into $SU(2)$. So the sphaleron can be viewed as a three-dimensional topological defect, whose strength is measured by the homotopy group $\Pi_3(SU(2))$ (given by non-equivalent mappings of the Universe $S^3$ into the order parameter space).
CHAPTER 1. EOPC: NEW EXERCISES

Fig. N1.9 Electroweak transition wall, sweeping at a high velocity \( v \), ending the symmetry-unbroken electroweak epoch and giving our particles masses. The wall interacts with the baryons (black) and antibaryons (white), letting those that impact the wall penetrate with probability \( T \). Deep inside the unbroken phase, sphalerons equilibrate the system and destroy any matter-antimatter asymmetry. Inside our world, sphalerons are inactive and baryon number is conserved.

This bubble wall is wildly out of equilibrium (Fig. N1.9). As it sweeps into the unbroken phase, it encounters baryons and antibaryons, but (because \( C \) is not a symmetry) the wall interacts differently with matter and antimatter. In particular, baryons are allowed to enter our universe (the broken-symmetry phase) with probability\(^{18} \), and antibaryons are allowed in with probability \( \overline{T} \). This leads to a pile-up of baryon density \( \rho_B(x) \) and antibaryon density \( \rho_{\overline{B}}(x) \) ahead of the front. The bubbles are huge, and so the bubble walls are nearly flat. We can thus study this pile-up as a one-dimensional problem, and change to a moving reference frame \( y = x - vt \) to find a time-invariant solution to the evolution laws.

We start in parts (c) through (f) by ignoring transitions between baryons and antibaryons, to work out the boundary conditions and the equations of motion in the moving reference frame. This means we can study \( \rho_B \) and \( \rho_{\overline{B}} \) independently. Let \( \rho_{\infty} \) be the density \( \rho_B(\infty) \) deep in the unbroken phase. Since the unbroken phase is presumed to be in equilibrium before the wall sweeps through,\(^{20} \rho_{\overline{B}}(x) \) will also be equal to \( \rho_{\infty} \) as \( x \to +\infty \).

(c) In steady state, ignoring any transitions changing baryons to antibaryons, can the density \( \rho_B(x) \) deep in our universe (as \( x \to -\infty \)) differ from \( \rho_{\infty} \)? (Hint: What is the current \( \mathcal{J}(y) \) in the moving reference frame?) Do the differing transmission probabilities suffice to explain the dearth of antimatter in our world?

Assume that baryons and antibaryons diffuse with diffusion constant \( D \). Let us begin

---

\(^{18}\)Warning: \( T \) is the probability of transmission, not the amplitude transmission constant \( T \) used in quantum mechanics and optics.

\(^{19}\)The spin of a massless particle becomes its helicity. So left-handed and right-handed photons carry circular polarization. If our quantum Hamiltonian had \( CP \) symmetry, left-handed massless antibaryons impinging from the unbroken symmetry phase would have the same transmission probability as right-handed baryons, leading to no net baryon number in our Universe. This is why Sakharov needs \( CP \) violation in addition to \( C \) violation.

\(^{20}\)That is, it is in metastable equilibrium, unstable only to our broken-symmetry universe.
by ignoring the sphaleron–induced reactions that change baryon number.

(d) Write the diffusion equation for baryons, and shift to the moving reference frame $y$, setting $y = 0$ to be the position of the bubble wall. Derive the evolution equation for $\hat{\rho}_B(y, t) = \rho_B(y + vt, t)$. Derive an ordinary differential equation for the stationary state, $\hat{\rho}_B(y)$ in the moving reference frame by setting $\partial \hat{\rho}_B(y, t)/\partial t = 0$. Note: $\hat{\rho}$ evaluated at $y = x - v(t + dt)$ is

$$
\hat{\rho}(y, t + dt) = \rho(y + v(t + dt), t + dt)
= \rho(y + vt, t) + v \frac{\partial \rho}{\partial x} \bigg|_{y+vt} dt + \frac{\partial \rho}{\partial t} \bigg|_{y+vt} dt.
$$

$$
\tag{N1.9}
$$

Fig. N1.10 Boundary conditions at electroweak wall. Near the wall, current conservation sets the boundary condition at the wall (eqn N1.10).

We also need to know the boundary condition at the wall $y = 0$. Consider a small region $0 < y < \delta$ on the unbroken symmetry side of the wall (Fig. N1.10). The net number of particles entering this region, divided by $\delta$, gives $\partial \hat{\rho}/\partial t$ just outside the wall. The current entering the region due to the moving reference frame is given by the velocity times $\hat{\rho}$. The current transmitted through the wall is the velocity times $T\hat{\rho}$ (the current impinging on the wall due to the moving reference frame, times the transmission fraction). Finally, the current exiting the region due to diffusion is $D$ times the slope.

(e) Why must the net current be zero for a stationary state? In the limit $\delta \to 0$, show that

$$
\frac{d\hat{\rho}_B}{dy} \bigg|_{0^+} = -(1 - T)(v/D)\hat{\rho}_B(0^+).
$$

$\tag{N1.10}$

Finally, argue that this boundary condition must hold even if the density has not reached a stationary state (and hence must hold even for the original evolution of $\rho(x, t)$.)
Show, in the absence of sphalerons, that the steady-state density of baryons is

\[ \hat{\rho}_B(y) = \rho_\infty (1 + (1/T - 1)e^{-(v/D)y}), \]  

(N1.11)

and hence \( \hat{\rho}_{\bar{B}}(y) = \rho_\infty (1 + (1/T - 1)e^{-(v/D)y}) \). (This should allow you to check your answers from part (c).)

Now let us add the sphaleron interaction which changes the net baryon number. For simplicity, let us ignore the leptons, and assume a baryon-nonconserving reaction that has an equal rate \( \Gamma \) for changing baryons to antibaryons and for changing antibaryons to baryons.

Write the coupled differential equations in the moving reference frame for the steady-states \( \hat{\rho}_B(y) \) and \( \hat{\rho}_{\bar{B}}(y) \). (For example, an extra term \( \Gamma (\hat{\rho}_B - \hat{\rho}_{\bar{B}}) \) is needed for \( \hat{\rho}_B \).) Add and subtract these equations to find uncoupled equations for \( \hat{\rho}(y) = (\hat{\rho}_B(y) + \hat{\rho}_{\bar{B}}(y))/2 \) and \( \delta \rho(y) = (\hat{\rho}_B(y) - \hat{\rho}_{\bar{B}}(y))/2 \). Ignoring the boundary conditions at zero, but applying the boundary conditions at \( y = \infty \), solve for the general solution of these coupled equations (retaining two unknown amplitudes). Show that \( \delta \rho(y) \propto \exp(-\Lambda y) \) decays exponentially, and determine \( \Lambda \).

If you have access and fluency in symbolic manipulation, you could now use the boundary conditions (eqn N1.10) at \( y = 0 \) for the baryons and the antibaryons to solve for the two unknown amplitudes in your equations. Using the densities at \( y = 0^+ \), you could then determine the transmitted densities. The algebra is a bit messy, so we are not asking you to do this. You would find in the end that the net baryon density in our Universe is predicted to be

\[ \rho_B - \rho_{\bar{B}} = \frac{2(\mathcal{T} - \overline{\mathcal{T}})(\sqrt{v^2 + 8D\Gamma} - v)\rho_\infty}{(\sqrt{v^2 + 8D\Gamma} - v)(\mathcal{T} + \overline{\mathcal{T}}) + 4\mathcal{T}\overline{\mathcal{T}}v}. \]  

(N1.12)

Find two choices of constants in the problem for which the baryon asymmetry is zero in our Universe. (You may assume \( D \) stays positive.) How do each of these relate to Sakharov’s conditions? What does the net baryon density become when \( \mathcal{T} = 0 \) (perfect exclusion of antibaryons)? Explain physically why this is to be expected.

At the time of the electroweak transition, there were many more baryons and antibaryons than there are now. Almost all the antibaryons were annihilated by baryons into photons, leaving the small density of baryons we have now. Various estimates for the original ratio suggest that it was less than \( 10^{-9} \) away from half and half, perhaps as small as \( 10^{-11} \).

Finally, one should note that the discovery of the Higgs boson produced a huge challenge for the electroweak baryogenesis. The sphaleron reaction rate \( \Gamma \) becomes very small when the Higgs mass becomes too high. As noted in part (g), the predicted net baryon density of our Universe vanishes as \( \Gamma \to 0 \) (when baryon number is conserved). Physicists used the current baryon number density and this feature of electroweak baryogenesis to predict a maximum value for the Higgs mass of 42 GeV. But the experimental mass turned out to be 125 GeV; the theory was not sufficient to explain the asymmetry. Where the antimatter went remains a mystery.
N1.9 Chiral waves: Fourier and Green. (3)

Recent research shows that many systems have chiral edge states—propagating modes in topological insulators and the quantized Hall effect that flow in only one direction. Such a system might be described by a field $\Xi(x,t)$ which evolves according to

$$\frac{\partial \Xi}{\partial t} = A \frac{\partial \Xi}{\partial x}. \quad (N1.13)$$

(see Exercise 9.13).

Our methods for solving the diffusion equation are most useful for equations which are both linear and translation invariant.

(a) Is the propagation law $N1.13$ linear? Is it translation invariant?

This equation can be solved by inspection, for any initial condition, as a traveling wave. In this exercise, we shall reproduce the traveling wave solution using more sophisticated methods. First, what is the traveling wave solution?

(b) Given an initial condition $\Xi_0(x) = \Xi(x,0)$, what will $\Xi(x,t)$ be? (Hint: Use parts (c) and (d) to check your answer.)

We can solve eqn $N1.13$ with Fourier transforms. Let $\hat{\Xi}_k(t)$ be the Fourier transform of $\Xi(x,t)$ with respect to $x$, using the conventions in the Appendix.

(c) Derive an equation for $d\hat{\Xi}_k/dt$. Solve it for $\hat{\Xi}_k(t)$ in terms of $\hat{\Xi}_k(0)$. Perform the inverse transform to evaluate $\Xi(x,t)$ in terms of $\Xi_0(x) = \Xi(x,0)$. Show your steps.

We can use the Greens function for eqn $N1.13$ to solve it.

(d) Use your general solution of part (b) to guess the Greens function $G(x,t)$ for eqn $N1.13$. Use the Greens function to evaluate $\Xi(x,t)$ for the initial condition $\Xi(x,0) = \Xi_0(x)$. Show your steps.

N1.10 Taste & smell with ensembles. (Biology) (3)

![Fig. N1.11 Receptor binding](image)

**Fig. N1.11 Receptor binding** in your nose or mouth leads to a signal that your brain interprets as taste or smell.

In Exercise 3.16, we modeled our sense of taste and smell in the microcanonical ensemble, and derived an expression for the signal involving the chemical potential. Using the grand canonical ensemble, we can now address this directly.

Figure N1.11 illustrates one of the ways our cells measure aspects of their environment—by placing receptors for given chemicals and measuring how often they are bound.
Model the receptor in the grand canonical ensemble, and assume it is in equilibrium with the fluid (air or saliva) surrounding it. Assume the fluid is at temperature $T$ and the chemical potential for the flavor molecules in the fluid is $\mu$. Let the energy of the receptor change by $\Delta E = E^B - E^U < 0$ as the receptor is bound, and let the entropy of the receptor change by $\Delta S = S^B - S^U$. Assume the volume in the system as a whole is unchanged when the flavor molecule binds, and for simplicity set $E^U = S^U = 0$.

Write the grand partition function $\Xi$ as a sum over the two states. What is the probability that the receptor is bound (and sending a signal to the brain)? Does your answer agree with your microcanonical answer, eqn 3.81?

N1.11 Entropy of Mastermind\textsuperscript{TM}.

Fig. N1.12 Mastermind\textsuperscript{TM} sample game, with two colors (B&O) and $N = 4$ pegs.

Mastermind is a game where a maker secretly picks a hidden code of $N$ colored pegs, and a breaker learns information about the positions and colors of the pegs through a sequence of guesses. After each guess, the maker responds with the number of guessed pegs which match in position and color (0r, 1r, \ldots, 4r), and the number of guessed pegs which share a color but are in the wrong position (0w, 1w, \ldots, 4w), as in Fig. N1.12.\textsuperscript{23}

\textsuperscript{21}To be fussy, we are summing over the bound and unbound Helmholtz free energies, which each sum over $e^{S/kB}$ states.

\textsuperscript{22}This exercise was developed in collaboration with Stephen Thornton.

\textsuperscript{23}You can view r and w as Right and Wrong, but they also match the red and white colors of the tiny pegs used in the response of the maker (Fig. N1.12).
In this exercise, we shall explore this game for the case of two colors, blue (B) and orange (O). Furthermore, instead of the breaker playing until the hidden code is matched, we shall play only one turn. The breaker’s job is to maximize the information about the hidden code after their guess, averaged over many games. The maker will eventually be allowed to choose a strategy optimized to frustrate the breaker, but for now we will assume that the maker chooses each of the possible codes with equal probability.

(a) If the breaker in a four-peg game guesses all blue and the response is 2r0w (Fig. N1.12), how many of the \(2^4 = 16\) possible hidden codes are still possible? What is the corresponding probability \(q_{20}\) that the response will be 2r0w, over many games where the breaker guesses all blue? How many hidden codes are still possible if BBBB yielded a response of 1r0w? 0r0w? 3r0w? 4r0w? What is the probability \(q_{rw}\) of getting each of these responses?

The breaker wins big in this case either if they guess correctly (4r0w) or if they guess completely wrong (0r0w) – they know precisely what the peg arrangement is. But how do we measure the information learned by guessing BBBB on average? Do we average the number of remaining peg arrangements over the possible responses? Do we do a weighted average of the number of remaining possible arrangements, based on the probability of getting each response?

In Section 5.3.2, we argued that the best way of measuring our ignorance is through entropy – the only measure that satisfies three key properties. (1) Entropy is maximum for equal probabilities. (2) Entropy is unaffected by extra states of zero probability. (3) Entropy changes properly for conditional probabilities.

(b) Which of the three properties is relevant for calculating the entropy of the breaker before the game starts? Which is relevant for calculating the entropy after they guess BBBB? Which tells us that we can also include legal but missing responses, like 2r1w for the guess BBBB?

(c) What is the starting entropy of the breaker, in bits? What is the breaker’s entropy \(S_{20}^{BBBB}\) after guessing BBBB and learning 2r0w? How many bits did the entropy decrease? What would their entropy \(S_{rw}^{BBBB}\) have been after each of the other possible responses \(rw = 00, 10, 30, \text{ and } 40\)? What is the average entropy \(S^{BBBB}\) averaging over all the responses, if all the hidden codes are equally likely? (Note: This last question is closely related to the discussion in Section 5.3.2 about your expected ignorance after asking your roommate where your keys were last seen.)

Now we must consider other possible initial guesses, to find the optimal choice. We can simplify our calculations by noticing some symmetries.

(d) Given that the maker picks codes with equal likelihood, can the average entropy after guessing BBBB and OOOO be different? Between BBBO and BOBB? Between BBBB and OBBB?

There is a permutation symmetry (reordering the pegs) and an inversion symmetry (B⇔O) in this problem; all initial guesses in a class related by these symmetries have the same entropy. This allows us to significantly reduce the number of breaker guesses
we need to calculate. To further reduce effort, we also reduce the number of pegs to \( N = 3 \).

The following table has two columns for breaker guesses and eight rows for the maker’s possible hidden codes (all equally likely).

<table>
<thead>
<tr>
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<th>BBB</th>
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<tr>
<td>BBB</td>
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<td>OBB</td>
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<td>BOO</td>
<td>2r0w</td>
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<td>OOB</td>
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<tr>
<td>OOO</td>
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</tbody>
</table>

(e) The other possibilities for breaker guesses should be related to the two shown. What is the multiplicity of BBB – the number of other breaker guesses in the same symmetry class? What is the multiplicity of BBO?

(f) Copy the table, and fill it in with the correct responses. Use the table to calculate \( q_{rw}^{BBB} \) and \( S_{rw}^{BBB} \) for the responses \( rw \) in column two, and \( q_{rw}^{BBO} \) and \( S_{rw}^{BBO} \) for column three. Calculate the average entropy after a guess in the BBB class and after one in the BBO class, as you did for BBBBB in part (c). Which will yield more information? (Hint: Use Fig. N1.13 to roughly check your answer.)

Now let us allow the maker to develop an optimal strategy to frustrate the breaker. What can we say about the optimal probability distribution choice for the breaker in this game? We assume that both breaker and maker can figure out the other’s optimal strategy. By playing a large number of games, any deviations from that optimal strategy can be detected and exploited.

Before the game begins, the maker’s choices have the same permutation and inversion symmetries as we saw for the breaker’s first guess.

(g) If the maker insists on a strategy that to break the symmetry – say, choosing BOB more often than OBB, can the breaker exploit knowing this to increase his knowledge after his guess? If the breaker insists on a strategy that guesses BOB more often than OBB, can the maker exploit it? Must the optimal strategies of the two satisfy the inversion and permutation symmetries?

We shall presume, whatever your answer for part (g), that the maker chooses BBB and OOO each with probability \( p \leq \frac{1}{2} \), and all other codes with probability \( 1/6 - p/3 \). Since the breaker know of their strategy and choice of \( p \), the breaker’s initial entropy changes.

(h) Calculate the breaker’s initial entropy as a function of \( p \), before they make a guess. If the maker chooses \( p \) near \( \frac{1}{2} \) and the breaker knows this, will the breaker’s optimal strategy change from the one you calculated in part (f)?
Since the maker’s strategy maintains the same permutation and inversion symmetries as the equal-weighted strategy, we can use the same table to calculate the breaker’s entropy as a function of $p$.

The table of codes, guesses, and responses now includes a column for the probabilities. The missing entries (unchanged from the earlier table) are filled in for one particular response $r = 2, w = 0$ for an initial guess of $BBO$.

<table>
<thead>
<tr>
<th></th>
<th>BBB</th>
<th>BBO</th>
</tr>
</thead>
<tbody>
<tr>
<td>$p$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$1/6 - p/3$</td>
<td>BBO</td>
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<tr>
<td>$1/6 - p/3$</td>
<td>BOB</td>
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<tr>
<td>$1/6 - p/3$</td>
<td>OBB</td>
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<tr>
<td>$1/6 - p/3$</td>
<td>BOO</td>
<td>2r0w</td>
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<tr>
<td>$1/6 - p/3$</td>
<td>OBO</td>
<td>2r0w</td>
</tr>
<tr>
<td>$1/6 - p/3$</td>
<td>OOB</td>
<td></td>
</tr>
</tbody>
</table>

Calculating the resulting average entropy after the two guesses is now both laborious and similar to your calculation in part (f). The particular response 2r0w for BBO, however, involves three outcomes with differing probabilities, so is worth investigation.

(i) What is the net probability for the response $q_{BBO}^{20}$ as a function of $p$? What is the entropy $S_{BBO}^{20}$ of the breaker after he gets that response to a BBO guess? In terms of these two quantities (without plugging in your formulas), what is the contribution of the 20 response to the average entropy of the breaker after the guess? Plot $S_{BBO}^{20}$ versus $p \in (0, \frac{1}{2})$.

Figure N1.13 shows the uncertainty of the breaker after the two possible classes of guesses. The entropy $S_{BBO}^{20}$ you calculated in part (i) is a necessary step in calculating $S^{BBO}$ in the plot below.

**Fig. N1.13 Breaker uncertainty** entropies, after one guess in Mastermind™.
In non-cooperative games, each player tries to choose from an ensemble of strategies that best optimizes their outcome. The breaker’s best strategy, in general, depends on the maker’s strategy (here the choice of $p$), and vice versa. Mathematician John Nash\(^{24}\) won the Nobel Prize in Economics for showing that such games will generally have a *Nash equilibrium*, when neither player has anything to gain by changing their strategy.

(j) **What is the optimal strategy for the breaker, for low $p$ and high $p$? Use Fig. N1.13 to estimate approximately at what value of $p$ the breaker’s strategy changes. Assuming the breaker is using the optimal strategy, estimate approximately at what value of $p$ the maker should use to frustrate the breaker as much as possible.** This combination of strategies is the Nash equilibrium for our game.

N1.12 **Interpolation and free energies.**

Alemi and Fischer [2] have unified a large number of deep neural network models into what they call TherML (Thermodynamics of Machine Learning). They combine four quantities $R$, $D$, $C$, and $S$ that various algorithms attempt to minimize into a single optimization goal

$$R + \delta D + \gamma C + \sigma S,$$

(N1.14)

weighing the importance of minimizing $R$, $D$, $C$, $S$ differently depending on the task. This sum is then minimized with respect to millions of parameters in the neural network, to achieve the user’s goals. Here $C$ measures the classification error, $R$ the complexity of the representation, $S$ the relative entropy in the parameters of the model, and $D$ the distortion measuring the ‘unsupervised’ learning performance. This looks remarkably like our free energy formulas like the grand free energy $\Phi = E - TS + PV$ and the energy $E = TS - PV + \mu N$. Note that $E$ weighs the relative ‘goals’ of $S$, $V$, and $N$ by their costs $T$, $-P$, and $\mu$.

\(^{24}\)This is the same John Nash that was featured in “A Beautiful Mind”.

---

**Fig. N1.14 Interpolations** for small step sizes usually have their maximum error near midway between fitted points.
Without pretending to explain these machine-learning terms, let us pursue a more tangible task with the same form: interpolating a function using polynomials (Fig. N1.14). Our goal is to minimize the error $E$ in the interpolation by varying computer time $T$ and storage $N$.

Consider approximating the periodic function $\sin(x)$ for $x$ in $(0, 2\pi)$ by linearly interpolating between $N + 1$ values at $x_n = n\epsilon$ with $\epsilon = 2\pi/N$ (Fig. N1.14). In each interval, the maximum error will approximately be in the center $x$ between the known values at $x \pm \epsilon/2$, where it is $|\sin(x) - \frac{1}{2}(\sin(x - \epsilon/2) + \sin(x + \epsilon/2))|$.

(a) Expanding about $x$ to second order in $\epsilon$, at what $x$ is the error biggest? Show that the maximum error $E$ goes as $E = A/N^2$ as $N \to \infty$. For $\sin(x)$, what is $A$? How many digits of accuracy $D(N) = -\log_{10}(E)$ will linear interpolation provide?

We will consider polynomial interpolation methods, that fit the nearest $M$ points separated by $\epsilon$ with a polynomial of degree $M - 1$. For large $N$ (small $\epsilon$), these fits match the first $M$ terms of the Taylor expansion about $x$, so their errors involve $\epsilon^M$ times the $M$th Taylor coefficient (which depends on the function being interpolated). For simplicity, we shall ignore this dependence, so the error $E \propto N^{-M}$ and the number of digits $D = -\log_{10} E \propto M \log(N)$. (You may check that this agrees with your answer to part (a)).

The time needed to evaluate this polynomial scales as $T \propto M^2$ [27, Section 3.2]. For convenience, we shall assume time $M = \sqrt{T}$ and digits $D = M \log N = \sqrt{T} \log N$ (by choosing suitable units).

We are willing to pay an amount $\chi$ per digit of accuracy. Let $\sigma$ be the cost of a unit of time, and $\mu$ be the cost of a unit of memory.

We can view the digits $D(N, T)$ as an analog of the microcanonical ensemble, where the amount of storage $N$ and the computer time per evaluation $T$ are fixed.

(b) How does the number of digits $D(N, T)$ increase as we increase $N \to N + dN$? Starting at $(N, T)$, how much are we willing to pay to increase $N \to N + dN$? How much would we pay for more computation time, $T \to T + dT$? Evaluate $\mu(N, T)$ and $\sigma(N, T)$, this ‘marginal cost’ of additional storage and time. Give a general Maxwell relation giving $(\partial \mu(N, T)/\partial T)|_N$ as a derivative of $\sigma$, and check it using your formulas for $\mu$ and $\sigma$.

So our general goal is to maximize

$$G(\chi, \sigma, \mu) = \chi D - \sigma T - \mu N$$

with respect to $T$ and $N$, in analogy with TherML’s optimization in eqn N1.14. Here $D$ is a function of $N$ and $T$. $G$ changes variables from $D$, $N$, and $T$ just as the Gibbs free energy $G(T, P)$ changes variables from the microcanonical entropy $S(E, V)$. There $1/T$ was analogous to the cost in entropy of buying a unit of energy, and $P/T$ analogous to the cost of a unit of volume. Here $\sigma$ and $\mu$ are literally the cost in dollars for buying computer time and memory.
Minimizing $G$ tells us that we have increased $N$ and $T$ until it costs $\chi$ per digit to make further improvements.

(c) Suppose we start at some non-optimized $N$ and $T$, and make a small change $(\Delta N, \Delta T)$ that produces a change $\Delta D$ in the digits of precision. How much are we willing to pay to benefit by the increased precision? How much does the change cost? Does the place where $G$ is minimized make the total cost equal to the total benefit? Or something else?

Let us imagine we have solved for $G$. What could we do with it? Just as derivatives of the Gibbs free energy $dG = -SdT + VdP + \mu dN$ allow us to write the entropy and the volume as functions of temperature and pressure, so our $G(\chi, \sigma, \mu)$ allows us to budget the computer time and storage in terms of their costs $\sigma$ and $\mu$ and the value per digit $\chi$. So, before plunging into the calculation of $G$...

(d) If you were planning a simulation, how would you use $G$ to determine the memory and computer time to budget, and the resulting number of digits you should purchase?

Now we calculate $G$.

(e) Extremize $G$ in eqn N1.15 with respect to $N$ to find a formula for $\mu$. Extremize with respect to $T$ to find a formula for $\sigma$.

You will need a new special function to solve for $G$. Let $w(A)$ be the solution to the equation $\log(w) = Aw$.\(^{25}\)

(f) Solve both formulas from part (e) for $\sqrt{T}$. Equate the solutions to give a self-consistent formula for $N$. Using the Lambert $W$ function to solve for $N$. Using one of your two formulas for $T$ to write $T$ as a power of $N$, and through that as a function of $\chi$, $\mu$, and $\sigma$.

(g) Let $\sigma = 3$ and $\mu = 0.1$. Plot $N$, $T$, and the number of digits for the range $\chi \in (3, 6)$. (Hint: $T$ should start near four, corresponding to the linear interpolation $M = 2$ in part (a).) How much does the order $M$ change as we add digits?

In principle, we could now combine this information into eqn N1.15 to get an explicit (but horrible) formula for $G$.

\(^{25}\)This can be written in terms the $-1$ branch of the Lambert $W$ function, $w(A) = -W(-A, k = -1)/A$. See the hints files [31] for implementations in Mathematica and Python.

N1.13 Convexity and phase separation. (Thermodynamics) 3

A piston, initially completely filled with water vapor, compresses the gas until it is completely liquid, moving between the marked positions in Fig. N1.15. The piston is held at temperature $T = 550K$ at all times.
Fig. N1.15 Pressure vs. volume for the van der Waals model applied to one mole of H₂O at T = 550K. The red line shows the vapor pressure $P_v$ at this temperature.

(a) Draw the path on Fig. N1.15 taken if the piston moves slowly enough that the system remains in thermal equilibrium at all times.

(b) Sketch a path on Fig. N1.15 taken if the piston moves fast enough so that the pressure rises past the vapor pressure (say, to $1.2 \times 10^8$ dynes/cm²) before a liquid water drop nucleates, but slowly enough so that the subsequent condensation of vapor into the water stays in equilibrium.

Figure N1.16 shows the chemical potential for the van der Waals model as a function of density at this same temperature. Remember that the Gibbs free energy is $\mu N$, so this is also the Gibbs free energy per molecule. Note that the van der Waals solution assumes that the system is filled with molecules at a uniform density $\rho$, not a mixture of liquid and gas.

Fig. N1.16 Chemical potential $\mu$ vs. density $\rho$ for the van der Waals model for H₂O at T = 550K.

(c) Sketch on a copy of Fig. N1.16 the free energy one would obtain by allowing for the separation of the water into coexisting liquid and gas. (Ignore the small contribution of
surface tension.)

If a system can be broken up into two weakly interacting subsystems, then the minimum free energy for the system in the limit of infinite size must be convex (see note 4 on page 324).

(d) In your solution to part (c), what are the two weakly interacting subsystems? Why did we need to take the limit of infinite size to ignore surface tension? Is your answer convex?

N1.14 Spinodals vs. Nucleation.

Here we explore the predicted edges of metastability for the liquid and gas, which are called spinodals in the older literature. Spinodals provide some insight into the behavior of materials near abrupt transitions, but should not be taken literally—fluctuations ignored by models like these cause the crossover from nucleation to ‘spinodal decomposition’ to become blurred.

Let us explore how the Gibbs free energy per particle \( G/N = \mu \) (Fig. N1.18) varies as we move between different points on the \( P-V \) diagram Fig. N1.17.

(a) At point \( \alpha \) in Fig. N1.17 at the highest pressure, how many other solutions are there with that pressure? Which of the labeled points in the free energy plots of Fig. N1.18 corresponds to the state \( \alpha \)?
Fig. N1.18 Chemical potential vs. density at fixed temperature $T = 550$ K, at some of the pressures corresponding to the points $\alpha - \zeta$ in Fig. N1.17. The labeled points $a-j$ here are solutions to the van der Waals model.

(b) Is the system at point $\delta$ in Fig. N1.17 stable, unstable, or metastable? Which point $a-j$ in Fig. N1.18 corresponds to $\delta$? Explain what about the free energy tells you whether a state is stable, unstable, or metastable.

We gradually compress a hot water vapor in a vertical piston, keeping the temperature at 550 K. We first study this in the $P-V$ plane and in terms of the chemical potential.

(c) At which labeled point in Fig. N1.17 does the metastable gas state go unstable? This is called a spinodal point. Which point $a-j$ in Fig. N1.18 corresponds to this spinodal point for the gas phase? In Exercise 11.3, we saw that the surface tension energy for nucleating a bubble is related to the barrier in free energy between the two phases. At the spinodal point, does the free energy barrier between the two phases disappear?

If one quickly changes the temperature or volume or other parameter across an abrupt phase transition line, the phase transition qualitatively happens in one of two ways. It can be nucleated by slow-forming bubbles or droplets. Or, if one moves into the unstable region, it can undergo spinodal decomposition, spontaneously separating into two phases without a nucleation barrier, with small random density fluctuations growing on many length scales. The boundary between these two is blurry except in mean-field theories like that of van der Waals.
The three points $\kappa$, $\lambda$, and $\mu$ in Fig. N1.19 correspond to our compression experiment in part (c). We prepare the state at $\lambda$ and $\mu$ by starting at state $\kappa$ at $t = 0$ and compressing the piston quickly while cooling to keep the temperature $T$ fixed. The states at $\lambda$ and $\mu$ begin in a uniformly compressed state with only microscopic fluctuations. Assume the system is in very low gravity (so the liquid water, when it forms, does not quickly fall to the bottom). For each, consider the density $\rho_\lambda(h,t)$ and $\rho_\mu(h,t)$ along a one-dimensional line $h$ rising from the bottom of the piston to the top.

(d) Roughly sketch $\rho_\lambda(h,t)$ as a function of $h$ for three times: $t = 0$ just after compression, a time $t_{\text{droplets}}$ when the line passes through a nucleated droplet or two (which have yet to grow to use up the extra water vapor, or to fall to the bottom), and a much later time $t_{\text{equilib}}$ when it is fully phase separated. (Hint: What fraction of the volume will be liquid?) Label your density axis with the equilibrium densities of the liquid and gas at 550 K, and with the initial density $\rho_\lambda$.

(e) Roughly sketch $\rho_\mu(h,t)$ as a function of $h$ at a time $t_{\text{spinodal}}$ when tiny random thermal density fluctuations have been noticably magnified by the instability, but have yet to approach the equilibrium densities of the two phases. Label your density axis with the equilibrium densities of the liquid and gas at 550 K, and with the initial density $\rho_\mu$.

After evolving for long times (ignoring gravity), a system which was launched quickly with spinodal decomposition or slowly by nucleation will evolve into rather similar states which coarsen with time.

N1.15 **Cell signaling and mutual information.** (Biology, Statistics) ③

To survive, living systems must accurately measure and respond to their environment. For example, *E. coli* bacteria famously execute a process called chemotaxis. By sensing gradients in nutrients, they decide whether to run (propel themselves forward) or tumble (change direction) using their propellor-like *flagella* (see Exercises 2.19 and 2.22). The environmental signal is transmitted from the receptors on the cell surface to the motors that control the flagella via a cascade of chemical reactions. These

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③This exercise was developed in collaboration with David Hathcock.
cascades involve kinases—protein enzymes that are activated (phosphorylated) by reactions that are catalyzed by upstream enzymes (the cell membrane receptors or another kinase). The activated kinase has a finite lifetime, with dephosphorylation reactions catalyzed by so-called phosphatase enzymes. Fluctuations in active kinase populations carry the signal through the cascade. Multi-level cascades can amplify signals, enabling sensitive detection of the environment, but also introduce noise due to the stochastic nature of enzymatic reactions. In this problem we will investigate how cells can mitigate this noise, optimizing the information transmitted by the signaling circuit.

![Diagram of a cellular signaling circuit](image)

**Fig. N1.20 The simplest cellular signaling circuit** involves: (i) the environmental signal from extracellular ligands X, (ii) an kinase population Y within the cell, activated by the membrane receptors. This minimal model is a coarse-grained representation of actual chemical circuits that regulate chemotaxis in *E. coli*, olfactory (smell) sensing in mammals, and yeast response to osmotic pressure [30, 14].

Let $X(t)$ be the local concentration of food as the bacteria swims forward. This concentration randomly fluctuates, driven by noise $\xi_x$ due to both to the bacteria’s movement and coupling to the environment, but changes relatively slowly, with a long correlation time $\gamma_x^{-1}$. The dynamics of $X$ are given by a Langevin equation, $\dot{X} = A_x - \gamma_x X + \xi_x(t)$, where $A_x$ keeps the mean food level $\overline{X} = A_x/\gamma_x$ positive. The cell’s measurement of this signal is noisy: small numbers of food molecules randomly bump into the receptors, which in turn causes activation of $R$ kinase within the cell. The dynamics of active kinase population $Y$ are given by, $\dot{Y} = A_y + R(X - \overline{X}) - \gamma_y Y + \xi_y(t)$, where $\xi_y$ is the measurement noise, $A_y$ is the background activation rate, and $\gamma_y$ is the deactivation rate, determined by phosphatase concentrations. As with $X$, $\gamma_y^{-1}$ sets the time-scale for fluctuations in $Y$. How can the cell tune $\gamma_y$ to filter out the measurement noise at the receptors and gain the most information about the food in its environment?

For our calculations, we will assume the terms $\xi_a(t)$ with $a = x, y$ are white noise with correlation\(^\text{27}\) $\langle \xi_a(t)\xi_b(t') \rangle = 2F_a\delta_{ab}\delta(t - t')$. Information is carried by the fluctuations in $X$ and $Y$ about the mean values, $\overline{X} = A_x/\gamma_x$ and $\overline{Y} = A_y/\gamma_y$; the bacteria wants to swim toward locations where there is more food than average. Subtracting off the

\(^{27}\text{If the number of molecules attaching to the receptor is the cause of the noise (this is called shot noise), then } F_x = A_x \text{ and } F_y = A_y.\)
means, \( x = X - \overline{X} \) and \( y = Y - \overline{Y} \), we are left with,

\[
\dot{x} = -\gamma_x x + \xi_x(t) \quad \dot{y} = -\gamma_y y + R x + \xi_y(t) \tag{N1.16}
\]

The calculations in this exercise are somewhat involved, so we shall provide many intermediate results to allow you to bypass parts if you get stuck. If you use a computer algebra system (Mathematica, SymPy, …) please provide your code. If you use paper and pencil, please show your steps and prune your dead ends.

(a) Fourier transform eqn (N1.16) and solve for \( \tilde{x}(\omega) \) and \( \tilde{y}(\omega) \). Express your answers in terms of the Fourier transformed noise \( \tilde{\xi}_x(\omega) \), \( \tilde{\xi}_y(\omega) \) as well as \( \gamma_x \), \( \gamma_y \), and \( R \).

If \( C_{ab}(\tau) = \langle a(t + \tau)b(t) \rangle \) is the correlation between \( a \) and \( b \), then \( \tilde{C}_{ab}(\omega) = \tilde{a}(\omega)\tilde{b}^*(\omega) \) (see Appendix A). For example, the noise correlation relation becomes \( \tilde{\xi}_a(\omega)\tilde{\xi}_b^*(\omega) = 2F\tilde{C}_{ab}(0) \) and the equal time correlation is \( C_{ab}(0) = 1/(2\pi) \int d\omega \tilde{a}(\omega)\tilde{b}^*(\omega) \). For the next parts, the following integrals are useful:

\[
\begin{align*}
\frac{1}{2\pi} & \int_{-\infty}^{\infty} e^{-i\omega t} d\omega = \frac{e^{-ct}}{2c} \\
\frac{1}{2\pi} & \int_{-\infty}^{\infty} \frac{d\omega}{\omega^2 + c^2} = \frac{1}{2c(c + d)} \\
\frac{1}{2\pi} & \int_{-\infty}^{\infty} \frac{d\omega}{\omega^2 + (c^2 + d^2)} = \frac{1}{2cd(c + d)}
\end{align*}
\]

(b) Show that \( x(t) \) has exponential time correlations, \( C_{xx}(\tau) = F_x/\gamma_x \exp(-\gamma_x |\tau|) \). As we mentioned above the signal has a correlation time scale set by \( \gamma_x^{-1} \).

(c) Show that the equal time correlations are \( C_{xx}(0) = F_x/\gamma_x \), \( C_{xy}(0) = F_x R/(\gamma^2_x + \gamma_x \gamma_y) \) and \( C_{yy}(0) = F_x R^2/(\gamma^2_x \gamma_y + \gamma_x \gamma^2_y) + F_y/\gamma_y \). To simplify the notation below, we will define \( \sigma^2_x = C_{xx}(0) \), \( \sigma^2_y = C_{yy}(0) \), and \( \text{cov}_{xy} = C_{xy}(0) \).

So far, we have understood the fluctuations and correlations in two chemical populations: food molecules in the environment and signaling proteins within the cell. How can we use these to quantify the information gained by the bacteria? Fortunately, Shannon explored this question in his seminal work on information theory. He developed the machinery of information entropy when working for the telephone company.\(^{28}\)

The phones in those days had lots of static. How could one calculate when the static noise made it impossible to hear the words? Shannon introduced the idea of a \textit{mutual information} between the original signal and the noisy final signal.

Suppose the probability distribution of an original message \( x \) is \( \rho_X(x) \), and the probability distribution of \( x \) being turned into a noisy final message \( y \) is \( \rho_T(y|x) \) (the probability of \( y \) given \( x \)). This transmitted probability distribution can be written in terms of the joint probability distribution \( \rho_{X,Y}(x,y) \), the likelihood of a random transmission having

\(^{28}\)This was in the olden days, when there was only one phone company in the US, Bell Telephone.
original message \( x \) and final noisy message \( y \). Note that \( \rho_X(x) = \int dy \rho_{X,Y}(x,y) \) and similarly for \( \rho_Y(y) \).

(d) Express in words the true statement \( \rho_T(y|x)\rho_X(x) = \rho_{X,Y}(x,y) \) in a way that would convince a non-scientist.

The mutual information\(^{29} \) is defined to be

\[
\mathcal{I}(X, Y) = k_S \int dx \int dy \rho_{X,Y}(x,y) \log \left( \frac{\rho_{X,Y}(x,y)}{\rho_X(x)\rho_Y(y)} \right),
\]

The mutual information is symmetric in \( X \) and \( Y \). How can it be used to study how much the noisy final signal \( y \) tells us about the original signal \( x \)?

(e) Show that \( \mathcal{I}(X, Y) = S(X) - S(X|Y) \), the entropy of the probability distribution of general noisy input signals \( \rho_X(x) \) minus the entropy of the noisy signals resulting in a particular received signal \( y \), \( \rho_T(x|y) \), averaged over \( y \) (Hint: the identity from part (d) also holds if we swap \( x \) and \( y \). Plug this in and separate the log into a difference of two terms). Explain in words why the first measures your ignorance of the message \( X \) before getting a telephone call \( Y \), and the latter is your average ignorance after a call.

To compute the mutual information for the cell signaling system, we require the joint distribution \( \rho_{X,Y}(x,y) \). In general, for systems driven by white-noise (with delta-function correlations), this is done by converting the Langevin equations into a Fokker-Planck equation (see Exercise 8.22), which can then be solved to obtain the equilibrium distribution. When the equations are linear (as is the case for our model, eqn (N1.16)), one finds that the joint distribution is a bivariate Gaussian\(^{30} \),

\[
\rho_{X,Y}(x,y) = \frac{1}{2\pi\sigma_x\sigma_y\sqrt{1-r^2}} \exp \left( -\frac{1}{2(1-r^2)} \left( \frac{x^2}{\sigma_x^2} + \frac{y^2}{\sigma_y^2} - \frac{2rxy}{\sigma_x\sigma_y} \right) \right).
\]

Here, the standard deviations \( \sigma_x = \langle x^2 \rangle^{1/2} \) and \( \sigma_y = \langle y^2 \rangle^{1/2} \) and correlation \( r = \langle xy \rangle/(\sigma_x\sigma_y) = \text{cov}_{xy}/(\sigma_x\sigma_y) \) are precisely those you computed in part (c).

(f) Argue that the marginal distributions \( \rho_X(x) \) and \( \rho_Y(y) \) are also Gaussian, with widths \( \sigma_x \) and \( \sigma_y \) respectively. Show that the mutual information for the bivariate Gaussian is \(-k_S/2 \log(1-r^2)\). Interpret this result for \( r \to 0 \) and \( r \to 1 \). Is the mutual information dependent on the magnitude of the fluctuations? (Hint: what happens if we scale \( x \to \alpha x \)).

\(^{29}\)The mutual information is also the distance in probability space between the joint distribution \( \rho_{X,Y}(x,y) \) and the uncorrelated case \( \rho_X(x)\rho_Y(y) \), using a distance measure called the Kullback-Liebler divergence (see Exercise 1.16).

\(^{30}\)This is a generalization of a familiar result from statistical mechanics: the Boltzmann distribution for an over-damped harmonic oscillator (which feels forces linear in the displacement) is Gaussian.
Biologically, the rate $\gamma_y$ is easiest to tune (e.g. through evolution and natural selection). This rate is controlled by concentrations of phosphatase in the cell, whereas the activation rate $R$ requires energy dissipation.

(g) Argue that maximizing mutual information is equivalent to maximizing $r^2$. Show that $r^2 = \frac{Z\Lambda}{((1 + Z)(1 + Z + \Lambda))}$, where $Z = \frac{\gamma_y}{\gamma_x}$ and $\Lambda = \frac{F_x R^2}{(F_y \gamma_x^2)}$. Maximize $r^2$ with respect to $\gamma_y$. Show that when the information transmission is optimized, $\gamma_y = \frac{\gamma_x \sqrt{1 + \Lambda}}{1}$ and $I = \frac{1}{2} k_S \log(1 + \sqrt{1 + \Lambda}) - 1$.

With all else fixed, there is an ideal response time: $\gamma_y^{-1}$ must be faster than $\gamma_x^{-1}$ by the factor $1/\sqrt{1 + \Lambda}$. If $\gamma_y$ is too small, $Y$ will integrate over the changes in $X$, filtering out too much of the signal. On the other hand, if $\gamma_y$ is too large, $Y$ will fail to filter out enough of the noise introduced at the cell receptors. The mutual information quantifies how much information the cell has about the current state of its environment, obtained by measuring the history of the signal and filtering out receptor noise. The dimensionless parameter $\Lambda$ serves as a measure of the signal fidelity. When $F_x \gg F_y$, then $\Lambda \gg 1$: it is easy to reproduce the signal if the magnitude of fluctuations in $X$ are large compared to the noise introduced by the receptors. On the other hand if $F_x \approx F_y$, $\Lambda = \frac{R^2}{\gamma_x^2}$, which measures the sensitivity of $Y$ to changes in the $X$ population over the timescale $\gamma_x^{-1}$. Real biological signaling circuits, like those in yeast and E. coli, tend to lie in the range $\Lambda = 100 - 1000$.

(h) For the range of $\Lambda$ listed above, how much information, in bits, can the cell learn about its environment from monitoring $Y(t)$?

N1.16 Emittance and particle beams.\(^{31}\) (Accelerator) \(^{3}\)

Particle accelerators take bunches of protons and antiprotons up to near the speed of light, and smash them head-on to see what happens. Electron microscopes take bunches of electrons and focus them to image materials at the atomic scale. X-ray sources accelerate electrons to near the speed of light, and use undulators to wiggle them to create X-rays (synchrotron X-ray sources) or free electron lasers (coherent beams of X-rays). In all of these applications, in addition to the energy per ion and the number of ions, the key property of a good bunch is its emittance. In this exercise, we shall explain why emittance is important, relate it to the entropy of the bunch, analyze its quantum limit, and explore the use of electron bunches to cool bunches of protons.

The 3D emittance of a bunch is loosely given\(^{32}\) by the product of the volume it occupies in position space and in momentum space:

$$\varepsilon = (\Delta q_x \Delta p_x)(\Delta q_y \Delta p_y)(\Delta q_z \Delta p_z).$$

\(^{31}\)This exercise was developed in collaboration with Michael Kaemingk. We have used real numbers as input for this exam, but our assumptions for the calculations are not reliable. For example, real bunches are not as tidy as our Gaussian model bunches, and their momentum spread is not thermal.

\(^{32}\)In part (a) you shall derive another more rigorous but less practical definition of the emittance in terms of the entropy per particle.
For our purposes, $\Delta q$ and $\Delta p$ will represent the standard deviations of the positions and momenta in the center of mass frame of the bunch. Here $z$ is the ‘longitudinal’ direction in which the beam is moving, and $x$ and $y$ are ‘transverse’. For synchrotrons, $y$ is the ‘vertical’ direction perpendicular to the plane of the circular orbit and $x$ the ‘horizontal’ direction in the plane of the circle but perpendicular to the motion. One also speaks of the 2D emittance of the transverse directions $(\Delta q_x \Delta p_x)(\Delta q_y \Delta p_y)$ perpendicular to the velocity of the bunch, or the 1D emittance along one of the axes. We shall see that many beams are strongly anisotropic, with different widths and momentum spreads along these three axes.

Emittance is a limiting parameter in the performance of any accelerator. In electron microscope/diffraction accelerators, the emittance limits the resolution; in colliders, it limits the luminosity; in free electron lasers, it limits the gain length and the minimum wavelength of the laser. Low beam emittance is therefore desirable, and reducing it is a central research goal of the Center for Bright Beams (CBB), a collaboration in which Cornell plays a leading role.

At the Large Hadron Collider (LHC), the bunches have $N = 1.2 \times 10^{11}$ protons, a bunch radius of $\sigma_q = 3.5 \mu$m, and a transverse emittance of $3.75 \mu$rad $= 1.88 \times 10^{-24}$ kg m$^2$/s, implying a RMS bunch momentum of $\sigma_p = 5.4 \times 10^{-19}$ kg m/s.

The ions in a bunch are often nearly noninteracting and uncorrelated, with all ions having nearly the same probability distribution in phase space. In this case, $\rho_N(\mathbb{P}, \mathbb{Q}) = \prod_{n=1}^N \rho(p_n, q_n)$. In this exercise, our bunches will mostly have Gaussian distributions. Here we very roughly approximate the LHC bunch as an isotropic, uncorrelated product of spherically symmetric Gaussian distributions

$$
\rho_{\text{LHC}}(p, q) = \frac{e^{-q^2/2\sigma_q^2}e^{-p^2/2\sigma_p^2}}{(2\pi\sigma_q^2)^{3/2}(2\pi\sigma_p^2)^{3/2}}.
$$

(a) Write the formula for the 3D emittance of the bunch in eqn N1.19. Note that the momenta have a thermal distribution. Write a formula for the temperature of the bunch. Write the entropy

$$
S = -k_B \langle \log \rho \rangle = -k_B \int \rho \log \rho
$$

$$
= -k_B \int d\mathbb{P} d\mathbb{Q} \rho(\mathbb{P}, \mathbb{Q}) \log \rho(\mathbb{P}, \mathbb{Q})
$$

$$
- 3Nk_B \log h.
$$

(Warning: There is an error in the corresponding eqn 5.20 in the print edition of the text.) Evaluate the temperature and the entropy of the bunch using the LHC parameters above. Write a formula for the emittance in terms of $\exp(S/Nk_B)$. We may view your last result as a more rigorous definition of the emittance.
Thus, the emittance, like the entropy, can only grow as the bunch passes through accelerating fields and focusing magnets. Making a low emittance bunch, and keeping the emittance low during its acceleration and focusing, is key to all accelerator applications. Why do we care about the momentum spread $\Delta p$ in eqn N1.19, if we want a dense beam or a sharp focus? The conservation of emittance forces a tradeoff between a narrow beam and one that stays narrow as it moves.

The angular dispersion of a beam is due to the momentum spread $\sigma_p$: the momentum in the transverse direction will make the beam grow in width. The angular spread in a beam is given by the ratio of the transverse momentum spread $\Delta p$ over the mean momentum of the ions in the forward direction. The latter, $mv$ for nonrelativistic motion (with $m$ the particle mass) becomes $mv/\sqrt{1-v^2/c^2} = \beta \gamma mc$ for a relativistic beam, with $\beta = v/c$ and $\gamma = 1/\sqrt{1-v^2/c^2}$. Thus the angular spread $\Delta \theta = \Delta p/\beta \gamma mc$ radians with respect to the direction of motion.

(b) An electron microscope has a beam with velocity $v = 0.62c$, a width $\sigma_q^i = 200 \mu m$ and a vertical emittance $\varepsilon = 2.7 \times 10^{-30} kg m^2/s$. What is its vertical momentum spread $\sigma_p$? What is its angular spread $\Delta \theta_i$ in radians? A lens system focuses the beam into a smaller width 40 $\mu m$, without increasing the entropy or changing its velocity. What is the new angular spread $\Delta \theta_f$, in terms of $\Delta \theta_i$? How far can it propagate before the new spread gives the beam a width larger than the original width $\sigma_q^i$? (Rough estimates are fine.)

One might be surprised that the ions in a bunch can be treated as non-interacting, given the strong Coulomb interactions between particles. Indeed, these ‘space-charge’ effects are important when the bunch is first formed. But as it reaches near the speed of light, these interactions become much less important. Using an ultrafast electron diffraction apparatus at Cornell as an example, the current state-of-the-art photoemission bunch starts out as a bunch of radius 20 $\mu m$ in the transverse directions, and a duration of 200 femtoseconds. Suppose this beam is now placed in an X-ray Free Electron Laser and accelerated to a speed very close to that of light (but leaving its duration fixed). The packet in the laboratory frame is 60 $\mu m$ long (roughly spherical). But the packet in the laboratory frame is Lorentz contracted by a factor of $\gamma \approx 34000$, so in the center of mass frame it is a long tube of length $\gamma \times 60 \mu m = 1.3 m$. The force on the electrons in the beam are mostly due to the charges within a distance along the tube roughly given by the distance of the electron from the center of the tube (20 $\mu m$), which in turn is roughly $1/\gamma$ of the total. Furthermore, the motion due to these forces in the lab frame takes $\gamma$ times longer to happen in the lab frame due to time dilation. The two effects combined imply that the Coulomb interactions are suppressed by $1/\gamma^2$, making them basically negligible.

Strong transverse bunch shape anisotropy is also seen in electron beams in synchrotrons. At the Cornell Electron Storage Ring (CESR), $\varepsilon_x \approx 500 eV m/c = 2.7 \times 10^{-25} kg m^2/s$ along the horizontal ($x$) direction, and $\varepsilon_y \approx 0.1 eV m/c = 5.3 \times 10^{-29} kg m^2/s$ along

---

33 The beam can lower its emittance by emitting X-rays, which is important in synchrotrons and undulators.
the vertical direction: a factor of 1000 anisotropy! Indeed, one of the ambitious goals in the accelerator community is to get the transverse emittance down to the quantum limit.

(c) What would the vertical emittance \( \Delta q_y \Delta p_y \) be at the quantum limit, set by the uncertainty principle? By what factor must CESR shrink their emittance to approach this goal?

Liouville’s theorem implies that reducing the entropy or the emittance cannot be done simply with the standard tools of accelerators (magnetic lenses, focusing solenoids, bunchers, etc.). Instead, the beam must be coupled to another system into which the entropy can be dumped. There are different methods for doing this, such as synchrotron cooling, electron cooling, and stochastic cooling (for which Simon van der Meer received the Nobel Prize in 1984). Here we will consider a simplified model of electron cooling.

Electron cooling is a mechanism by which an electron beam is used to reduce the momentum spread, and therefore the entropy, of a beam of heavier ions. It is being used in the Relativistic Heavy Ion Collider (RHIC), colliding gold ions to create a quark-gluon plasma. The electron and gold ion beams are overlapped with nearly the same velocity. Here again we assume the momenta of the two beams have a Gaussian distribution, characterized by a temperature.

(d) Suppose the center-of-mass momenta in a gold ion bunch has initial temperature \( T_{Au} \), and is put into contact with a co-moving electron bunch of the same radius \( \sigma_q \), the same number of particles \( N \) and a temperature \( T_e \ll T_{Au} \). What will the final temperature \( T_f \) of the two ion beams be, if they have enough time to exchange kinetic energies and thermalize? (You may ignore the interactions between the various ions.) Calculate the ratio of the one-dimensional emittances \( \varepsilon_{Au}/\varepsilon_e \) and the beam angular dispersion \( \Delta \theta_{Au}/\Delta \theta_e \) of the resulting gold ion and electron bunches.

As the gold ion beam travels through the electron gas, it will experience two effects. One is a drag force proportional to the velocity with respect to the center of mass. The other is fluctuations due to individual scattering events with the electrons. In Exercise 6.18, you found a fluctuation-response relationship between the noise and the drag coefficient in simulations of Langevin dynamics for one-dimensional motion. The relationship you found in that exercise was

\[
k_B T = \langle \Delta p^2 \rangle / 2 \eta \Delta t. \tag{N1.22}
\]

We express this relation in terms of \( \eta \), the inverse of the mobility, so the force on a moving gold ion is \( F = -\eta \dot{v} \). (The mobility is called \( \gamma \) in Exercise 6.18.) For us, \( \Delta p \) is a momentum change due to a collision, \( T \) is the temperature of the electron beam, and \( \Delta t \) is the time between collisions.

So, to find the drag force for a moving gold ion, we shall calculate the noise on a stationary ion and use this relationship.

---

\footnote{The horizontal width is large because of the randomness introduced by the emission of X-ray photons as the beam circles around.}
The geometry of collisions between electrons and ions via the Coulomb interaction is complicated. One simplifying concept is that of a scattering cross section—the effective circular disk that would suffer the same forces. For the densities of electrons and gold ions in our two bunches, you may assume that a gold ion near the center of the electron bunch has a cross section $\Sigma = 0.1 \text{nm}^2 = 10^{-19} \text{m}^2$. The momentum exchange in a collision with that disk will solely depend on the momentum $p_x$ of the electron, where $x$ is the axis perpendicular to that disk.

How big a momentum change will a gold ion have during a collision with an electron? (How much momentum is exchanged when a ping-pong ball hits a car?)

(e) Taking the limit $m_{\text{Au}}/m_e \to \infty$, what is the net change $\Delta p$ for a stationary gold disk, given a head-on elastic collision with an electron with $x$ momentum $p_x$? Assume that the particles are non-relativistic in the center-of-mass frame.

Let us now calculate the other terms in eqn N1.22.

(f) Give a formula for the average time $\Delta t$ between collisions from the cross section $\Sigma$ of the stationary ion disk, the electron number density $n$, and the momentum distribution of the electrons. Give a formula for the average squared momentum transfer $\langle \Delta p^2 \rangle$ for these collisions. Give the formula for the temperature $k_B T$ of the electrons. (Rough estimates are fine. You may assume the electrons have a Gaussian momentum distribution with width $\sigma_p$.)

At the Relativistic Heavy Ion Collider (RHIC) work is underway to cool gold ions with an electron beam. In one scheme, a gold ion beam circles a ring with circumference $C = 3.9 \text{km}$, and each turn passes through a segment of length $0.0078 C$ with a co-moving electron gas of density $10^8 \text{cm}^{-3} = 10^{14} \text{m}^{-3}$ and momentum spread $1.2 \times 10^{-23} \text{kg m/s}$. The electrons are refreshed continuously using a Cornell-invented energy recovery Linac (ERL). The bunches are moving fast: their value of $\gamma = 1/\sqrt{1 - v^2/c^2} = 105$. Time dilation makes the interaction time needed larger by a factor of $\gamma$.

(g) Using your answers from part (f) and eqn N1.22 from Exercise 6.18, find a formula for the inverse mobility $\eta$. Using $F = -\eta v = \dot{p}$, find a formula for the exponential decay time $\tau$ for the gold ion velocity in the center of mass frame. Evaluate it for RHIC’s electron cooling scenario.

(h) What is the exponential decay time for the gold ion beam in the lab frame of RHIC?

N1.17 Nonabelian defects. (Mathematics) 35

In this problem, we will try to understand the defects that occur in a liquid crystal composed of a planar tri-headed molecule (Figure N1.21). The ordered states of this system have all molecules in the same orientation, but with liquid translational order (randomly placed centers). This exercise will explore subtle questions regarding classification of nonabelian defects. It will explore braiding with mathematics related to the problems of anyon statistics and topological quantum computing. And it will provide

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35 This exercise was developed in collaboration with Stephen Thornton.
a complete analysis of the transformation of one defect as it is pulled around another, explored in the case of fingerprints in Exercise 9.17.

![Fig. N1.21 Dihedral molecules.](image)

(a) The tri-headed molecule studied in this problem. (b) The molecular structure of 1, 3, 5-Trichlorobenzene, with a representation of a tri-headed molecule superimposed [4]. If these molecules were to form a liquid crystal, the liquid crystal would have the types of defects described in this exercise.

If the molecule 1, 3, 5-Trichlorobenzene (Fig. N1.21b), which has the same symmetry as our cartoon molecule, had a liquid crystalline phase\textsuperscript{36} where they oriented parallel to one another, one would find the defects we study here [4]. These systems are generally called \textit{dihedral} liquid crystals for reasons that we will explore shortly.\textsuperscript{37}

![Fig. N1.22 Defects with molecules confined to the plane.](image)

Fig. N1.22 Defects with molecules confined to the plane. Note that the molecules are at random positions; the orientations are shown on a square lattice for convenience.

Figure N1.22 shows two defects (red points) around which the molecules swirl. Recall that the winding number is defined as the fraction of a full rotation $\Delta \phi/2\pi$ an object does as we travel around a defect counterclockwise in real space. $\Delta \phi$ has a sign associated with it. Is the defect labeled by the winding number? In part (a) we find that the answer is yes if the molecular “legs” are confined to the plane, but otherwise no:

(a) \textit{What are the winding numbers of the defects shown in Figure N1.22? If we allow the molecules to rotate in three dimensions, are these two defects topologically equivalent? Why or why not?} (Hint: Try applying a rotation about various axes to all the molecules.)

We saw in part (a) that the molecular orientation, after following the path around a defect...\textsuperscript{36}There is currently no experimental evidence for this specific type of liquid crystal.\textsuperscript{37}Note: Most papers on dihedral liquid crystals confine the molecules to the plane. We are allowing them to rotate in 3 dimensions.
defect, must return to an orientation related by the symmetry group of the molecule.
Our molecule has a dihedral symmetry group $D_3$ (hence the name dihedral liquid crystal). We imagine that the homotopy group (the nonequivalent possible circular paths in the order parameter space) will be related somehow to $D_3$.

$D_3$ is a six element group, generated\(^{38}\) by two elements: a $2\pi/3$ counter-clockwise rotation $a$ about an axis normal to the plane of the molecule (taken to face out of the page for the planar configurations), and a flip $x$, rotating by $\pi$ about a particular leg of the molecule (the left-hand portion of the group presentation N1.23 below). The multiplication table (Table N1.1) can be derived from the three equality relations on the right side of presentation N1.23:

$$D_3 = \left\langle x, a \mid x^2 = 1, a^3 = 1, xax^{-1} = a^{-1}\right\rangle.$$ (N1.23)

The group element $gh$ is found in the row labeled by $g$ and the column labeled by $h$. This is important because $D_3$ is nonabelian.

![Multiplication table for the dihedral group $D_3$.](image)

Table N1.1: Multiplication table for the dihedral group $D_3$. Note that the group is non-abelian: $gh$ is not equal to $hg$. The element $gh$ would be found in the row labeled by $g$ and the column labeled by $h$.

(b) What are the symmetry group elements given by the paths in Fig. N1.22 (a) and (b)? Explain in words why $xax^{-1} = a^{-1}$. Use the multiplication rules to calculate $xa \otimes xa$ and $xa \otimes xa^2$. Compare your answers to those in the table.

We know from part (a) that two different dihedral symmetry group elements can be topologically equivalent—so the defect strength is not just given by the element of $D_3$. We shall explore this further below. But first, let us see if the same dihedral symmetry group element can correspond to two non-equivalent topological defects.

---

\(^{38}\)A group is *generated* by a subset of elements (called *generators*) if every member of the group can be represented by repeatedly multiplying generators and their inverses.
Fig. N1.23 Two equivalent defects, one with the molecules confined to the plane, and the other with one leg rotated to point ‘upward’.

(c) What is the winding number of the defect shown in Fig. N1.23(a)? If $D_3$ were the homotopy group, and we were to assign this defect an element of $D_3$, which element would it be? What does this suggest about the defect? Can we heal this defect by rotating into the third dimension? Does the rotation into the third dimension shown in Fig. N1.23(b) heal the defect in the order parameter field?

We need a homotopy group that somehow has a category for a defect that returns the molecule to its original orientation trivially (the identity element in $D_3$), and somehow also allows a defect to return the molecule differently (nontrivially) to the same labelling of the legs (the element $a^3$). There is a wonderful treatment of the topological theory of defects by Mermin [21], which we draw upon here.

How can a $2\pi$ rotation in three dimensions be different from the identity? For those experienced in quantum physics, rotating a spin $\frac{1}{2}$ electron by $2\pi$ changes the sign of the wavefunction. This is because the spin wavefunction is represented not by a vector, but by a $2 \times 2$ unitary matrix in $SU(2)$, and there are two $SU(2)$ matrices for every rotation in $SO(3)$. Similarly, in our problem for every rotation $g$ in $D_3$ there are two rotations $\pm g$ in $Dic_3$: the 12-element dicyclic group, which is the first homotopy group for our dihedral liquid crystal. The multiplication rules for $Dic_3$ are

$$Dic_3 = \left\{ x, a \mid x^2 = -1, a^3 = -1, xax^{-1} = a^{-1} \right\} \quad (N1.24)$$

Here we presume that $a$ corresponds to a $1/3$ counter-clockwise turn. (We will not write out the whole multiplication table.)

(d) View the defect in Fig. N1.23(a) as three $1/3$ counterclockwise turns. What is the homotopy group element in $Dic_3$ for the defect? How did lifting to $Dic_3$ fix the problem we had in part (c)?

The rotation of the molecular orientation by $2\pi$ gives a net minus sign, just as for an electron spin.

---

[21] Mermin tells us that the order parameter space of our dihedral liquid crystal is $SO(3)/D_3$ (the continuous broken symmetry modulo the residual symmetry group). The first homotopy group of a simply-connected group modulo a discrete subgroup is the discrete subgroup—but $SO(3)$ is not simply connected. He prescribes using the simply-connected universal cover $SU(2)$, and the corresponding discrete group $Dic_3$. Thus $\Pi_1(SO(3)/D_3) \cong \Pi_1(SU(2)/Dic_3) \cong Dic_3$. 

---
Now we still need to understand how the two defects in Fig. N1.22, with different group elements (in both $D_3$ and $Dic_3$), can be the same defect. Mermin tells us that defects with non-abelian homotopy groups are not classified by their homotopy group elements, but by the *conjugacy classes* of the group. The conjugacy classes are subsets of the group whose elements are related by conjugation. Two group elements $g$ and $h$ are said to be conjugate if there exists a group element $\gamma$ such that $h = \gamma g \gamma^{-1}$.

(e) Show in complete generality that the identity element $g = 1$ of a homotopy group is the only element in its conjugacy class. Interpret this physically: can there be two different ways of having no defect? One can check that $-1$ commutes with all other group elements, implying that it too is the only element in its class.

Do the two defects in Fig. N1.22 indeed lie in the same conjugacy class?

(f) What is the homotopy group element $\beta \in Dic_3$ for the defect in Fig. N1.22(a)? The two defects in that figure combine to form no defect, so the defect in Fig. N1.22(b) must have homotopy group element $\hat{\beta} = \beta^{-1}$. Find a group element $\gamma$ that shows $\hat{\beta} = \gamma \beta \gamma^{-1}$.

Is there a physical reason for this peculiar conjugacy class criterion? Indeed: it is precisely the transformation of the defect $\beta$ when it *braids* around another defect $\gamma$.

Braiding in three dimensions is associated with hair-styling: you take two or more strings of hair or string and weave them over and under one another to form a kind of rope. If we move a point defect with strength $\beta$ in two dimensions around another defect $\alpha$ (with the necessary continuous readjustments of the order parameter fields), we can view the process as braiding in space-time.

40The conjugacy classes of $Dic_3$ are as follows:

\[
C_0 = \{1\}, \quad C_0^* = \{-1\}, \quad C_1 = \{a, -a^2\}, \quad C_2 = \{a^2, -a\},
\]

\[
C_{\text{even}} = \{x, xa^2, -xa\}, \quad C_{\text{odd}} = \{xa, -x, -xa^2\}.
\]

41Braiding is important for the study of anyons. Nonabelian braiding is important for topological quantum computing.
Fig. N1.24 Braiding two defects. (a-d) can represent the motion of a red two-dimensional point defect around a brown defect in time, or a red line defect curling around a brown line defect in three dimensions. (a) The red defect has strength $\beta$, measured by the path beginning and ending at $X$. (The connection to $X$ makes this a based homotopy class, which is important for the argument but not crucial here.) (b-c) The defect is continuously dragged around the brown defect of strength $\alpha$. The measured strength cannot change, either as the red defect continuously is dragged, or as the path anchored at $X$ is continuously modified to surround $\beta$ without touching $\alpha$. (d) When the red defect returns to its original position, the original path 2 may measure a different strength $\tilde{\beta}$ than $\beta$ measured by the deformed path $1 \to 2 \to 3$.

Figure N1.24 visually argues that a defect $\beta$ encircling a defect $\alpha$ indeed changes its homotopy group element.\(^{42}\) It argues that the original path $\beta$ deforms into another path (in the same homotopy class) as the defect circumnavigates $\alpha$. This path becomes the product of three simpler loops.

(g) In Fig. N1.24(d), what homotopy group elements $g_1$, $g_2$, and $g_3$ correspond to the three segments of the path? (The labels $\tilde{\beta}$ and $\alpha$ suggest the answers for two of the three.) What is $\beta$, in terms of $g_1$, $g_2$, and $g_3$?\(^{43}\) What is $\tilde{\beta}$ in terms of $\beta$ and $\alpha$?

The schematic of braiding in Figure N1.24 is translated into a real configuration of our order parameter with two defects in Figure N1.25.

---

\(^{42}\)We found in Fingerprints (Exercise 9.17) that moving a dislocation around a disclination can change the sign of its Burger’s vector. This too is due to a non-abelian homotopy group, discussed by Poenaru et al.\(^{26}\) in one of the founding papers in this field. They find that the combined order parameter space of rotations and translations relevant for fingerprints is the Klein bottle. Its first homotopy group is nonabelian, which provides an explanation for the dislocation annihilation in Fig. 9.36.

\(^{43}\) It turns out that there is ambiguity in whether a concatenation of two loops (traveling along 1 and then 2) with homotopy elements $g_1$ and $g_2$ corresponds to a single combined loop with homotopy element $g_1g_2$ or $g_2g_1$. However, once a convention is chosen, the element assigned to the concatenation of several loops $g_1, \ldots, g_n$ is fixed. You are free to choose this convention.
Fig. N1.25 Braiding in practice. (a) The order parameter field in the vicinity of two defects. (b) The order parameter field after one defect has been pulled a quarter-way around the other. (c) The resulting order parameter field after the braiding process. Also, view the evolution of the order parameter in time at the course Web site [31].

(h) What is the homotopy group element $\beta \in \text{Dic}_3$ of the left defect in Fig. N1.25(a)? What is the element $\alpha$ for the defect in the center of the figure?44 The central brown defect does not change during the braiding process, just as in Figure N1.25. What is the homotopy group element $\hat{\beta}$ for the left defect in frame (c)? Use the group multiplication rules (eqn N1.24) to verify that your prediction in part (g) for $\hat{\beta}$ is correct.

N1.18 Light proton tunneling. (Dimensional analysis) 3

Fig. N1.26 Atom tunneling. A hydrogen atom tunnels a distance $a_0$, breaking a bond of strength $E_{\text{bind}}$ equal to its ionization energy.

In this exercise, we continue to examine a parallel world where the proton and neutron masses are equal to the electron mass, instead of $\sim 2,000$ times larger.

With everything two thousand times lighter, will atomic tunneling become important? Let’s make a rough estimate of the tunneling suppression (given by the approximate WKB formula $\exp(-\sqrt{2MVQ}/\hbar)$).

Imagine an atom hopping between two positions, breaking and reforming a chemical bond in the process—an electronic energy barrier, and an electronic-scale distance. The distance will be some fraction of a Bohr radius $a_0$ and the barrier energy will be some fraction of a Rydberg, but the the atomic mass would be some multiple of the proton mass.

44 The element assigned to this defect also depends on multiple conventions, including whether the molecules in Fig. N1.25 rotate up or down in the third dimension. Your choice of convention will affect the element $\alpha$, but not the result of the braiding $\hat{\beta}$. 
In our world, what would the suppression factor be for an hydrogen atom of mass \( \sim M_p \) tunneling through a barrier of height \( V \) of one Rydberg = \( \hbar^2/(2m_e a_0^2) \), and width \( Q \) equal to the Bohr radius \( a_0 \)? How would this change in the parallel world where \( M_p \to m_e \)? (Simplify your answer as much as possible.) (Use the real-world\(^{45} \) \( a_0 \) and Rydberg for the parallel world, not your answers from a previous exercise. Also please use the simple formula above: don’t do the integral. Your answer should involve only two of the fundamental constants.)

\[ \text{N1.19 Beer.} \]

We shall experimentally explore the statistical mechanics of beer. To explore all aspects, you will need

- Warm and cold non-alcoholic beer, in bottles or cans. It should not be too filtered, so that the head of beer is stable for some time. (Sodas will not work well.)
- Disposable transparent cups, spoons, bottle opener if necessary,
- Salt, sugar, and sand,
- A tub or pail to dispose of liquids.

Avoid shaking the beer before opening. (We will shake it later.) Carefully open a beer, catching any overflow in the cup. Notice there is a pressure release upon opening.

There are a variety of ideas one might explore.

**Rigidity transition.** Is beer foam a solid, liquid, or gas? Are beer bubbles in the beer a solid, liquid, or gas? What is the nature of the transition as the bubbles approach the surface, losing the fluid between them?

**Coarsening.** Are the bubbles in the beer changing in size or arrangement before they pop at the top? Can you observe what the mechanisms are for bubble evolution?

**Nucleation.** Are new bubbles formed deep inside the liquid? Or do they form along the edges of the container? Does this change if you shake the container? Do you remember where and how bubbles of boiling water form?

Try adding salt. Are the effects chemical, or physical? Speculate. Is there a relation to cloud seeding?

If you are feeling quantitative, what should the nucleation rate be, according to theory? (The surface tension of water is \( \sigma \sim 7.56 \text{ N/m at } 0^\circ \text{C} \). The prefactor for the nucleation rate is about \( 10^{13}/\text{s per water molecule} \).)

**Thermodynamics.** Which is more foamy, a cold bottle of beer or a warm one? What does that mean about the temperature dependence of the chemical potential of \( \text{CO}_2 \) in water? Do you remember seeing small bubbles form as you heat water, long before it boils?

\(^{45}\)The reduced mass effects you found in the earlier exercise will be much less important for larger atoms and molecules, so we shall not include them here.
Examine an unopened, unshaken bottle of beer. Are there any bubbles?

Take a mostly empty container of beer. Holding it shut, shake it to get as much foam as possible. Can you get bubbles with a larger volume than that of the original beer? How much larger? If you can, find some non-alcoholic beer (like Guinness) that uses nitrogen rather than CO₂. Is the behavior different?

At what partial pressure of CO₂ in the beer will bubbles stop forming? At what partial pressure will it reach equilibrium with the air in the room? How does this relate to the boiling and evaporation of water?

Consider a bubble of gaseous CO₂ in equilibrium with the beer. What determines whether molecules of CO₂ will move into or out of the bubble? If you are feeling quantitative, the beer acts as an ideal gas of CO₂ (or N₂) except for a free energy per molecule due to interactions of a dissolved molecule with the surrounding water. The beer foam forms at nearly constant temperature when the pressure changes as the bottle is opened.

N1.20 **Distinguished and undistinguished particles.**

In quantum physics, all electrons are fundamentally indistinguishable. In classical physics, we often have systems where the particles are in principle distinguishable, but nothing in our experiment depends on which particle is which. (Examples might include colloidal particles, grains of sand, atoms differing only in their isotopes or nuclear spins, etc.) In Section 3.5, we introduced the idea of “undistinguished” particles, and in Section 7.5 we argued that non-interacting undistinguished particles obey Maxwell–Boltzmann statistics. How can we derive the properties for these undistinguished particles, and why is it useful?

In statistical mechanics, when our experiments do not depend on some microscopic details about a system, we create a free energy by “integrating out” the microscopic details. If we promise to ignore the differences between particles that are in principle distinguishable, we must sum over all the possible ways of labeling the particles.

(a) Suppose we have a system composed of \( M \) different types of particles, with \( N_1 \) indistinguishable particles of type 1, \ldots, \( N_M \) indistinguishable particles of type \( M \). The system has a total of \( N \) particles, in a microcanonical ensemble with total energy \( E \) in a volume \( V \), which has an energy shell volume of

\[
\Omega_{\text{dist}}(E, V, N_1, N_2, \ldots, N_M) .
\]  

(N1.26)

Assume the energy of the particles is independent of their identities – that all Hamiltonians describing our experiment is the same under a permutation \( P_n \) of the positions and their momenta,

\[
\mathcal{H}(x_1, \ldots, x_N, p_1, \ldots, p_N) = \\
\mathcal{H}(x_{P_1}, \ldots, x_{P_N}, p_{P_1}, \ldots, p_{P_N}) .
\]  

(N1.27)

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46 This exercise was developed in collaboration with Stephen Thornton.

47 See also Cates and Manoharan [7], who discuss several colloidal paradoxes connected to this exercise.
Calculate $\Omega^{\text{summed}}(E,V,N)$ by summing $\Omega^{\text{dist}}$ over all permutations $P$, and write it in terms of factorials and $\Omega^{\text{dist}}(E,V,N,0,\ldots,0)$, as if all particles were of the first type. Show that, if all particles are distinct, so $M = N$ and $N_m \equiv 1$, that $\Omega^{\text{summed}}(E,V,N) = N! \Omega^{\text{dist}}(E,V,N,0,\ldots,0)$.

Given our experiments do not care about the identities, $\Omega^{\text{summed}}$ will give exactly the same predictions as did $\Omega^{\text{dist}}$ (as is always true of free energies formed in this way). But is that what we want?

Remember Nernst’s law (the “third law of thermodynamics”), that the quantum entropy per particle vanishes at zero temperature? This is true if all of the particles are identical, or if all of the particles are different in ways important to the Hamiltonian (so the ground state is not invariant under permutations). This is because there are usually only a small number of states with the minimum possible energy $E_{\text{min}}$ (see footnote 20 on page 151).

(b) Consider a quantum system with $N$ distinguishable particles, with a Hamiltonian invariant under permutations of the particles, and with a crystalline ground state with energy $E_{\text{min}}$. What is the entropy per particle $S^{\text{summed}}(E_{\text{min}},V,N)/N$ at zero temperature? Use Stirling’s formula to simplify it. Does it go to zero for large $N$? Is it at least extensive? (Hint: Assume $\Omega^{\text{dist}}(E_{\text{min}},V,N,0,\ldots,0)$ has only one lowest energy configuration.)

This is technically correct. If the particles are distinct, even at zero temperature they can be swapped around in lots of ways. But if our experiments do not care about this swapping, perhaps we can drop the $N!$ and forget about this annoying permutation entropy?

This drastic step is precisely how we defined Maxwell–Boltzmann statistics in eqn 7.48, $\Omega(N)^{\text{MB}} = \Omega(N)^{\text{dist}}/N!$.

The fact that entropy should be extensive was a consequence of the third key property of Shannon entropy (Section 5.3.2). The entropy $S^{\text{summed}}$ is not extensive: the sum $S_1 + S_2$ for two weakly interacting subsystems (and hence the product $\Omega_1 \Omega_2$ of their energy shell volumes) is not anywhere near that of the combined system. In particular, if you double the total number of particles, energy, and volume, $S^{\text{summed}}$ does not double: the permutation entropy grows much faster than linearly in $N$. Maxwell–Boltzmann statistics cures this second annoyance.

In section 3.5, we divided the distinguishable energy shell volume $\Omega^{\text{crude}}$ for the ideal gas (eqn 3.48) by $N!$ to get our final formula for the entropy (eqn 3.57).

(c) Show that the three terms in the last line of eqn 3.49 for the ‘crude’ entropy $S^{\text{crude}}$ in the limit of large systems scale in the same way with system size, and are not extensive. (Hint: Use Stirling’s formula.) Show that our final formula for the ideal gas entropy (eqn 3.57) is extensive. In detail, fill in the steps between eqn 3.55 and 3.57 (ignoring Planck’s constant) to see how dividing $\Omega^{\text{crude}}$ in eqn 3.48 by $N!$ fixes this.

Maxwell–Boltzmann statistics also is needed to talk about separating weakly-interacting subsystems – separating our experiment from the rest of the world. We use this exten-
sively in our model of temperature and chemical potential as connecting our subsystem to an external bath. Here we illustrate this problem, not with a classical phase space, but with a system with discrete energy levels. We shall see that Maxwell–Boltzmann statistics in this case does not have a simple interpretation in terms of counting how to place undistinguished particles into energy levels, but does serve to fix another serious problem with applying $\Omega^{\text{diff}}$ to systems with varying number of particles.

Figure N1.27 shows a system at fixed energy $E$ and particle number $N$, with a subsystem with energy $E_s$ and number $N_s$, and a bath with the remaining energy $E_b = E - E_s$ and $N_b = N - N_s$. All particles in our bath are in a state with energy zero. Particles in the subsystem occupy one of two states, with energies $+\Delta$ or $-\Delta$.

![Diagram of a system with a subsystem and a bath](image)

Fig. N1.27 Microcanonical, fixed $N$ system composed of a subsystem potentially exchanging energy and particles with a bath. The current state has system and subsystem energy $E = E_s = -\Delta$, $N_s = 5$ and $N = 10$.

Let us begin with $N = 3$ and $E = 0$. We shall be interested in the volume of the energy shells $\Omega(E, N)$, which here is given by the number of different configurations that the particles can take. Usually we would use something like eqn 3.23, giving the probability $\rho_s(E_s, N_s)$ of the subsystem to have energy $E_s$, should generalize to

$$\rho_s(E_s, N_s) \doteq \frac{\Omega_s(E_s, N_s)\Omega_b(E - E_s, N - N_s)}{\Omega(E, N)};$$

(N1.28)

the probability should be the fraction of the energy surface that has the desired energy and number in the subsystem. Here $\Omega_s$ and $\Omega_b$ are calculated by assuming that they are isolated from one another.

First consider distinguishable particles. Here, to calculate $\Omega_s^{\text{dist}}(E_s, N_0, N_1, N_2)$ for the isolated subsystem we fill it with particles 0 and 1, and the bath with particle 2.

(d) What is the number of states $\Omega_s^{\text{dist}}(0, 1, 1, 0)$ for when the subsystem has zero energy and has particles 0 and 1 inside? What is $\Omega_b^{\text{dist}}(0, 0, 0, 1)$ when particle 2 is in the bath? What is the total number of states $\Omega^{\text{dist}}$ with energy zero and three distinguishable particles, assuming the two sides are allowed to exchange energy and particles? Compute
the probability \( \rho^\text{frac}_s(0,2) \) from the fraction of configurations of the whole system that have zero energy and two particles in the subsystem. Compute the probability \( \rho^\text{pred}_s(0,2) \) predicted by eqn N1.28, assuming \( \Omega_s(E_s, 2) \) and \( \Omega_b(E - E_s, 1) \) are given by the uncoupled calculations \( \Omega^\text{dist}(E, 1, 1, 0) \) and \( \Omega^\text{dist}_b(E, 0, 0, 1) \). Do they agree? Which is correct? Why are they different?

This too is fixed by using Maxwell–Boltzmann statistics.

(e) What are \( \Omega^\text{MB}_s(0,2) \) and \( \Omega^\text{MB}_b(0,1) \)? What is \( \Omega^\text{MB}(0,3) \), including contributions from other particle arrangements of zero energy? (Hint: Use eqn 7.48 and your answers for \( \Omega^\text{dist} \).) Which of the three has no natural interpretation as a number of ways of filling the states with undistinguished particles? Compute the probability given by eqn N1.28. Does it agree with the correct answer from part (d)?

These \( N! \) factors might remind you of the entropy of mixing in Section 5.2.1. There we started with a symmetric split, with the same number of atoms in the subsystem and the bath, and with all the interaction energies between the atoms the same (the Hamiltonian is independent of permutations). If the atoms on the left and right are different (orange and blue), when the partition is lifted the entropy jumped. But if they are the same, the entropy hardly shifted.

Of course, the entropy of mixing is real! If we had a membrane that would let only orange particles through, half of them would diffuse into the blue half leading to a big pressure difference. We could then extract work by moving the wall between the two subsystems. We can still use Maxwell–Boltzmann statistics for the orange atoms and the blue atoms separately. But we should not use Maxwell–Boltzmann statistics ignoring the orange/blue difference, if we plan later to use a membrane whose Hamiltonian distinguishes them.

N1.21 Localization.\(^{48}\) (Quantum, Condensed Matter) 3

In Section 7.4, we discussed how non-interacting electrons provide a useful model for metals, even though the electron-electron interactions are strong. The Fermi liquid of quasiparticles is a kind of adiabatic continuation of the noninteracting electron system, connected by perturbation theory. Here we shall study how a one-dimensional non-interacting metal responds to disorder. We shall discuss how metals with weak disorder are understood by perturbing around the clean state. We shall discover that strong disorder leads to an insulating system whose eigenstates are not extended, but localized. We will describe these localized states explicitly by perturbing about a state of isolated atomic states.

Consider a one–dimensional chain of atoms \( n \), each with one noninteracting electron state \(|n\rangle\) of energy \( U_n \), that can be occupied by either zero or one spinless electrons.

\(^{48}\)Hints for the computations can be found at the book web site [31].
Electrons can hop between atoms with matrix element $t$, leading to the Hamiltonian

$$\mathcal{H} = \sum_{n=1}^{N} U_n |n\rangle \langle n|$$

(N1.29)

$$-t \sum_{n=1}^{N-1} (|n\rangle \langle n+1| + |n+1\rangle \langle n|)$$

$$= \begin{pmatrix}
U_1 & -t & 0 & \ldots & 0 & 0 \\
-t & U_2 & -t & \ldots & 0 & 0 \\
0 & -t & U_3 & \ldots & 0 & 0 \\
0 & 0 & -t & \ldots & 0 & \vdots \\
\vdots & \vdots & \vdots & \ddots & \vdots & -t \\
0 & 0 & 0 & \ldots & -t & U_N
\end{pmatrix}$$

We shall take the random energies $U_n$ as uniformly distributed between $-W$ and $W$. Without disorder ($W = 0$), this is a textbook model used to describe energy bands in crystals. Three dimensional analogs of this ‘tight-binding’ model are quite realistic models of Fermi surfaces and energy bands in real materials.\(^{49}\)

(a) Write a function that builds the Hamiltonian matrix of eqn N4.80 with size $N$, bandwidth $2W$, and hopping matrix element $t$. Studying zero disorder $W = 0$, find the eigenvectors for $t = 1$, and $N = 100$, sorted by their eigenvalues. Plot the eigenvectors for the four lowest energies. Check numerically that these four are sinusoidal with wavevectors $k_\alpha = \pi\alpha/(N+1)$ appropriate for a box of size $N$ with hard-wall boundary conditions half a grid spacing to either side. Check that their four eigenvalues are the corresponding $E_{k_\alpha} = -2t \cos(k_\alpha)$

Imagine a 1D metal at zero temperatures with electrons filling the states up to a Fermi surface, here just two points at some $\pm k_{\text{Fermi}}$. Consider a packet of electrons made up of eigenstates near $k_{\text{Fermi}}$ traveling to the right. The wavepacket\(^{50}\) will travel, as usual, at the group velocity $dE_k/dk|_{k_{\text{Fermi}}}$, without dissipation.

Now let us explore what happens when we add a weak disorder.

(b) Build a Hamiltonian with weak disorder $W = W_{\text{weak}} = 0.04$, $t = 1$, and $N = 100$. Plot the lowest four eigenvectors. Are the eigenstates still extensive (reaching from one side of the box to the other)?

---

\(^{49}\)We shall see that even a small disorder changes the metallic behavior of one-dimensional electrons in a qualitative and interesting way. Indeed, one dimensional electrons are unstable in many interesting ways. Adding interactions between electrons, they become Luttinger liquids, with emergent scale invariance. Adding interactions with lattice vibrations, they can become topological insulators, with solitons and fractional charges.

\(^{50}\)Wavepackets are used to connect waves to particle-like motion. In a non-disordered system, one superimposes states with similar momenta to make a spatially localized wavefunction, which then moves with the group velocity of the wave. We discuss wavepackets to motivate the effects of disorder, but no knowledge about them is required to do this exercise.
At this point, we could use perturbation theory to calculate the disordered eigenstates and energy levels. We could then create wavepackets and see how they evolve. In three dimensions, the scattering off of the disorder changes the electron transport qualitatively. Instead of wavepackets moving forever in one direction (ballistic transport, infinite conductivity), one gets diffusive motion of the electron probability through space (disorder providing an elastic scattering length, and a finite conductivity). In three dimensions, this is a good model for metals with impurities or dopants, illustrating how one can understand complex behavior by perturbing around solvable special cases.

Instead, let us examine what happens at large disorder $W$, or equivalently, small hopping $t$. (All of our eigenvectors depend only on $W/t$, and we will perturb in $t$ to study the localized states.)

(c) Set $t = t_{\text{Weak}} = 0.1$, $W = 1$, and $N = 100$, plotting the ten eigenvectors with lowest energies. Also do a log-linear plot of the probability density (absolute square of the wavefunctions) for these eigenvectors. Do the eigenstates still look as if the will be extensive (stretching from one end of a macroscopic wire to the other)? What solvable special limit of $H$ should we use to capture this new behavior?

Here we find the eigenstates appear localized – fixed in space near individual ‘atoms’. The probabilities in these states fall exponentially with distance from their centers. A wavepacket formed from localized states like these cannot transport current: for large disorder, our model describes an insulator.

Just as one can use perturbation theory to describe dirty metals in three dimensions from models like ours, we can use perturbation theory to calculate and understand these localized states. You should remember the use of second-order perturbation theory to describe the energies of a Hamiltonian $H = H_0 + V$ for small $V$. You may not remember that the first step was to use first-order perturbation theory to determine the eigenvectors. If $|\Psi_i^{(0)}\rangle$ has unperturbed eigenvalue $E_i^{(0)}$, then to first order

$$|\Psi_i^{(1)}\rangle = |\Psi_i^{(0)}\rangle + \sum_{i\neq j} \frac{\langle \Psi_j^{(0)} | V | \Psi_i^{(0)} \rangle}{E_i^{(0)} - E_j^{(0)}} |\Psi_j^{(0)}\rangle.$$  \hspace{1cm} (N1.30)

If the hopping is small compared to the disorder, let us perturb in $t$.

(d) What are the eigenenergies for $H$ in eqn N4.80 with $t = 0$? Argue that, to first order in $t$, the new eigenstates will be confined to three adjacent sites.

(e) Write a function, given $H$, $i$, $t$, and $N$, that gives the perturbed eigenstate to lowest order in $t$ that is centered at site $i$. Find the site of the ground state with largest probability. Plot the true ground state and your first-order approximation to it. (Hint: You may be unlucky, and happen to have a neighbor site with a near degenerate energy. Just create a new Hamiltonian and try again.)

What controls whether our model is a metal or an insulator? For a given $W/t$, are all the states either extended or all localized? Or could there be some mixture?
We can examine this by defining a rough measure of how spread out the wavefunction is, called the participation ratio:

$$P(\psi) = \frac{(\sum_n |\psi(n)|^2)^2}{\sum_n |\psi(n)|^4} = \frac{1}{\sum_n |\psi(n)|^4}. \quad (N1.31)$$

(f) Show that a state whose probability is spread uniformly among $M$ sites has $P = M$. At zero disorder, what is the participation ratio for the lowest energy state? For the long-wavelength next few states? What is the ratio for a localized state that decays exponentially, $\psi(j) \propto \exp(-|i - j|/\lambda)$, in an infinite chain, with $\lambda$ much larger than one?

If the participation ratio $P \ll N$ we can reasonably expect that the eigenstate is localized.

(g) Calculate the participation ratio for all the eigenstates for intermediate disorder $W_{\text{inter}} = 0.5$, $t = 1$, $N = 100$, and plot them against the energy. Is there a systematic variation? Plot the wavefunction for an energy in the middle of the energy band (eigenvalue $E$ near zero), and one at the top and bottom of the band. Which are less localized – the states near the edges of the band, or the states in the center?

In experiments, one finds a region of localized states at the edges of a band, and extended states in the middle of the band. Between these is a mobility edge, where a metal-insulator transition occurs as more electrons are added.

Finally, can we find a mobility edge for our model? One thing to check is if the wavefunctions might have decay lengths larger than our system (so they just look extended at $N = 100$).

(h) Find the eigenvectors and eigenvalues for the same parameters as in part (g), $W = W_{\text{inter}} = 0.5$ and $t = 1$, except for a much larger system ($N = 2000$ if it is feasible on your system). Plot the participation ratio verses energy, and plot eigenstate in the center of the band and at the two edges. Do the states in the middle of the band now appear localized? Are their participation ratios larger than 100 – the system size in part (g)? Does it make sense that they looked extended in the smaller system, but clearly in an infinite system are localized?

As it happens, disordered electrons in one dimension are always localized, even for tiny disorder. The spinless, noninteracting electrons we study here are also always localized in two dimensions. In two dimensions, they can become extended when interactions, spin orbit scattering, or strong magnetic fields are added. In particular, 2D electrons in a strong magnetic field exhibit the quantum Hall effect (with extended states around the

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51 Each time we add one order to perturbation theory, we get a wavefunction extending outward by one atom. It appears from the first two terms that each order multiplies the terminal amplitude by a factor $|\psi(i+n)/\psi(i+(n-1))| = |t/(U_i - U_{i+n})|$. Roughly speaking, if $U_i$ is at the edge of the band, this factor is twice as small as if $U_i$ is in the center, so there is less localization at the center. Notice, though, that this is useful only when $t > |U_i - U_{i+n}|$. Rare, nearly degenerate states can mix strongly, even at long distances, making the arguments subtle.
edges). Even more interesting, interacting electrons in a strong magnetic field exhibit the fractional quantum Hall effect – our first example of an experimental system with fractional charges and fractional statistics.

N1.22 Correlation matching.\textsuperscript{52}  

Figure N1.28 shows the 2D correlation function $C(r)$ and its 2D Fourier transform $\tilde{C}(k)$ for a periodic stripe pattern (for which an 8x8 block is shown).

![Figure N1.28: Stripe pattern correlations](image)

\textbf{Fig. N1.28 Stripe pattern correlations.} Tile: An 8x8 Ising pattern $s(x, y)$ that is repeated to fill space. Black is $-1$, white is $+1$. $C(r)$: The 2D correlation function for the tiled space. $\tilde{C}(k)$: The Fourier transform of the correlation function $\tilde{C}(k)$. Note that the zero of both the correlation function and its Fourier transform are not centered; you may imagine invisible rows $-4 \equiv 4$ and $-\pi \equiv \pi$.

Figure N1.29 shows three other 8x8 blocks of spins.

![Figure N1.29: Tiles](image)

\textbf{Fig. N1.29 Tiles.} Snapshot tiles of an Ising model with period 8 in the x and y directions.

\begin{itemize}
  \item For each of the tiles in Fig. N1.29, identify which of the choices in Fig. N1.30 corresponds to $C(r)$, and which corresponds to $\tilde{C}(k)$. The color scale and axes for each are given by Fig. N1.28.
\end{itemize}

\textsuperscript{52}This exercise was developed in collaboration with Stephen Thornton.
Consider an Ising model on a square lattice, starting in an infinite-temperature random configuration, and shifted to a low temperature. It will coarsen with time, as shown in the two panels in Fig. N1.31.

These two pictures were created with different dynamical rules. In one of them, we use a traditional Metropolis algorithm, flipping one spin at a time. This does not properly describe, say, oil and water separating from one another, because one cannot "flip" an oil molecule into a water molecule – the molecules are locally conserved. The other uses an algorithm invented by Kawasaki, which swaps the spins of neighboring

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This exercise was developed in collaboration with Stephen Thornton.
sites (with a probability determined by their six neighbors). This allows up-spin “oil”
molecules to diffuse through down-spin “water” regions.
It probably is not obvious from the two snapshots which evolved without conservation
and which was conserved. We shall use correlation functions to determine which is
which.
What correlation function should we use? Our systems are translation invariant, so
the correlation function is expected to depend only on the separation between
spins, \( r = x - x' \). The system is qualitatively changing in time (as it coarsens),
so \( C(r, t) = \langle s(x, t)s(x + r, t) \rangle \) is a natural choice. The insets of the two panels
in Fig. N1.32 show a 2D plot of \( C(r, t) \) for the evolution conditions of the corre-
sponding snapshots in Fig. N1.31, and the main panels plot the angular average\(^{55}\)\( C(r, t) = (1/2\pi) \int_0^{2\pi} d\theta C(r \cos(\theta), r \sin(\theta), t) \).

**Fig. N1.32 Coarsening correlation functions.** The two angular-averaged spin-spin cor-
relation functions, each for the corresponding snapshots in Fig. N1.31(a) & (b). Distances
are measured in lattice spacings. (a) The noise in the 2D inset would disappear with an en-
semble average over initial conditions, but the oscillations in the \( C(r) \) plot are real. (b) The
correlation functions are an average of many ensemble snapshots prepared like Fig. N1.31(b).
Thinking about how the two systems grow should allow you to deduce which is which
without mathematics.

(a) *Note the oscillations in \( C(r, t) \) on the left, that are absent on the right. Decide
whether oscillations are more clearly necessary for conserved or non-conserved growth.
Explain your reasoning physically, without calculations.* (In particular, your reasoning
should be different from the argument below!)
Let us now produce a definitive argument using \( C(r, t) \) to settle which figure has non-
conserved dynamics. Consider \( C^s(\delta) = \langle s(x) s(x + \delta) \rangle_x \), the correlation function for
a snapshot of a particular coarsening system, and \( C^s(r, t) \), its angular average.
(b) *Show, for an \( N \)-spin Ising model, that*
\[
\int d\delta_x d\delta_y C^s(\delta_x, \delta_y, t) = M(t)^2 / N, \tag{N1.32}
\]
*where \( M \) is the total magnetization of the snapshot at time \( t \). Go to polar coordinates,
and relate this formula to \( \int_{r>0} r C^s(r, t) dr \).*

\(^{54}\)They have periodic boundary conditions.

\(^{55}\)Here and elsewhere we approximate the discrete lattice of spins as a continuous integral over space, for
simplicity. Also, the apparent rotation invariance in these snapshots is only approximate. It is true that the
Ising model at \( T_c \) has an emergent rotational invariance, but this is not true for low temperatures.
CHAPTER 1. EOPC: NEW EXERCISES

Fig. N1.33 Using \( rC(r) \) to study coarsening. The correlation functions shown in Fig. N1.32, multiplied by \( r \). We shall use them to determine which is conserved and which non-conserved, and use one of them to measure the coarsening length scale.

(c) Presuming both conserved and non-conserved simulations start with zero magnetization at infinite temperature,\(^{56}\) which of the two snapshots in Fig. N1.31 evolved with conserved dynamics? Use Fig. N1.33 and eqn N1.32 to explain why.

Now let us explore the coarsening dynamics using our understanding of emergent scaling behavior. When the features in the system are large compared to the lattice and small compared to the size of the system, the equal-time correlation function is expected to take a scaling form

\[
C(r, t) = C(r/L(t)).
\]  

(N1.33)

The system looks statistically similar to itself at previous times, but rescaled by a growing lengthscale \( L(t) \). Our scaling assumptions suggest that \( L(t) \propto t^\rho \), where the critical exponent \( \rho \) depends on the type of dynamics (conserved, non-conserved, surface diffusion, . . . ).

In testing whether universal scaling functions hold, we often use scaling collapses. Usually we would plot \( C(r, t) \) versus \( r/t^\rho \), especially if we know what power to expect. But in many cases we would like to use our simulations or experiments to measure \( L(t) \), see if they obey our scaling assumption (eqn N1.33), and measure \( \rho \). So, we would like a reliable way of measuring \( L(t) \).

(d) Show that the scaling assumption of eqn N1.33 implies that

\[
\int d^2 r C(r, t) = \alpha L(t)^2,
\]  

(N1.34)

where \( \alpha \) is a time-independent constant. What is \( \alpha \) in terms of the scaling function \( C(x, y) \)?

This suggests that, so long as \( \alpha \) is nonzero,\(^ {57}\) a good measure of the length scale is

\[
L(t) \propto \sqrt{2\pi \int_0^\infty d r r C(r, t)},
\]  

(N1.35)

\(^{56}\) We could be more fussy. At \( T = \infty \) in the finite Ising model, \( M \) is the sum of \( N \) numbers \( \pm 1 \), so random walks tell us that \( \langle M^2 \rangle = N \), which is precisely the contribution to \( C(r, 0) \) at \( r = 0 \). So the negative regions for must cancel the positive regions for \( r > 0 \).

\(^{57}\) Note that we learned from parts (b) and (c) that \( \alpha \) is zero for coarsening with a conserved order parameter. All is not lost. We could use a different moment of \( C(r, t) \), another feature (like the first zero-crossing), or just rescale the curves until they overlap.
**Fig. N1.34 Coarsening correlation collapse.** (a) When plotted versus \( r/L(t) \), the correlation functions \( C(r, t) \) at late times collapse onto a single curve, validating our scaling ansatz in eqn N1.33. (b) The measured \( L(t) \) curves are roughly straight on a log-log plot, implying a power law \( L(t) \sim t^\rho \), as predicted by emergent scale self-similarity. The fit is to the late-time behavior.

To test that our scaling ansatz works, we can measure \( L(t) \) for various times \( t \), and plot all the curves \( C(r, t) \) versus \( r/L(t) \). Shown in Fig. N1.34(b) is a plot of \( L(t) \) versus \( t \) for the evolution used in panel (b) of the previous figures in this exercise. Fig. N1.34(a) shows \( C(r, t) \) plotted against \( r/L(t) \) for various times \( t \).

At late times, we expect the 2D scaling form in eqn N1.33 to hold, and we observe in Fig. N1.34(a) that the 1D angular averages \( C(r, t) \) curves collapse well when plotted versus \( r/L(t) \).

(e) Argue that the common curve \( C(r, t) \) vs. \( r/L(t) \) at late times is the angular average of \( C \). Scaling collapses allow one to measure universal scaling functions with experiments or simulations. *Is the power law found in Fig. N1.34(b) closer to that predicted for conserved, or non-conserved coarsening?*

**N1.24 Ising critical correlations.**

Here we examine numerical simulations of the equilibrium, equal-time, spin-spin correlation function

\[
C(r) = \langle s(x)s(x+r) \rangle = C(r)
\]  

(N1.36)

near the Ising critical point (where the 2D correlation function becomes isotropic, so it only depends on \( r = |r| \)).

Figure N1.35 shows snapshots of the Ising model at \( T < T_c \), at \( T = T_c \), and at \( T > T_c \), together with their equilibrium, equal-time, radially averaged correlation functions \( C(r) \).

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\(^{58}\) This exercise was developed in collaboration with Stephen Thornton.
It is challenging to estimate the correlation length from snapshots of systems that are approaching critical points.

(a) Estimate the correlation length using the snapshots for $T < T_c$ and $T > T_c$, and mark the corresponding point on the corresponding curve in the correlation function plots. Discuss how you determined the correlation length in each case. What feature of $C(r)$ in the curves roughly corresponds to your estimates of the correlation length from the snapshots?

(b) Use the plots of the correlation function at $T_c$ to estimate the critical exponent $\eta$ for the 2D Ising model, where $C(r) \sim r^{-\eta}$ at $T_c$ in $d = 2$. Compare with the known exact value.

(c) Sketch the connected correlation function $C_{\text{conn}}(r) = \langle s(\mathbf{r} + \mathbf{x})s(\mathbf{x}) \rangle - \langle s(\mathbf{x}) \rangle^2$ for the $T < T_c$ curve in the linear-linear plot.
Chapter 2

Advanced Statistical Mechanics

These exercises cover topics in more advanced statistical mechanics and the renormalization group, algorithms, and computer science.

N2.1 Singular corrections to scaling and the RG. ③

In this exercise, we derive the form of the scaling function (eqn 12.102) for the effects of irrelevant operators on the properties of systems near critical points (see Exercise 12.31). Remember that irrelevant directions shrink under coarse-graining. Let \( \chi \) be the susceptibility of the Ising model, as a function of the reduced temperature \( t = T_c - T \) and some irrelevant operator \( u \):

\[
\begin{align*}
\frac{d\chi}{d\ell} &= -(\gamma/\nu)\chi, \\
\frac{dt}{d\ell} &= t/\nu, \\
\frac{du}{d\ell} &= -yu.
\end{align*}
\] (N2.1)

How do we derive the universal scaling function \( X(z) \) from these renormalization-group flows? Consider the flows illustrated in Fig. 12.8, except now with a third dimension involving the prediction \( \chi \). Consider a point \((t_0, u_0, \chi_0)\) in the system space, and the invariant curve defined by \( z = u_0 t_0^\omega \) (dashed lines). Our renormalization group allows us to calculate \( \chi(\ell, t, u) \) along these curves—relating the behavior everywhere near the critical manifold (vertical swath flowing toward \( S^* \)) to the properties along the outgoing trajectories, which approach closer and closer to the unstable manifold (the horizontal swath flowing away from \( S^* \)).

For example, we can define the universal scaling function \( X(z) \) (for positive time \( t \)) to be the \( \chi_{t^*} \) where the flow crosses \( t_{t^*} = 1 \).

(a) Solve eqns N2.1 for \( u_\ell \) and \( t_\ell \). Setting \( t_{t^*} = 1 \), what is \( u_{t^*} \) in terms of your invariant combination \( z \)?

So we label each invariant scaling curve by the value of the vertical position \( u_{t^*} \) where it crosses \( t_{t^*} = 1 \).
N2.2 Nonlinear RG flows and analytic corrections.\footnote{This exercise was developed in collaboration with Colin Clement}

We consider the effects of nonlinear terms in renormalization-group flows. The Ising model in zero field has one relevant variable (the deviation $t$ of the temperature from $T_c$). To calculate the specific heat, we shall also consider the flow of the free energy per spin $f$ under the renormalization group. Instead of a discrete coarse-graining by a factor $b$, here we use a continuous coarse-graining measured by $\ell$. One can think of one coarse-graining step by $b = (1 + \epsilon)$ as incrementing $\ell \to \ell + \epsilon$; equivalently, coarse-graining to $\ell$ changes length scales by $\exp(\ell)$.

Consider the particular flow equations\footnote{Note that these are total derivatives. So the first equation tells us the total change in $f$ after coarsening by a factor $1 + d\ell$. $f(t)$ then will coarse-grain to $f_\ell(t_\ell)$ without needing to worry about the chain rule $df(t)/d\ell = \partial f/\partial t + \partial f/\partial t dt/d\ell$.}

\begin{align}
\frac{df_\ell}{d\ell} &= D f_\ell - a t_\ell^2 \\
\frac{dt_\ell}{d\ell} &= t_\ell/v,
\end{align}

(N2.2)

where $D$ is the dimension of space and $a t_\ell^2$ is a nonlinear term that will be important in two dimensions.

We shall call the free energy per spin of the actual system $f_0$, and the temperature of the actual system $t_0$. The coarse-grained free energy and temperature are $f_\ell$ and $t_\ell$, after being coarse-grained by a factor $\exp(\ell)$. (Hence at $\ell = 0$ we have not yet coarse-grained, so $f_0 = f$ and $t_0 = t$.) Notice here that the free energy of our system is the initial condition $f_0(t_0)$ of this differential equation, and $f_\ell = f(\ell)$ and $t_\ell = t(\ell)$ are the renormalization-group flows of the two variables. To derive the scaling behavior, we shall coarse-grain to $\ell^*$ where $t_\ell^* = 1$, at which point the coarse-grained free energy is $f_\ell^*$.

Let us start with the linear case $a = 0$.

(a) Solve for $f_\ell$ and $t_\ell$ for $a = 0$. Setting $t_\ell^* = 1$, solve for $f_0$ in terms of $f_\ell^*$, $t_0$, $D$, and $v$. Solve for the specific heat per spin $c = T \partial^2 f_0/\partial T^2$, where $t_0 = (T - T_c)/T_c$. Show that the specific heat near $T_c$ has a power-law singularity $c \propto t^{-\alpha}$, with $\alpha = 2 - D v$. (For example, in $D = 3$, $v \approx 0.63$, so $\alpha = 2 - D v \approx 0.11$; the specific heat diverges at $T_c$.) Writing

\begin{equation}
\alpha = t^{-\alpha} (c_0 + c_1 t + c_2 t^2 + \ldots),
\end{equation}

(N2.3)

what is the first correction $c_1$ to the specific heat near $t = 0$ in the absence of the nonlinear term?

Why is the linear term in the free-energy flow equal to the dimension, $df/d\ell = D f + \ldots$, where all other terms are hard-to-compute critical exponents? There is no completely
general answer to this question (although there are arguments for specific models). Indeed, in other models of disordered systems and glasses, and models above the upper critical dimension, the linear term is not given by the dimension. The relation \(2 - \alpha = D\nu\) is called a hyperscaling relation (to emphasize they involve the dimension \(D\)), and these other models are said to violate hyperscaling. One physical description of hyperscaling involves the singular part of the free energy \(f\) inside a correlated region.

(b) In the case \(a = 0\), show that the singular free energy contained in a correlated volume \(\xi^D\) near the critical temperature becomes independent of the distance to the critical point.

Glassy and disordered systems become extremely sluggish as they are cooled. In at least some cases, this is precisely because the energy barriers needed to continue equilibration diverge as their correlation lengths grow—they are glassy because their RG flows violate the hyperscaling relation.

So much for the power law singularity—what about the correction term \(c_1\) in part (a)? It is an analytic correction to scaling.\(^3\) Here it is subdominant—near the critical temperature where \(t \to 0\), it is less singular than the leading term. One expects in a real physical system that the microscopic bond free energy \(J\) between spins will be some analytic function of temperature, and the physical free energy and specific heat will be multiplied by \(J\). Expanding \(J\) in a Taylor series about \(t = 0\) would also give us terms like those in eqn N2.3.

Does the introduction of the higher-order nonlinear term \(at_\ell^2\) in eqn N2.2 change the behavior in an important way? Rather than exercising your expertise in analytic solutions of nonlinear differential equations, eqn N2.4 provides not \(f_\ell\) and \(t_\ell\) as functions of \(\ell\), but the relation between the two:

\[
f_\ell(t_\ell) = f_0 \left( \frac{t_\ell}{t_0} \right)^{D\nu} - \frac{avn_\ell^2}{(2 - D\nu)} (1 - (t_\ell/t_0)^{(2-D\nu)}).
\]

(c) Show that \(f_\ell(t_\ell)\) in eqn N2.4 satisfies the differential equation given by eqn N2.2, using

\[
\frac{df_\ell}{dt_\ell} = \frac{df_\ell}{d\ell} / \frac{dt_\ell}{d\ell}.
\]

Show that it has the correct initial conditions at \(\ell = 0\). What is \(f_\ell^*\) at \(\ell^*\), where \(t_\ell = 1\)? Show your method.

Again, it is important to remember that \(f_\ell(t_\ell)\) is not the free energy as a function of temperature—it is the coarse-grained free energy as a function of the coarse-grained temperature of a system starting at a free energy \(f_0\) at a temperature \(t_0\). It is \(f_0(t_0)\)

---

\(^3\)These are distinct from singular corrections to scaling that arise from irrelevant terms under the renormalization group (Exercise 12.31).
that we want to know. Since here we have only one relevant variable (in zero field), all the flows lead to the same \(^4\) final point \(f_\ell (t_\ell^* = 1)\)

(d) Solve for \(f_0\) in terms of \(f_\ell\) and \(t_0\). Solve for the specific heat \(c = T \partial^2 f / \partial T^2\), where \(t = (T - T_c) / T_c\). Show that it can be written in the form

\[
c = c_{\text{+ analytic}}(t) + t^{-\alpha} c_{\text{* analytic}}(t)
\]

where the additive correction \(c_{\text{+ analytic}}(t)\) and the multiplicative correction \(c_{\text{* analytic}}(t)\) have a simple Taylor series about \(t = 0\). Write these two corrections, in terms of \(f_\ell^*, T_c, a, \nu,\) and \(D\).

Here the nonlinear term \(a\) gives us both a smooth multiplicative term and a smooth additive background to the specific heat. This makes sense physically. An additive term would come from other degrees of freedom like atomic vibrations, or a box holding the magnet. A multiplicative term would arise, for example, from an effective temperature dependence of the coupling between spins (say, due to thermal expansion separating the atoms).

N2.3 **Beyond power laws: 2D Ising logs.**

The two-dimensional Ising model has a logarithmic singularity in the specific heat. The exact result shows that the specific heat per spin is [20, eqn 3.119]

\[
c(T) = k_B \frac{2}{\pi} \left( \frac{2J}{k_B T_c} \right)^2 \left[ - \log(1 - T / T_c) \\
+ \log(k_B T_c / (2J) - (1 + \pi / 4)) \right] \\
= - \frac{8}{\pi k_B T_c^2} \log \left( \frac{t}{\sqrt{2 k_B T_c \exp(-(1 + \pi / 4))}} \right)
\]

where \(t = (T - T_c) / T_c\) and we set \(J = 1\). (Remember \(\log(t)\) is negative for small \(t\).) Linearizing the flows around the renormalization-group fixed point predicts a power law \(c \sim t^{-\alpha}\). When \(\alpha \to 0\) one often observes logarithmic corrections, as we see in eqn N2.7. But such corrections are not predicted by the RG flows after linearizing! The key nonlinear term is the term \(at_\ell^2\) of eqn N2.2 we studied for 3D in Exercise N2.2.

(a) Is the solution for \(f_\ell (t_\ell)\) in eqn N2.4 useful in \(D = 2\)? Why or why not? (Hint: The exponent \(\nu = 1\) for the two-dimensional Ising model.)

Again, we provide the solution of the nonlinear RG eqns N2.2, now for \(D = 2\)

\[
f_\ell (t_\ell) = f_0 (t_\ell / t_0)^2 - at_\ell^2 \log(t_\ell / t_0).
\)

\(^4\)Remember for systems with more than one variable (say \(t\) and \(h\)), the free energy depends on the invariant curve departing from the fixed point, labeled, say, by \(h / t^\beta = h_\ell^* (t_\ell^* = 1)\). We solve for \(f_0 (t_0, h_0)\) in terms of \(f_\ell^*(1, h_\ell^*)\) just as we do in part (b).

\(^5\)This exercise was developed in collaboration with Colin Clement.
(b) Show that $f_\ell(t_\ell)$ in eqn N2.8 satisfies the differential equation given by eqn N2.2, with the correct initial conditions. Solve for $f_\ell$ in terms of $f_\ell^*$ and $t_0$, where $t_\ell^* = 1$. Solve for the specific heat $c = T\partial^2 f / \partial T^2$, where $t = t_0 = (T - T_c)/T_c$ and $f = f_0$. (Remember the chain rule: $\partial f / \partial T = (\partial f / \partial t)(\partial t / \partial T)$.) Does it agree asymptotically with the exact result in eqn N2.7? What are $c_0$ and $\tau$, in terms of $a$ and $f_\ell^*$?

Thus for the 3D Ising model (Exercise N2.2), nonlinear terms in the renormalization-group flow equations give only analytic corrections to scaling, where in the 2D Ising model they introduce logarithms in the specific heat. Normal form theory (see Exercise 12.4) can be used to determine when one may safely linearize these equations, and to organize the other critical points into universality families [28].

N2.4 Eigenvectors near the renormalization-group fixed point. ③

The critical exponents in the renormalization group are given by the eigenvalues of the RG transformation linearized near the fixed point. What do the eigenvectors mean?

Consider a two-dimensional Ising model with two parameters, a nearest-neighbor bond $K = J/T$ and a next-neighbor interaction $K_2 = J_2/T$ lying along the diagonal bonds.

$$H = -K \sum_{i,j} S_{i,j} S_{i+1,j} + S_{i,j} S_{i,j+1}$$

$$-K_2 \sum_{i,j} S_{i,j} S_{i+1,j+1} + S_{i,j} S_{i+1,j-1} \quad \text{(N2.9)}$$

If we decimate to the black squares of a checkerboard (say, $i + j$ even), we get a new square-lattice Hamiltonian rotated by 45° coarse-grained by a factor $b = \sqrt{2}$. The next-neighbor bond basically becomes a nearest-neighbor bond—it mostly renormalizes to zero in one step, and contributes its value to the new nearest-neighbor coupling. The deviation of the nearest-neighbor bond from the critical point $K^*$, we may crudely assume, rescales by a factor $b^{1/\nu}$ under coarse-graining (remember $K \sim J/T$) and then is increased by $K_2$. So under one coarse-graining step

$$K' - K^* = b^{1/\nu}(K - K^*) + K_2,$$

$$K_2' = 0. \quad \text{(N2.10)}$$

(a) Our crude renormalization-group flow is already linear. What is the fixed point? What is the Jacobian $J$ about the fixed point? What are the eigenvalues $\lambda_0$ and $\lambda_1$? (Let $\lambda_1$ be the relevant eigenvalue, greater than one.)

Our Jacobian matrix is not symmetric (or Hermitian), so it has two sets of eigenvectors—left eigenvectors $\ell_\alpha J = \lambda_\alpha \ell_\alpha$, and right eigenvectors $J r_\alpha = \lambda_\alpha r_\alpha$.

(b) What are the left and right eigenvectors? Are the left eigenvectors orthonormal? Are they normal to the right eigenvector that has a different eigenvalue?

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6Instead of thinking of a Hamiltonian space with temperature as an extra parameter, it is convenient to work at fixed temperature, and mimic raising temperature by lowering the overall scale of the energy.
(c) Draw the flow in the \((K, K_2)\) plane near the fixed point. Indicate the directions of the left eigenvectors and right eigenvectors in different colors. Also draw the boundary between the ferromagnetic and paramagnetic phase. How is this boundary related to the stable manifold of the fixed point? Is it related to any of the eigenvectors?

Consider a new set of scaling variables \(u_\alpha\), given by the dot products of the displacement from the fixed point with the left eigenvectors:

\[
u_\alpha = \ell_\alpha \cdot (K - K^*, K_2) \tag{N2.11}\]

(d) Show that the phase boundary has \(u_1 = 0\) (using the convention that \(\lambda_1\) is the relevant direction). How do the coordinates \(u_\alpha\) flow under the renormalization group?

In general, there is a nonlinear transformation between the parameters \(T, H, J_2, \ldots\) in a Hamiltonian and the natural coordinates \(t(T, H, J_2), h(T, H, J_2), u(T, H, J_2)\) which flow simply under the renormalization group. This coordinate change is one of the contributors to analytic corrections to scaling.

(e) Are \(u_0\) and \(u_1\) relevant, irrelevant, or marginal? Which coordinate, \(u_0\) or \(u_1\), is the scaling variable corresponding to the reduced temperature \(t(K, K_2)\)? If we write a property of our system \(X(K, K_2) = X(K(u_0, u_1), K_2(u_0, u_1)) = u_1^* \mathcal{X}(u_0 / u_1^*)\), can there be any dependence on \(u_0\), within our crude model? How does \(X\) vary near the phase boundary?

N2.5 Is the fixed point unique? Period doubling.\(^7\) (Dynamical systems) \(^3\)

Is the fixed point of the renormalization group unique? (It seems unlikely that coarse-graining the Ising model in momentum space gives the same fixed point Hamiltonian as real-space decimation on a square lattice. One would be spherically symmetric, the other has a square symmetry. Naturally, both look the same on long length and time scales, but their short-distance behavior is different.) If not, and there are many alternative fixed points in system space describing a phase transition, can any system at the critical point be a fixed point, for a suitable renormalization group?

We shall answer this question for the particular case of the period-doubling onset of chaos. In particular, we shall investigate what happens to the renormalization-group fixed point as we change coordinates. There is no reason to expect that Nature measures distances \(x\) in the same way, though, as we do. We could equally well decimate and rescale in a different coordinate \(y = \phi(x)\), where we assume \(\phi\) is smooth, monotone increasing, has a smooth inverse \(\phi^{-1}\). We will be considering one-humped maps with a maximum at \(x = 0\), so we shall assume \(\phi(0) = 0\) to keep the maximum at zero.

(a) Give the formula for the function \(\tilde{g}(y)\) corresponding to the map \(g(x)\), in terms of \(g\) and \(\phi\). Hint: You need to find the \(x\)-value from \(y\), then apply \(g\) to get the new \(x\), and then find the new \(y\). Show that the inverse formula is for \(g\) in terms of \(\phi\) and \(\tilde{g}\) is

\[
g(x) = \phi^{-1}\tilde{g}(\phi(x)) \tag{N2.12}\]

\(^7\)There are many exercises exploring this chaotic logistic map (see Index).
If \( g(x) \) is at the onset of chaos (showing universal scaling behavior), will \( \tilde{g}(y) \) also be at the onset?

Recall from Exercises 12.9 and 12.29 that the transition to chaotic motion (Fig. 12.17) of one-humped maps \( g(x) \) is understood using a renormalization group that decimates in time by a factor of two using \( g(g(x)) \), and rescaling the coordinate \( x \) by a factor of \( \alpha \approx -2.5 \):

\[
T[g](x) = \alpha g(\alpha^{-1}x)).
\]

Let \( g^* \) be the fixed point \( T[g^*] = g^* \) of our renormalization group in the space of one-humped maps.

(b) Write a renormalization-group transformation \( \tilde{T} \) for which \( \tilde{g}^* \) is a fixed point. (Hint: Use the formula \( T[g^*] = g^* \), and the formula for \( g^* \) in terms of \( \tilde{g}^* \) and \( \phi \) from part (a). Change variables to \( y \) and solve for \( \tilde{g}(y) \) on the right-hand side.) Show that \( \tilde{T} \) can be viewed as a decimation plus a nonlinear "stretching" function \( \tilde{\alpha}(y) \), \( \tilde{T} = \tilde{\alpha} \circ \tilde{g} \circ \tilde{g} \circ \tilde{\alpha}^{-1}(y) \). What is \( \tilde{\alpha}(y) \)?

So there is an infinite family of plausible, nonlinear renormalization-group transformations, with an infinite family of fixed points given taking our original fixed point \( g^*(x) \) and changing variables \( x \to \phi(x) \).

Can we change variables so that our fixed point is a parabola (as in the logistic map)? Can we make any one-humped map at the onset of chaos a fixed point, by choosing an appropriate change of coordinates?

There is an elegant proof that this is not possible. Our one-humped map has a fixed point \( g^*(x^*) = x^* \), and also many periodic orbits\(^8\) \( g^{[2^n]}(x_n) = g^*(\ldots(x_n)\ldots) = x_n \) with period 2, 4, 8, \ldots (Beware! \( g^* \) is a fixed point in function space under the transformation \( T \). \( x^* \) is a fixed point on the real line under the function \( g \).

(c) Show that \( dx/dy = \phi^{-1}(\phi(x)) = 1/(dy/dx) = 1/\phi'(x) \). If \( x^* \) is a fixed point of \( g^* \), what is the fixed point \( y^* \) of \( \tilde{g}^* \)? Find the slope \( d\tilde{g}^*/dy \) at the new fixed point, and show that it equals the slope \( g^*(x^*) \) at the old fixed point. Similarly, show that \( dg^{[2^n]}(x_n)/dx \) is \( g^{[2^n]}(x_n)/dy \). (Note that \( dg(\ldots(x)\ldots)/dx = g'(g(\ldots))g'(\ldots) \ldots \), so the product of the derivatives along any period orbit is invariant under coordinate changes.)

So, unless the logistic map happens to have the same derivative as \( g^* \) at their respective fixed points, there can be no coordinate transform taking one to the other, and there is no reason to think there is some nonlinear renormalization-group transformation that has the logistic map as a fixed point. (Later we shall find a more direct way to see that the logistic map has different behavior than any fixed point.) Indeed, all the fixed points of all the iterates of a map would have to agree with the iterates of \( g^* \) to allow for such a transformation. So our critical surface in function space, where one-humped maps

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8Remember the period doubling bifurcations, which start with a stable \( 2^n \) cycle and end with a stable \( 2^{n+1} \) cycle surrounding an unstable \( 2^n \) cycle, forming \( 2^n \) little pitchforks. All of these unstable \( 2^n \) cycles survive to the onset of chaos.
are poised at the onset of chaos, has two subsurfaces—the maps that can be formed under coordinate changes (and could be fixed points), and the maps that cannot be fixed points.

We can gain more insight into these two types of critical points by considering the renormalization-group flow of functions $\tilde{g}^*$. (It is a fixed point of $\tilde{T}$ in part (b), but it must flow to $g^*$ under our original renormalization-group transformation $T$.) Consider maps near to $g^*$, formed by picking coordinate transformations $\phi$ that are near to the identity:

$$\phi(x) = x + \epsilon \sum_{p=1}^{\infty} \phi_p x^p.$$  \hfill (N2.14)

(d) Write $\phi^{-1}(y)$ to linear order in $\epsilon$, as a similar sum. Calculate the change $\tilde{g}^*(x) - g^*(x)$ to linear order in $\epsilon$. (Hint: Start with the equation $\phi(g^*(x)) = \tilde{g}^*(\phi(x))$.) Express it as a sum $\epsilon \sum_{p=1}^{\infty} \phi_p \Psi_p(x)$.

(e) Show that $g^*(g^*(x/\alpha)) = g^*(x)/\alpha$. By differentiating $T[g^*][x]$, show that $g''^*(x) = g''^*(g(x/\alpha))g''^*(x/\alpha)$. Calculate the change $T[\tilde{g}^*](x) - g^*$ to linear order in $\epsilon$. Use your first two formulas to show that the change can be written $\epsilon \sum_{p=1}^{\infty} \phi_p \alpha^{1-p} \Psi_p(x)$.

You have shown that $\Psi_p(x)$ is an eigenfunction of $T$, with eigenvalue $\alpha^{1-p}$. You have also shown that only these eigenvalues and eigenfunctions can be generated by infinitesimal changes of coordinates. Feigenbaum conjectured in his early paper [12, p.687] that these eigenfunctions were the only ones. Our argument in part (c) suggests that there must be functions near to $g^*$ that cannot be reached by a change of coordinates.

The lowest few eigenvalues $\lambda_n$ of $T$, calculated for perturbations in the even subspace, are approximately

$$\{\lambda_1, \lambda_2, \ldots\} = \{4.67, 1.00, 0.160, -0.124, -0.0573, 0.0255, -0.0101, \ldots\}$$ \hfill (N2.15)

For your convenience, we provide the powers of $\alpha \approx -2.503$, so

$$\alpha^{1-p} = \{1.00, -0.400, 0.160, -0.0638, 0.0255, -0.0102, 0.00407, \ldots\}$$ \hfill (N2.16)

(f) Is $\lambda_4$ a power of $\alpha$? Was Feigenbaum correct? Give the next few eigenvalues $\lambda_n$ that are not given by coordinate changes. Which powers $p$ are missing from the list? Do those powers correspond to even, or odd perturbations to $g^*$?

So, at least at the period-doubling onset of chaos, there are two kinds of critical points, those that can be reached by a smooth change of coordinates (and could be fixed points of a different renormalization group), and those that cannot. We propose a new nomenclature, based on the analogy with gauge transformations in electromagnetism and field theory. In electromagnetism, a gauge transformation can be viewed as a spatially varying change in the convention of how we measure (gauge) the phase of the wavefunction for charged particles: $\Psi(x_1, x_2, \ldots) \rightarrow \prod_j e^{i\chi(x_j)}\Psi$. Here smooth changes of coordinates acts to change the way we measure position $x$ in our chaotic system; more
generally we might consider also changing the way we measure the control parameter \( \mu \), or (for an Ising model) changing the definitions of magnetization \( M \), temperature \( T \), external field \( H \), etc. We shall call the first \textit{gauge} critical points and the second (more common) category \textit{singular} critical points. More generally, there are three sources of corrections to scaling. There are \textit{singular corrections} to scaling, which are generated by irrelevant eigenvectors of the renormalization group that are new, irrational numbers not related to the other exponents (perhaps like \( \lambda_4 \) in part (f)). Smooth coordinate transformations in the control variables (here the parameter \( \mu \) in the logistic map) generate what are traditionally termed analytic corrections to scaling. Finally, there are what are traditionally termed redundant corrections to scaling, which correspond to systems that could have been fixed points of a different renormalization group. We shall call both of these last two \textit{gauge} corrections to scaling: they can give corrections with integer power laws (like the analytic background in the specific heat), or corrections that are combinations of integers and the relevant and gauge-irrelevant critical exponents.

How in practice can we tell the special “gauge” critical points from the “singular” ones?

\[ \text{Fig. N2.1 Expanded bifurcation diagram.} \] The attractor at the fixed point \( g^* \) has a perfect self-similarity upon flipping and rescaling about the function maximum \( x = 0.5 \) (green-dashed line). The lower first-tier branch of the bifurcation diagram maps onto the entire diagram when rescaled vertically by \( \alpha \) and horizontally by \( \delta \); the same happens to the lower branch (whose upper second-tier branch maps onto the entire lower first-tier branch), and so on. Any other map will have this property only asymptotically, near the maximum (green dashed line) and near the critical point.

Suppose we study a map at its critical point, which deviates from \( g^* \) primarily along the eigendirection corresponding to \( \lambda_4 = -0.124 \). This deviation will lead to changes in the spatial patterns seen in the attractor (Fig. N2.1); for example, the ratio \( \alpha_n \) of the widths of the \( n^{th} \) and \( (n+1)^{st} \) tier branch will not be precisely \( \alpha \).

Let the \( \psi_n(x) \) be the \( n^{th} \) eigenfunction of \( T \).\(^9\)

\[ \text{(g) Consider an infinitesimal perturbation of the fixed point } g^* + \epsilon \psi_4(x) \text{ along this direction. How does the leading correction to } \alpha_n - \alpha \text{ scale with } \lambda_n? \] (Hint: How does

\[ \text{We already have a name for the special eigenfunctions, so for example the eigenfunction associated with } \lambda_2 = 1 = \alpha^{1-p} \text{ with } p = 1 \text{ is } \psi_2(x) = \Psi_1(x) = \psi_2(x)). \]
\( \alpha_n \) change under one application of the renormalization group operator \( T \)?

Thus the special, gauge critical points have corrections to scaling that are expressable solely in terms of the relevant critical exponents.

**Conjectures:**

N2.1 The allowed renormalization-group fixed point functions are precisely those accessible by analytic coordinate transformations \( y = \phi(x) \), which are precisely those generated by moving along the scaling variables \( u^n \) associated with the eigenvalues \( \lambda_n \) not given by powers of \( \alpha \) (in a way similar to how infinitesimal Lie algebra symmetries are related to Lie group operations). In particular, a large change of variables will have corrections to scaling at the critical point only involving powers of \( \alpha \).

N2.2 We conjecture that all corrections to scaling for one-humped maps [19] will be given by changing to *normal form* coordinates. So, for the logistic map \( f(x) = \mu x(1-x) \), the traditional analysis suggests there are nonlinear scaling coordinates that satisfy \( T(u_n) = \lambda u_n \); the curve in function space traced by the logistic map then has coordinates \( u_1(\mu) = \mu - \mu_\infty + b_1(\mu - \mu_\infty)^2 + \ldots, \ u_2(\mu) = a_2 + b_2(\mu - \mu_\infty) + c_2 \ldots, \ldots \). We are led to conjecture that a further change of coordinates \( x \to \phi(x) \) allows us to remove all corrections involving the special eigendirections with \( \lambda = \alpha^{1-p} \) for integer \( p > 1 \). That is, we conjecture that not only is there a gauge submanifold in the surface of critical points, but that the critical points can be foliated into surfaces where the singular corrections are the same.

N2.3 We conjecture that the separation between gauge and singular fixed points we define here have analogies in the Ising model and other critical points. (Indeed, we took the name from that literature, where it involves changes in the coordinates describing the fields.) See in particular reference [13], which answers the analogous question as to whether there is a renormalization group that can make any Ising critical point into its fixed point.

N2.4 We note that, in the case of thermodynamic critical points, the distinction between control variables (like \( t = T - T_c \) and \( h = H/T \)) and “results” (like magnetization \( M(t, h) \), entropy \( S(t, h) \), and energy \( E(t, h) \)) are rather artificial; under a Legendre transformation we can take a system described by the Gibbs free energy \( G(T, H) \) to a system described by the microcanonical entropy \( S(E, H) \) or the Helmholtz free energy \( A(T, M) \). We conjecture that Legendre transformations will convert the analytic corrections to gauge corrections in our nomenclature. If so, it may be better to treat these on an equal footing, and term them both analytic corrections to scaling.

N2.5 We conjecture that a subset of the irrelevant eigenvalues for other critical points will similarly be given in terms of combinations of relevant eigenvalues. We conjecture that analytic changes in the results variables can remove these corrections to scaling. (Again, Nature does not tell you how you should measure your magnetization, just as it did not tell you how to measure your applied field.)
Many of the ideas here were prompted by an analysis of the onset of chaos from quasiperiodic motion [24, 29]. There the coordinate transformation taking the map to a simple rotation was numerically straightforward, central to understanding maps below the onset of chaos, and led to self-similar, nonanalytic maps at the onset of chaos. The coordinate transformations that connected two points on the critical manifold were not analytic, but numerically had a continuous first derivative. We conjecture that the corrections to scaling at the quasiperiodic onset of chaos will again separate into gauge and singular components.

A fair split? Number partitioning.10 (Computer science, Mathematics, Computation, Statistics) ③

A group of $N$ kids want to split up into two teams that are evenly matched. If the skill of each player is measured by an integer, can the kids be split into two groups such that the sum of the skills in each group is the same?

This is the number partitioning problem (NPP), a classic and surprisingly difficult problem in computer science. To be specific, it is NP −−complete—a category of problems for which no known algorithm can guarantee a resolution in a reasonable time (bounded by a polynomial in their size). If the skill $a_j$ of each kid $j$ is in the range $1 \leq a_j \leq 2^M$, the “size” of the NPP is defined as $NM$. Even the best algorithms will, for the hardest instances, take computer time that grows faster than any polynomial in $MN$, getting exponentially large as the system grows.

In this exercise, we shall explore connections between this numerical problem and the statistical mechanics of disordered systems. Number partitioning has been termed “the easiest hard problem”. It is genuinely hard numerically; unlike some other NP −−complete problems, there are no good heuristics for solving NPP (i.e., that work much better than a random search). On the other hand, the random NPP problem (the ensembles of all possible combinations of skills $a_j$) has many interesting features that can be understood with relatively straightforward arguments and analogies.

We start with the brute-force numerical approach to solving the problem.

(a) Write a function `ExhaustivePartition(S)` that inputs a list $S$ of $N$ integers, exhaustively searches through the $2^N$ possible partitions into two subsets, and returns the minimum cost (difference in the sums). Test your routine on the four sets [22] $S_1 = [10, 13, 23, 6, 20]$, $S_2 = [6, 4, 9, 14, 12, 3, 15, 15]$, $S_3 = [93, 58, 141, 209, 179, 48, 225, 228]$, and $S_4 = [2474, 1129, 1388, 3752, 821, 2082, 201, 739]$. Hint: $S_1$ has a balanced partition, and $S_4$ has a minimum cost of 48. You may wish to return the signs of the minimum-cost partition as part of the debugging process.

What properties emerge from studying ensembles of large partitioning problems? We find a phase transition. If the range of integers ($M$ digits in base two) is large and there are relatively few numbers $N$ to rearrange, it is unlikely that a perfect match

10This exercise draws heavily from [22, chapter 7]. Hints for the computations can be found at the book website [31].
can be found. (A random instance with \( N = 2 \) and \( M = 10 \) has a one chance in \( 2^{10} = 1,024 \) of a perfect match, because the second integer needs to be equal to the first.) If \( M \) is small and \( N \) is large it should be easy to find a match, because there are so many rearrangements possible and the sums are confined to a relatively small number of possible values. It turns out that it is the ratio \( \kappa = M/N \) that is the key; for large random systems with \( M/N > \kappa_c \) it becomes extremely unlikely that a perfect partition is possible, while if \( M/N < \kappa_c \) a fair split is extremely likely.

(b) Write a function \( \text{MakeRandomPartitionProblem}(N,M) \) that generates \( N \) integers randomly chosen from \( \{1, \ldots, 2^M\} \), rejecting lists whose sum is odd (and hence cannot have perfect partitions). Write a function \( p\text{Perf}(N,M,\text{trials}) \), which generates \( \text{trials} \) random lists and calls \( \text{ExhaustivePartition} \) on each, returning the fraction \( p_{\text{perf}} \) that can be partitioned evenly (zero cost). Plot \( p_{\text{perf}} \) versus \( \kappa = M/N \), for \( N = 3, 5, 7 \) and \( 9 \), for all integers \( M \) with \( 0 < \kappa = M/N < 2 \), using at least a hundred trials for each case. Does it appear that there is a phase transition for large systems where fair partitions go from probable to unlikely? What value of \( \kappa_c \) would you estimate as the critical point?

Should we be calling this a phase transition? It emerges for large systems; only in the “thermodynamic limit” where \( N \) gets large is the transition sharp. It separates two regions with qualitatively different behavior. The problem is much like a spin glass, with two kinds of random variables: the skill levels of each player \( a_j \) are fixed, “quenched” random variables for a given random instance of the problem, and the assignment to teams can be viewed as spins \( s_j = \pm 1 \) that can be varied (“annealed” random variables)\(^{11}\) to minimize the cost \( C = |\sum_j a_j s_j| \).

(c) Show that the square of the cost \( C^2 \) is of the same form as the Hamiltonian for a spin glass, \( H = \sum_{i,j} J_{ij} s_i s_j \). What is \( J_{ij} \)?

The putative phase transition in the optimization problem (part (b)) is precisely a zero-temperature phase transition for this spin-glass Hamiltonian, separating a phase with zero ground-state energy from one with nonzero energy in the thermodynamic limit.

We can understand both the value \( \kappa_c \) of the phase transition and the form of \( p_{\text{perf}}(N,M) \) by studying the distribution of possible “signed” costs \( E_a = \sum_j a_j s_j \). These energies are distributed over a maximum total range of \( E_{\text{max}} - E_{\text{min}} = 2 \sum_{j=1}^N a_j \leq 2N 2^M \) (all players playing on the plus team, through all on the minus team). For the bulk of the possible team choices \( \{s_j\} \), though, there will be some cancellation in this sum. The probability distribution \( P(E) \) of these energies for a particular NPP problem \( \{a_j\} \) is not simple, but the average probability distribution \( \langle P(E) \rangle \) over the ensemble of NPP problems can be estimated using the central limit theorem. (Remember that the central limit theorem states that the sum of \( N \) random variables with mean zero and

---

\(^{11}\) Quenched random variables are fixed terms in the definition of the system, representing dirt or disorder that was frozen in as the system was formed (say, by quenching the hot liquid material into cold water, freezing it into a disordered configuration). Annealed random variables are the degrees of freedom that the system can vary to explore different configurations and minimize its energy or free energy.
standard deviation $\sigma$ converges rapidly to a normal (Gaussian) distribution of standard deviation $\sqrt{N}\sigma$.)

(d) Estimate the mean and variance of a single term $s_ja_j$ in the sum, averaging over both the spin configurations $s_j$ and the different NPP problem realizations $a_j \in [1, \ldots, 2^M]$, keeping only the most important term for large $M$. (Hint: Approximate the sum as an integral, or use the explicit formula $\sum_1^K k^2 = K^3/3 + K^2/2 + K/6$ and keep only the most important term.) Using the central limit theorem, what is the ensemble-averaged probability distribution $P(E)$ for a team with $N$ players? Hint: Here $P(E)$ is nonzero only for even integers $E$, so for large $N$ $P(E) \approx (2/\sqrt{2\pi}\sigma) \exp(-E^2/2\sigma^2)$; the normalization is doubled.

Your answer to part (d) should tell you that the possible energies are mostly distributed among integers in a range of size $\sim 2^M$ around zero, up to a factor that goes as a power of $N$. The total number of states explored by a given system is $2^N$. So, the expected number of zero-energy states should be large if $N \gg M$, and go to zero rapidly if $N \ll M$. Let us make this more precise.

(e) Assuming that the energies for a specific system are randomly selected from the ensemble average $P(E)$, calculate the expected number of zero-energy states as a function of $M$ and $N$ for large $N$. What value of $\kappa = M/N$ should form the phase boundary separating likely from unlikely fair partitions? Does that agree well with your numerical estimate from part (b)?

The assumption we made in part (e) ignores the correlations between the different energies due to the fact that they all share the same step sizes $a_j$ in their random walks. Ignoring these correlations turns out to be a remarkably good approximation. We can use the random-energy approximation to estimate $p_{\text{perf}}$ that you plotted in part (b).

(f) In the random-energy approximation, argue that $p_{\text{perf}} = 1 - (1 - P(0))^{2^{N-1}}$. Approximating $(1 - A/L)^L \approx \exp(-A)$ for large $L$, show that

$$p_{\text{perf}}(\kappa, N) \approx 1 - \exp \left[ -\sqrt{\frac{3}{2\pi N}} 2^{-N(\kappa - \kappa_c)} \right].$$

(Rather than plotting the theory curve through each of your simulations from part (b), we change variables to $x = N(\kappa - \kappa_c) + (1/2) \log_2 N$, where the theory curve

$$p_{\text{scaling}}^{\text{scaling}}(x) = 1 - \exp \left[ -\sqrt{\frac{3}{2\pi}} 2^{-x} \right].$$

More precisely, we ignore correlations between the energies of different teams $s = \{s_i\}$, except for swapping the two teams $s \rightarrow -s$. This leads to the $N - 1$ in the exponent of the exponent for $p_{\text{perf}}$ in part (f). Notice that in this approximation, NPP is a form of the random energy model (REM, Exercise 3.19), except that we are interested in states of energy near $E = 0$, rather than minimum energy states.
is independent of $N$. If the theory is correct, your curves should converge to $p_{\text{perf}}^{\text{scaling}}(x)$ as $N$ becomes large.

(g) Reusing your simulations from part (b), make a graph with your values of $p_{\text{perf}}(x, N)$ versus $x$ and $p_{\text{perf}}^{\text{scaling}}(x)$. Does the random-energy approximation explain the data well?

Rigorous results show that this random-energy approximation gives the correct value of $\kappa_c$. The entropy of zero-cost states below $\kappa_c$, the probability distribution of minimum costs above $\kappa_c$ (of the Weibull form, Exercise 1.9), and the probability distribution of the $k$ lowest cost states are also correctly predicted by the random-energy approximation. It has also been shown that the correlations between the energies of different partitions vanish in the large $(N, M)$ limit so long as the energies are not far into the tails of the distribution, perhaps explaining the successes of ignoring the correlations.

What does this random-energy approximation imply about the computational difficulty of NPP? If the energies of different spin configurations (arrangements of kids on teams) were completely random and independent, there would be no better way of finding zero-energy states (fair partitions) than an exhaustive search of all states. This perhaps explains why the best algorithms for NPP are not much better than the exhaustive search you implemented in part (a); even among $\textbf{NP} - \text{-complete}$ problems, NPP is unusually unyielding to clever methods.\(^{13}\) It also lends credibility to the conjecture in the computer science community that $\textbf{P} \neq \textbf{NP} - \text{-complete}$; any polynomial-time algorithm for NPP would have to ingeniously make use of the seemingly unimportant correlations between energy levels.

N2.7 Cardiac dynamics.\(^{14}\) (Computation, Biology, Complexity) \footnote{The computational cost does peak near $\kappa = \kappa_c$. For small $\kappa \ll \kappa_c$ it is relatively easy to find a good solution, but this is mainly because there are so many solutions; even random search only needs to sample until it finds one of them. For $\kappa > \kappa_c$ showing that there is no fair partition becomes slightly easier as $\kappa$ grows $[22, \text{Fig. 7.3}].$} \footnote{This exercise and the associated software were developed in collaboration with Christopher Myers.}


The cardiac muscle is an excitable medium. In each heartbeat, a wave of excitation passes through the heart, compressing first the atria which pushes blood into the ventricles, and then compressing the ventricles pushing blood into the body. In this exercise we will study simplified models of heart tissue, that exhibit spiral waves similar to those found in arrhythmias.

An excitable medium is one which, when triggered from a resting state by a small stimulus, responds with a large pulse. After the pulse there is a refractory period during which it is difficult to excite a new pulse, followed by a return to the resting
state. The FitzHugh-Nagumo equations provide a simplified model for the excitable heart tissue:\textsuperscript{15}

\[
\frac{\partial V}{\partial t} = \nabla^2 V + \frac{1}{\epsilon} (V - V^3/3 - W)
\]

\[
\frac{\partial W}{\partial t} = \epsilon (V - \gamma W + \beta),
\]

(N2.19)

where \(V\) is the transmembrane potential, \(W\) is the recovery variable, and \(\epsilon = 0.2, \gamma = 0.8, \text{ and } \beta = 0.7\) are parameters. Let us first explore the behavior of these equations ignoring the spatial dependence (dropping the \(\nabla^2 V\) term, appropriate for a small piece of tissue). The dynamics can be visualized in the \((V,W)\) plane.

(a) Find and plot the nullclines of the FitzHugh-Nagumo equations: the curves along which \(dV/dt\) and \(dW/dt\) are zero (ignoring \(\nabla^2 V\)). The intersection of these two nullclines represents the resting state \((V^*,W^*)\) of the heart tissue. We apply a stimulus to our model by shifting the transmembrane potential to a larger value—running from initial conditions \((V^* + \Delta,W^*)\). Simulate the equations for stimuli \(\Delta\) of various sizes; plot \(V\) and \(W\) as a function of time \(t\), and also plot \(V(t)\) versus \(W(t)\) along with the nullclines. How big a stimulus do you need in order to get a pulse?

Excitable systems are often close to regimes where they develop spontaneous oscillations. Indeed, the FitzHugh-Nagumo equations are equivalent to the van der Pol equation (which arose in the study of vacuum tubes), a standard system for studying periodic motion.

(b) Try changing to \(\beta = 0.4\). Does the system oscillate? The threshold where the resting state becomes unstable is given when the nullcline intersection lies at the minimum of the \(V\) nullcline, at \(\beta_c = 7/15\).

Each portion of the tissue during a contraction wave down the heart is stimulated by its neighbors to one side, and its pulse stimulates the neighbor to the other side. This triggering in our model is induced by the Laplacian term \(\nabla^2 V\). We simulate the heart on a two-dimensional grid \(V(x_i,y_j,t), W(x_i,y_j,t)\), and calculate an approximate Laplacian by taking differences between the local value of \(V\) and values at neighboring points.

There are two natural choices for this Laplacian. The five-point discrete Laplacian is generalization of the one-dimensional second derivative, \(\partial^2 V/\partial x^2 \approx (V(x + dx) - 2V(x) + V(x - dx))/dx^2\):

\[
\nabla_5^2 V(x_i,y_i) \approx (V(x_i,y_{i+1}) + V(x_i,y_{i-1}) + V(x_{i+1},y_i) + V(x_{i-1},y_i) - 4V(x_i,y_i))/dx^2
\]

\[
\leftrightarrow \frac{1}{dx^2} \begin{pmatrix}
0 & 1 & 0 \\
1 & -4 & 1 \\
0 & 1 & 0
\end{pmatrix}
\]

\textsuperscript{15}Nerve tissue is also an excitable medium, modeled using different Hodgkin-Huxley equations.
where \( dx = x_{i+1} - x_i = y_{i+1} - y_i \) is the spacing between grid points and the last expression is the \textit{stencil} by which you multiply the point and its neighbors by to calculate the Laplacian. The nine-point discrete Laplacian has been fine-tuned for improved circularly symmetry, with stencil

\[
\nabla^2_{[9]} V(x_i, y_i) \leftrightarrow \frac{1}{dx^2} \begin{pmatrix}
1/6 & 2/3 & 1/6 \\
2/3 & -10/3 & 2/3 \\
1/6 & 2/3 & 1/6
\end{pmatrix}.
\]

\tag{N2.21}

We will simulate our partial-differential equation (PDE) on a square \( 100 \times 100 \) grid with a grid spacing \( dx = 1 \).\footnote{Smaller grids would lead to less grainy waves, but slow down the simulation a lot.} As is often done in PDEs, we will use the crude Euler time-step scheme \( V(t + \Delta) \approx V(t) + \Delta \partial V / \partial t \) (see Exercise 3.12): we find \( \Delta \approx 0.1 \) is the largest time step we can get away with. We will use “no-flow” boundary conditions, which we implement by setting the Laplacian terms on the boundary to zero (the boundaries, uncoupled from the rest of the system, will quickly turn to their resting state). If you are not supplied with example code that does the two-dimensional plots, you may find them at the book website [31].

(c) \textit{Solve eqn N2.19 for an initial condition equal to the fixed point \((V^*, W^*)\) except for a 10 \times 10 square at the origin, in which you should apply a stimulus \( \Delta = 3.0 \).} (Hint: Your simulation should show a pulse moving outward from the origin, disappearing as it hits the walls.)

If you like, you can mimic the effects of the sinoatrial (SA) node (your heart’s natural pacemaker) by stimulating your heart model periodically (say, with the same 10 \times 10 square). Realistically, your period should be long enough that the old beat finishes before the new one starts.

We can use this simulation to illustrate general properties of solving PDEs.

(d) \textbf{Accuracy}. \textit{Compare the five and nine-point Laplacians. Does the latter give better circular symmetry? Stability. After running for a while, double the time step \( \Delta \). How does the system go unstable? Repeat this process, reducing \( \Delta \) until just before it goes nuts. Do you see inaccuracies in the simulation that foreshadow the instability?}

This checkerboard instability is typical of PDEs with too high a time step. The maximum time step in this system will go as \( dx^2 \), the lattice spacing squared—thus to make \( dx \) smaller by a factor of two and simulate the same area, you need four times as many grid points and four times as many time points—giving us a good reason for making \( dx \) as large as possible (correcting for grid artifacts by using improved Laplacians). Similar but much more sophisticated tricks have been used recently to spectacularly increase the performance of lattice simulations of the interactions between quarks [11].

As mentioned above, heart arrhythmias are due to spiral waves. To generate spiral waves we need to be able to start up more asymmetric states—stimulating several rectangles at different times. Also, when we generate the spirals, we would like to
emulate electroshock therapy by applying a stimulus to a large region of the heart. We can do both by writing code to interactively stimulate a whole rectangle at one time. Again, the code you have obtained from us should have hints for how to do this.

(e) Add the code for interactively stimulating a general rectangle with an increment to $V$ of size $\Delta = 3$. Play with generating rectangles in different places while other pulses are going by: make some spiral waves. Clear the spirals by giving a stimulus that spans the system.

There are several possible extensions of this model, several of which involve giving our model spatial structure that mimics the structure of the heart. (One can introduce regions of inactive “dead” tissue. One can introduce the atrium and ventricle compartments to the heart, with the SA node in the atrium and an AV node connecting the two chambers . . . ) Niels Otani has an exercise with further explorations of a number of these extensions, which we link to from the Cardiac Dynamics website.

N2.8 Quantum dissipation from phonons. (Quantum) ②

fig. N2.2 Atomic tunneling from a tip. Any internal transition among the atoms in an insulator can only exert a force impulse (if it emits momentum, say into an emitted photon), or a force dipole (if the atomic configuration rearranges); these lead to nonzero phonon overlap integrals only partially suppressing the transition. But a quantum transition that changes the net force between two macroscopic objects (here a surface and a STM tip) can lead to a change in the net force (a force monopole). We ignore here the surface, modeling the force as exerted directly into the center of an insulating elastic medium.\textsuperscript{17} See “Atomic Tunneling from a STM/AFM Tip: Dissipative Quantum Effects from Phonons” Ard A. Louis and James P. Sethna, Phys. Rev. Lett. 74, 1363 (1995), and “Dissipative tunneling and orthogonality catastrophe in molecular transistors”, S. Braig and K. Flensberg, Phys. Rev. B 70, 085317 (2004).

Electrons cause overlap catastrophes (X-ray edge effects, the Kondo problem, macroscopic quantum tunneling); a quantum transition of a subsystem coupled to an electron bath ordinarily must emit an infinite number of electron-hole excitations because the bath states before and after the transition have zero overlap. This is often called an infrared catastrophe (because it is low-energy electrons and holes that cause the zero overlap), or an orthogonality catastrophe (even though the two bath states aren’t just orthogonal, they are in different Hilbert spaces). Phonons typically do not produce overlap catastrophes (Debye–Waller, Frank–Condon, Mössbauer). This difference is usually attributed to the fact that there are many more low-energy electron-hole pairs
(a constant density of states) than there are low-energy phonons ($\omega_k \sim ck$, where $c$ is the speed of sound and the wave-vector density goes as $(V/2\pi)^3d^3k$).

However, the coupling strength to the low energy phonons has to be considered as well. Consider a small system undergoing a quantum transition which exerts a net force at $x = 0$ onto an insulating crystal:

$$H = \sum_k p_k^2/2m + 1/2m\omega_k^2 q_k^2 + F \cdot u_0.$$  \hspace{1cm} (N2.22)

Let us imagine a kind of scalar elasticity, to avoid dealing with the three phonon branches (two transverse and one longitudinal); we thus naively write the displacement of the atom at lattice site $x_n$ as $u_n = (1/\sqrt{N}) \sum_k q_k \exp(-ikx_n)$ (with $N$ the number of atoms), so $q_k = (1/\sqrt{N}) \sum_n u_n \exp(ikx_n)$.

Substituting for $u_0$ in the Hamiltonian and completing the square, find the displacement $\Delta_k$ of each harmonic oscillator. (Physically, the force $F$ adds a small linear term to the phonon mode with wavevector $k$, whose minimum becomes displaced by some amount $\Delta_k$.) Let $|F\rangle$ be the ground state of the harmonic oscillators under the force $F$. Write the formula for the likelihood $\langle F|0 \rangle$ that the phonons will all end in their ground states, as a product over $k$ of the phonon overlap integral $\exp(-\Delta_k^2/8\sigma_k^2)$ (with $\sigma_k = \sqrt{\hbar/2m\omega_k}$ the zero-point motion in that mode). Converting the product to the exponential of a sum, and the sum to an integral $\sum_k \sim (V/(2\pi)^3 \int d^3k$, do we observe an overlap catastrophe?

Note that you’ve calculated the probability of a zero-phonon transition—the likelihood that the quantum transition can happen without emitting any phonons is zero. But the same argument shows that there is zero probability of emitting one phonon, or any finite number of phonons. The only allowed transitions emit an infinite number of low-energy phonons. The initial and final ground states are in different Hilbert spaces—no finite number of excitations can connect them.

N2.9 Ising lower critical dimension. (Dimension dependence) \(3\)

What is the lower critical dimension of the Ising model? If the total energy $\Delta E$ needed to destroy long-range order is finite as the system size $L$ goes to infinity, and the associated entropy grows with system size, then surely long-range order is possible only at zero temperature.

(a) Ising model in $D$ dimensions. Consider the Ising model in dimension $D$ on a hypercubic lattice of length $L$ on each side. Estimate the energy\(^{18}\) needed to create a domain wall splitting the system into two equal regions (one spin up, the other spin down). In what dimension will this wall have finite energy as $L \to \infty$? Suggest a bound for the lower critical dimension of the Ising model.

The scaling at the lower critical dimension is often unusual, with quantities diverging in ways different from power laws as the critical temperature $T_c$ is approached.

\(^{18}\)Energy, not free energy! Think about $T = 0$. 

(b) **Correlation length in 1D Ising model.** Estimate the number of domain walls at temperature $T$ in the 1D Ising model. How does the correlation length $\xi$ (the distance between domain walls) grow as $T \to T_c = 0$? (Hint: Change variables to $\eta_i = S_i S_{i+1}$, which is $-1$ if there is a domain wall between sites $i$ and $i+1$.) The correlation exponent $\nu$ satisfying $\xi \sim (T - T_c)^{-\nu}$ is $1, \sim 0.63, \text{ and } 1/2$ in dimensions $D = 2, 3, \text{ and } \geq 4$, respectively. Is there an exponent $\nu$ governing this divergence in one dimension? How does $\nu$ behave as $D \to 1$?

**N2.10 XY lower critical dimension and the Mermin-Wagner Theorem.** (Dimension dependence) ③

Consider a model of continuous unit-length spins (e.g., XY or Heisenberg) on a $D$-dimensional hypercubic lattice of length $L$. Assume a nearest-neighbor ferromagnetic bond energy

$$-JS_i \cdot S_j.$$  \[ \text{(N2.23)} \]

Estimate the energy needed to twist the spins at one boundary $180^\circ$ with respect to the other boundary (the energy difference between periodic and antiperiodic boundary conditions along one axis). In what dimension does this energy stay finite in the thermodynamic limit $L \to \infty$? Suggest a bound for the lower critical dimension for the emergence of continuous broken symmetries in models of this type.

Note that your argument produces only one thick domain wall (unlike the Ising model, where the domain wall can be placed in a variety of places). If in the lower critical dimension its energy is fixed as $L \to \infty$ at a value large compared to $k_B T$, one could imagine most of the time the order might maintain itself across the system. The actual behavior of the XY model in its lower critical dimension is subtle.

On the one hand, there cannot be long-range order. This can be seen convincingly, but not rigorously, by estimating the effects of fluctuations at finite temperature on the order parameter, within linear response. Pierre Hohenberg, David Mermin and Herbert Wagner proved it rigorously (including nonlinear effects) using an inequality due to Bogoliubov. One should note, though, that the way this theorem is usually quoted (“continuous symmetries cannot be spontaneously broken at finite temperatures in one and two dimensions”) is too general. In particular, for two-dimensional crystals one has long-range order in the crystalline orientations, although one does not have long-range broken translational order.

On the other hand, the XY model does have a phase transition in its lower critical dimension at a temperature $T_c > 0$. The high-temperature phase is a traditional paramagnetic phase, with exponentially decaying correlations between orientations as the distance increases. The low-temperature phase indeed lacks long-range order, but it does have a stiffness—twisting the system (as in your calculation above) by $180^\circ$ costs a free energy that goes to a constant as $L \to \infty$. In this stiff phase the spin–spin correlations die away not exponentially, but as a power law.

The corresponding *Kosterlitz-Thouless phase transition* has subtle, fascinating scaling properties. Interestingly, the defect that destroys the stiffness (a vortex) in the
Kosterlitz–Thouless transition does not have finite energy as the system size $L$ gets large. We shall see that its energy grows $\sim \log L$, while its entropy grows $\sim T \log L$, so entropy wins over energy as the temperature rises, even though the latter is infinite.

N2.11 Long-range Ising. (Dimension dependence)

The one-dimensional Ising model can have a finite-temperature transition if we give each spin an interaction with distant spins.

**Long-range forces in the 1d Ising model.** Consider an Ising model in one dimension, with long-range ferromagnetic bonds

$$\mathcal{H} = \sum_{i>j} J \sigma_i \sigma_j.$$  \hspace{1cm} (N2.24)

For what values of $\sigma$ will a domain wall between up- and down-spins have finite energy? Suggest a bound on $\sigma$ analogous to the lower critical dimension (the maximum power $\sigma$, above which a ferromagnetic state is only possible when the temperature is zero). (Hint: Approximate the sum by a double integral. Avoid $i = j$.)

The long-range 1D Ising model at the lower critical power law has a transition that is closely related to the Kosterlitz-Thouless transition. It is in the same universality class as the famous (but obscure) Kondo problem in quantum phase transitions. And it is less complicated to think about and less complicated to calculate with than either of these other two cases.

N2.12 Equilibrium Crystal Shapes. (Condensed matter)

What is the equilibrium shape of a crystal? There are nice experiments on single crystals of salt, gold, and lead crystals (see http://www.lassp.cornell.edu/sethna/Crystal Shapes/Equilibrium_Crystal_Shapes.html). They show beautiful faceted shapes, formed by carefully annealing single crystals to equilibrium at various temperatures. The physics governing the shape involves the anisotropic surface tension $\gamma(\hat{n})$ of the surface, which depends on the orientation $\hat{n}$ of the local surface with respect to the crystalline axes.

We can see how this works by considering the problem for atoms on a 2D square lattice with near-neighbor interactions (a lattice gas which we map in the standard way onto a conserved-order parameter Ising model). Here $\gamma(\theta)$ becomes the line tension between the up and down phases—the interfacial free energy per unit length between an up-spin and a down-spin phase. We draw heavily on Craig Rottman and Michael Wortis, Phys. Rev. B 24, 6274 (1981), and on W. K. Burton, N. Cabrera and F. C. Frank, Phil. Trans. R. Soc. Lond. A 243 299-358 (1951).

(a) Interfacial free energy, $T = 0$. Consider an interface at angle $\theta$ between up-spins and down-spins. Show that the energy cost per unit length of the interface is

$$\gamma_0(\theta) = 2J(\cos(|\theta|) + \sin(|\theta|)).$$  \hspace{1cm} (N2.25)
where length is measured in lattice spacings.

(b) Interfacial free energy, low $T$. Consider an interface at angle $\theta = \arctan N/M$, connecting the origin to the point $(M, N)$. At zero temperature, this will be an ensemble of staircases, with $N$ steps upward and $M$ steps forward. Show that the total number of such staircases is $(M + N)!/(M!N!)$. Hint: The number of ways of putting $k$ balls into $\ell$ jars allowing more than one ball per jar (the “number of combinations with repetition”) is $(k + \ell - 1)!/(k!(\ell - 1)!))$. Look up the argument. Using Stirling’s formula, show that the entropy per unit length is

$$s_0(\theta) = k_B \left( \frac{1}{2} \log \left( \frac{\cos(|\theta|) + \sin(|\theta|)}{\cos(|\theta|) \log(\cos(|\theta|))} \right) - \cos(|\theta|) \log(\cos(|\theta|)) - \sin(|\theta|) \log(\sin(|\theta|)) \right).$$

(c) Wolff construction. Using the energy of eqn N2.25 and approximating the entropy at low temperatures with the zero-temperature form eqn N2.26, plot the Wulff shape.

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19If you put on an external field favoring up-spins, then large clusters will grow and small clusters will shrink. The borderline cluster size is the critical droplet (see Entropy, Order Parameters, and Complexity, section 11.3). Indeed, the critical droplet will in general share the equilibrium crystal shape.

20See Burton, Cabrera, and Frank, Appendix D. This is their equation D4, with $p \propto \gamma(\theta)$ given by equation D7.
for $T = 0.01, 0.1, 0.2, 0.4,$ and $0.8J$ for one quadrant ($0 < \theta < \pi/2$). Hint: Ignore the parts of the face outside the quadrant; they are artifacts of the low temperature approximation. Does the shape become circular\(^{21}\) near $T_c = 2J/\log(1 + \sqrt{2}) \sim 2.269J$? Why not? If you’re ambitious, Rottman and Wortis’s article above gives the exact interfacial free energy for the 2D Ising model, which should fix this problem. What is the shape at low temperatures, where our approximation is good? Do we ever get true facets? The approximation of the interface as a staircase is a solid-on-solid model, which ignores overhangs. It is a good approximation at low temperatures.

Our model does not describe faceting—one needs three dimensions to have a roughening transition, below which there are flat regions on the equilibrium crystal shape.\(^ {22}\)

\(^{21}\)This is the emergent spherical symmetry at the Ising model critical point.

\(^{22}\)Flat regions demand that the free energy for adding a step onto the surface become infinite. Steps on the surfaces of three-dimensional crystals are long, and if they have positive energy per unit length the surface is faceted. To get an infinite energy for a step on a two-dimensional surface you need long-range interactions.
Chapter 3

Numerical Methods

These exercises cover numerical methods, and are designed to complement the text *Numerical Recipes*, by William H. Press, Saul A. Teukolsky, William T. Vetterling, and Brian P. Flannery.

N3.1 Condition number and accuracy.¹ (Numerical) ³

You may think this exercise, with a 2x2 matrix, hardly demands a computer. However, it introduces tools for solving linear equations, condition numbers, singular value decomposition, all while illustrating subtle properties of matrix solutions. Use whatever linear algebra packages are provided in your software environment.

Consider the equation \( Ax = b \), where

\[
A = \begin{pmatrix}
0.780 & 0.563 \\
0.913 & 0.659
\end{pmatrix}
\quad \text{and} \quad
b = \begin{pmatrix}
0.217 \\
0.254
\end{pmatrix}.
\]  \( \text{(N3.1)} \)

The exact solution is \( x = (1, -1) \). Consider the two approximate solutions \( x_\alpha = (0.999, -1.001) \) and \( x_\beta = (0.341, -0.087) \).

(a) Compute the residuals \( r_\alpha \) and \( r_\beta \) corresponding to the two approximate solutions. (The residual is \( r = b - Ax \).) Does the more accurate solution have the smaller residual?

(b) Compute the condition number² of \( A \). Does it help you understand the result of (a)? (Hint: \( V^T \) maps the errors \( x_\alpha - x \) and \( x_\beta - x \) into what combinations of the two singular values?)

(c) Use a black-box linear solver to solve for \( x \). Subtract your answer from the exact one. Do you get within a few times the machine accuracy of \( 2.2 \times 10^{-16} \)? Is the problem

¹Adapted from Saul Teukolsky, 2003.
²See section 2.6.2 for a technical definition of the condition number, and how it is related to singular value decomposition. Look on the Web for the more traditional definition(s), and how they are related to the accuracy.
the accuracy of the solution, or rounding errors in calculating $A$ and $b$? (Hint: try calculating $Ax - b$.)

N3.2 **Sherman–Morrison formula.** (Numerical) 3

Consider the 5x5 matrices

$$T = \begin{pmatrix} E - t & t & 0 & 0 & 0 \\ t & E & t & 0 & 0 \\ 0 & t & E & t & 0 \\ 0 & 0 & t & E & t \\ 0 & 0 & 0 & t & E - t \end{pmatrix} \quad (N3.2)$$

and

$$C = \begin{pmatrix} E & t & 0 & 0 & t \\ t & E & t & 0 & 0 \\ 0 & t & E & t & 0 \\ 0 & 0 & t & E & t \\ t & 0 & 0 & t & E \end{pmatrix} \quad (N3.3)$$

These matrices arise in one-dimensional models of crystals.3 The matrix $T$ is *tridiagonal*: its entries are zero except along the central diagonal and the entries neighboring the diagonal. Tridiagonal matrices are fast to solve; indeed, many routines will start by changing basis to make the array tridiagonal. The matrix $C$, on the other hand, has a nice periodic structure: each basis element has two neighbors, with the first and last basis elements now connected by $t$ in the upper-right and lower-left corners. This periodic structure allows for analysis using Fourier methods (Bloch waves and $k$-space).

For matrices like $C$ and $T$ which differ in only a few matrix elements4 we can find $C^{-1}$ from $T^{-1}$ efficiently using the Sherman-Morrison formula (section 2.7).

*Compute the inverse5 of $T$ for $E = 3$ and $t = 1$. Compute the inverse of $C$. Compare the difference $\Delta = T^{-1} - C^{-1}$ with that given by the Sherman-Morrison formula

$$\Delta = \frac{T^{-1} u \otimes v T^{-1}}{1 + v \cdot T^{-1} \cdot u}.$$ \quad (N3.4)

---

3As the Hamiltonian for electrons in a one-dimensional chain of atoms, $t$ is the hopping matrix element and $E$ is the on-site energy. As the potential energy for longitudinal vibrations in a one-dimensional chain, $E = -2t = K$ is the spring constant between two neighboring atoms. The tridiagonal matrix $T$ corresponds to a kind of free boundary condition, while $C$ corresponds in both cases to periodic boundary conditions.

4More generally, this works whenever the two matrices differ by the outer product $u \otimes v$ of two vectors. By taking the two vectors to each have one nonzero component $u_i = u \delta_{ia}, v_j = v \delta_{ib}$, the matrices differ at one matrix element $\Delta_{ab} = uv$; for our matrices $u = v = 1, 0, 0, 0, 1$ (see section 2.7.2).

5Your software environment should have a *solver* for tridiagonal systems, rapidly giving $u$ in the equation $T \cdot u = r$. It likely will not have a special routine for inverting tridiagonal matrices, but our matrix is so small it is not important.
N3.3 Methods of interpolation. (Numerical)  
We’ve implemented four different interpolation methods for the function \( \sin(x) \) on the interval \((-\pi, \pi)\). On the left, we see the methods using the five points \( \pm\pi, \pm\pi/2, \) and \(0\); on the right we see the methods using ten points. The graphs show the interpolation, its first derivative, its second derivative, and the error. The four interpolation methods we have implemented are (1) Linear, (2) Polynomial of degree three \((M = 4)\), (3) Cubic spline, and (4) Barycentric rational interpolation. Which set of curves (A, B, C, or D) in Fig. N3.1 corresponds with which method?

N3.4 Numerical definite integrals. (Numerical)  
In this exercise we will integrate the function you graphed in the first, warmup exercise:

\[
F(x) = \exp(-6 \sin(x)). 
\]  
(N3.5)

As discussed in Numerical Recipes, the word integration is used both for the operation that is inverse to differentiation, and more generally for finding solutions to differential equations. The old-fashioned term specific to what we are doing in this exercise is quadrature.

(a) Black-box. Using a professionally written black-box integration routine of your choice, integrate \( F(x) \) between zero and \( \pi \). Compare your answer to the analytic integral\(^6\) \((\approx 0.34542493760937693)\) by subtracting the analytic form from your numerical result. Read the documentation for your black box routine, and describe the combination of algorithms being used.

(b) Trapezoidal rule. Implement the trapezoidal rule with your own routine. Use it to calculate the same integral as in part (a). Calculate the estimated integral \( \text{Trap}(h) \) for \( N + 1 \) points spaced at \( h = \pi/N, \) with \( N = 1, 2, 4, \ldots, 2^{10} \). Plot the estimated integral versus the spacing \( h \). Does it extrapolate smoothly to the true value as \( h \to 0 \)? With what power of \( h \) does the error vanish? Replot the data as \( \text{Trap}(h) \) versus \( h^2 \). Does the error now vanish linearly?

Numerical Recipes tells us that the error is an even polynomial in \( h \), so we can extrapolate the results of the trapezoidal rule using polynomial interpolation in powers of \( h^2 \).

(c) Simpson’s rule (paper and pencil). Consider a linear fit \((i.e., A+Bh^2)\) to two points at \( 2h_0 \) and \( h_0 \) on your \( \text{Trap}(h) \) versus \( h^2 \) plot. Notice that the extrapolation to \( h \to 0 \) is \( A \), and show that \( A \) is \( 4/3 \text{Trap}(h_0) - 1/3 \text{Trap}(2h_0) \). What is the net weight associated with the even points and odd points? Is this Simpson’s rule?

(d) Romberg integration. Apply \( M \)-point polynomial interpolation (here extrapolation) to the data points \( \{h^2, \text{Trap}(h)\} \) for \( h = \pi/2, \ldots, \pi/2^M \), with values of \( M \) between two and ten. (Note that the independent variable is \( h^2 \).) Make a log–log plot of the absolute value of the error versus \( N = 2^M \). Does this extrapolation improve convergence?

\(^6\pi(\text{BesselI}[0, 6] - \text{StruveL}[0, 6]), \) according to Mathematica.
Interpolated \( \sin(x) \approx \) 

First derivative 

Second derivative 

Error \( \sin(x) - \) interpolation 

Fig. N3.1 Interpolation methods.
(e) **Gaussian Quadrature.** Implement Gaussian quadrature with $N$ points optimally chosen on the interval $(0, \pi)$, with $N = 1, 2, \ldots, 5$. (You may find the points and the weights appropriate for integrating functions on the interval $(-1, 1)$ on the course website; you will need to rescale them for use on $(0, \pi)$.) Make a log–log plot of the absolute value of your error as a function of the number of evaluation points $N$, along with the corresponding errors from the trapezoidal rule and Romberg integration.

(f) **Integrals of periodic functions.** Apply the trapezoidal rule to integrate $F(x)$ from zero to $2\pi$, and plot the error on a log plot (log of the absolute value of the error versus $N$) as a function of the number of points $N$ up to $N = 20$. (The true value should be around $422.44623805153909946$.) **Why does it converge so fast?** (Hint: Don’t get distracted by the funny alternation of accuracy between even and odd points.)

The location of the Gauss points depend upon the class of functions one is integrating. In part (e), we were using Gauss-Legendre quadrature, appropriate for functions which are analytic at the endpoints of the range of integration. In part (f), we have a function with **periodic boundary conditions**. For functions with periodic boundary conditions, the end-points are no longer special. What corresponds to Gaussian quadrature for periodic functions is just the trapezoidal rule: equally-weighted points at equally spaced intervals.

**N3.5 Numerical derivatives.** (Rounding errors, Accuracy) 

Calculate the numerical first derivative of the function $y(x) = \sin(x)$, using the centered two-point formula $\frac{dy}{dx} \sim (y(x+h) - y(x-h))/(2h)$, and plot the error $y'(x) - \cos(x)$ in the range $(-\pi, \pi)$ at 100 points. (Do a good job! Use the step-size $h$ described in Numerical Recipes section 5.7 to optimize the sum of the truncation error and the rounding error. Also, make sure that the step-size $h$ is exactly representable on the machine.) **How does your actual error compare to the fractional error estimate given in NR section 5.7?** Calculate and plot the numerical second derivative using the formula

$$
\frac{d^2y}{dx^2} \sim \frac{y(x+h) - 2y(x) + y(x-h)}{h^2},
$$

again optimizing $h$ and making it exactly representable. Estimate your error again, and compare to the observed error.

**N3.6 Summing series.** (Efficiency)

Write a routine to calculate the sum

$$
s_n = \sum_{j=0}^{n} (-1)^j \frac{1}{2j+1}.
$$

As $n \to \infty$, $s_\infty = \pi/4$. About how many terms do you need to sum to get convergence to within $10^{-7}$ of this limit? Now try using Aitken’s $\Delta^2$ process to accelerate the convergence:

$$
s_n' = s_n - \frac{(s_{n+1} - s_n)^2}{s_{n+2} - 2s_{n+1} + s_n}.
$$
About how many terms do you need with Aitken’s method to get convergence to within $10^{-7}$?

N3.7 Random histograms. (Random numbers) ②

(a) Investigate the random number generator for your system of choice. What is its basic algorithm? Its period?

(b) Plot a histogram with 100 bins, giving the normalized probability density of 100,000 random numbers sampled from (a) a uniform distribution in the range $0 < x < 2\pi$, (b) an exponential distribution $p(x) = 6 \exp(-6x)$, and (c) a normal distribution of mean $\bar{x} = 3\pi/2$ and standard deviation $\sigma = 1/\sqrt{6}$. Before each plot, set the seed of your random number generator. Do you now get the same plot when you repeat?

N3.8 Monte Carlo integration. (Random numbers, Robust algorithms) ③

How hard can numerical integration be? Suppose the function $f$ is wildly nonanalytic, or has a peculiar or high-dimensional domain of integration? In the worst case, one can always try Monte Carlo integration. The basic idea is to pepper points at random in the integration interval. The integration volume times the average of the function $V\langle f \rangle$ is the estimate of the integral.

As one might expect, the expected error in the integral after $N$ evaluations is given by $1/\sqrt{N-1}$ times the standard deviation of the sampled points (NR equation 7.7.1).

(a) Monte Carlo in one dimensional integrals. Use Monte Carlo integration to estimate the integral of the function introduced in the preliminary exercises

$$y(x) = \exp(-6\sin(x))$$  \hspace{1cm} (N3.9)

over $0 \leq x < 2\pi$. (The correct value of the integral is around 422.446.) How many points do you need to get 1% accuracy? Answer this last question both by experimenting with different random number seeds, and by calculating the expected number from the standard deviation. (You may use the fact that $\langle y^2 \rangle = (1/2\pi) \int_0^{2\pi} y^2(x)dx \approx 18,948.9$.)

Monte Carlo integration is not the most efficient method for calculating integrals of smooth functions like $y(x)$. Indeed, since $y(x)$ is periodic in the integration interval, equally spaced points weighted equally (the trapezoidal rule) gives exponentially rapid convergence; it takes only nine points to get 1% accuracy. Even for smooth functions, though, Monte Carlo integration is useful in high dimensions.

(b) Monte Carlo in many dimensions (No detailed calculations expected). For a hypothetical ten-dimensional integral, if we used a regular grid with nine points along each axis, how many function evaluations would we need for equivalent accuracy? Does the number of Monte Carlo points needed depend on the dimension of the space, presuming (perhaps naively) that the variance of the function stays fixed?

Our function $y(x)$ is quite close to a Gaussian. (Why? Taylor expand $\sin(x)$ about $x = 3\pi/2$.) We can use this to do importance sampling. The idea is to evaluate the integral of $h(x)g(x)$ by randomly sampling $h$ with probability $g$, picking $h(x) = y(x)/g(x)$. The
variance is then $\langle h^2 \rangle - \langle h \rangle^2$. In order to properly sample the tail near $x = \pi/2$, we should mix a Gaussian and a uniform distribution:

$$g(x) = \frac{\epsilon}{2\pi} + \frac{1 - \epsilon}{\sqrt{\pi/3}} \exp(-6(x - 3\pi/2)^2/2).$$

(N3.10)

I found minimum variance around $\epsilon = 0.005$.

(c) Importance Sampling (Optional for 480). Generate 1,000 random numbers with probability distribution $g$.\(^{7}\) Use these to estimate the integral of $y(x)$. How accurate is your answer?

N3.9 **Washboard potential.** (Solving) \(^{2}\)

Consider a washboard potential\(^{8}\)

$$V(r) = A_1 \cos(r) + A_2 \cos(2r) - Fr$$

(N3.11)

with $A_1 = 5$, $A_2 = 1$, and $F$ initially equal to 1.5.

(a) Plot $V(r)$ over $(-10, 10)$. Numerically find the local maximum of $V$ near zero, and the local minimum of $V$ to the left (negative side) of zero. What is the potential energy barrier for moving from one well to the next in this potential?

Usually finding the minimum is only a first step—one wants to explore how the minimum moves and disappears.

(b) Increasing the external tilting field $F$, graphically roughly locate the field $F_c$ where the barrier disappears, and the location $r_c$ at this field where the potential minimum and maximum merge. (This is a saddle-node bifurcation.) Give the criterion on the first derivative and the second derivative of $V(r)$ at $F_c$ and $r_c$. Using these two equations, numerically use a root-finding routine to locate the saddle-node bifurcation $F_c$ and $r_c$.

N3.10 **Sloppy minimization.** (Statistics) \(^{3}\)

"With four parameters I can fit an elephant. With five I can make it waggle its trunk." This statement, attributed to many different sources (from Carl Friedrich Gauss to Fermi), reflects the problems found in fitting multiparameter models to data. One almost universal problem is sloppiness—the parameters in the model are poorly constrained by the data.

---

\(^{7}\)Take $(1 - \epsilon)M$ Gaussian random numbers and $\epsilon M$ random numbers uniformly distributed on $(0, 2\pi)$. The Gaussian has a small tail that extends beyond the integration range $(0, 2\pi)$, so the normalization of the second term in the definition of $g$ is not quite right. You can fix this by simply throwing away any samples that include points outside the range.

\(^{8}\)A washboard is what people used to hand-wash clothing. It is held at an angle, and has a series of corrugated ridges; one holds the board at an angle and rubs the wet clothing on it. Washboard potentials arise in the theory of superconducting Josephson junctions, in the motion of defects in crystals, and in many other contexts.
Consider the classic ill-conditioned problem of fitting exponentials to radioactive decay data. If you know that at $t = 0$ there are equal quantities of $N$ radioactive materials with half-lives $\theta_\alpha$, the radioactivity that you would measure is

$$y_\theta(t) = \sum_{\alpha=0}^{N-1} \theta_\alpha \exp(-\theta_\alpha t).$$  (N3.12)

Now, suppose you do not know the decay rates $\theta_\alpha$. Can you reconstruct them by fitting the data to experimental data $d(t)$?

Start with just two radioactive decay elements $N = 2$. Suppose the actual decay constants for $d(t)$ are $\theta_0 = [1, 2]$ (so the data fall on the curve $d(t) = \exp(-t) + 2 \exp(-2t) = y_{\theta_0}(t)$). For convenience, suppose we have perfect data at all times, with uniform error bars, so the cost is an integral over all times of the square of the error

$$C[\theta] = \int_0^\infty (y_\theta(t) - d(t))^2 \, dt.$$  (N3.13)

(a) Draw a contour plot of $C$ in the square $0.5 < \theta_\alpha < 2.5$, with contours at $C = \{2^{-12}, 2^{-11}, \ldots, 2^0\}$. You may need to set the number of grid points per side to 40 to see the two minima.

One can see from the contour plot that measuring the two rate constants separately would be a challenge. This is because the two exponentials have similar shapes, so increasing one decay rate and decreasing the other can almost perfectly compensate for one another.

(b) If we assume both elements decay with the same decay rate $\theta = \theta_0 = \theta_1$, minimize the cost to find the optimum choice for $\theta$. Where is this point on the contour plot? Plot $d(t)$ and $y(t)$ with this single-exponent best fit on the same graph, over $0 < t < 2$. Do you agree that it would be difficult to distinguish these two fits?

This problem can become much more severe in higher dimensions. The banana-shaped ellipses in your contour plot can become needle-like, with aspect ratios of more than a thousand to one (about the same as a human hair). The relative widths of the ellipses are given by the square roots of the eigenvalues of the cost $C$.

(c) For our exercise, where the data are perfectly fit by $\theta = \theta_0$, show that the cost Hessian is a continuous integral

$$H_{\alpha\beta} = (J^T J)_{\alpha\beta} = J_{t\alpha} J_{t\beta} = \int_0^\infty J(t, \alpha) J(t, \beta) \, dt$$  (N3.14)

where the Jacobian is now the $\infty \times N$ “matrix” $J(t, \alpha) = \exp(-\theta_\alpha t)(1 - \theta_\alpha t)$.

(d) Write a routine to calculate $H(\theta)$ by doing the indefinite integral in eqn N3.14. Find the eigenvalues and the eigenvectors for the cost Hessian $H$ for your plot in part (b), evaluated at $\theta_0$, and check them against your contour plot. What is the ratio of the
long axis to the short axis, as predicted from your eigenvalues? For a sum of nine exponentials, with \( \theta_0 = [1, 2, 3, \ldots, 9] \), construct the Hessian, find its eigenvalues. By what factor does each successive eigenvalue shrink? Are they sloppy (roughly equally spaced in log)?

N3.11 Sloppy monomials.\(^9\) (Statistics) \(^3\)

The same function \( f(x) \) can be approximated in many ways. Indeed, the same function can be fit in the same interval by the same type of function in several different ways! For example, in the interval \([0, 1]\), the function \( \sin(2\pi x) \) can be approximated (badly) by a fifth-order Taylor expansion, a Chebyshev polynomial, or a least-squares (Legendre\(^{10}\)) fit:

\[
\begin{align*}
f(x) &= \sin(2\pi x) \\
f_{\text{Taylor}} &\approx 0.000 + 6.283x + 0.000x^2 - 41.342x^3 \\
&\quad + 0.000x^4 + 81.605x^5 \\
f_{\text{Chebyshev}} &\approx 0.007 + 5.652x + 9.701x^2 - 95.455x^3 \\
&\quad + 133.48x^4 - 53.39x^5 \\
f_{\text{Legendre}} &\approx 0.016 + 5.410x + 11.304x^2 - 99.637x^3 \\
&\quad + 138.15x^4 - 55.26x^5
\end{align*}
\]

It is not a surprise that the best fit polynomial differs from the Taylor expansion, since the latter is not a good approximation. But it is a surprise that the last two polynomials are so different. The maximum error for Legendre is less than 0.02, and for Chebyshev is less than 0.01, even though the two polynomials differ by

\[
\text{Chebyshev} - \text{Legendre} = \begin{align*}
&- 0.009 + 0.242x - 1.603x^2 \\
&\quad + 4.182x^3 - 4.67x^4 + 1.87x^5
\end{align*}
\]

a polynomial with coefficients two hundred times larger than the maximum difference! This flexibility in the coefficients of the polynomial expansion is remarkable. We can study it by considering the dependence of the quality of the fit on the parameters. Least-squares (Legendre) fits minimize a cost \( C^{\text{poly}} \), the integral of the squared difference between the polynomial and the function:

\[
C^{\text{poly}} = (1/2) \int_0^1 (f(x) - y_\theta(x))^2 \, dx,
\]

\[
y_\theta(x) = \sum_{m=0}^{M-1} \theta_m x^m
\]

\(^9\)Thanks to Joshua Waterfall, whose research is described here.

\(^{10}\)The orthogonal polynomials used for least-squares fits on \([-1,1]\) are the Legendre polynomials, assuming continuous data points. Were we using orthogonal polynomials for this exercise, we would need to shift them for use in \([0,1]\).
How quickly does this cost increase as we move the \( M \) parameters \( \theta_m \) away from their best-fit values? Varying any one monomial coefficient will of course make the fit bad. But apparently certain coordinated changes of coefficients do not cost much—for example, the difference between least-squares and Chebyshev fits given in eqn N3.15.

How should we explore the dependence in arbitrary directions in parameter space? We can use the eigenvalues of the Hessian to see how sensitive the fit is to moves along the various eigenvectors...

(a) Note that the first derivative of the cost \( C_{\text{poly}} \) is zero at the best fit. Show that the Hessian second derivative of the cost is

\[
H_{mn}^{\text{poly}} = \frac{\partial^2 C_{\text{poly}}}{\partial \theta_m \partial \theta_n} = \frac{1}{m + n + 1}.
\]

This Hessian is the Hilbert matrix, famous for being ill-conditioned (having a huge range of eigenvalues).\(^\text{11}\) Tiny eigenvalues of \( H^{\text{poly}} \) correspond to directions in polynomial space where the fit does not change.

(b) Calculate the eigenvalues of the \( 6 \times 6 \) Hessian for fifth-degree polynomial fits. Do they indeed span a large range? How big is the condition number (the ratio of the largest to the smallest eigenvalue)? Calculate the eigenvalues of larger Hilbert matrices. At what size do your eigenvalues seem contaminated by rounding errors?

Notice from Eqn N3.17 that the dependence of the polynomial fit on the monomial coefficients is independent of the function \( f(x) \) being fitted. We can thus vividly illustrate the sloppiness of polynomial fits by considering fits to the zero function \( f(x) \equiv 0 \). A polynomial given by an eigenvector of the Hilbert matrix with small eigenvalue must stay close to zero everywhere in the range \([0, 1]\). Let us check this.

(c) Calculate the eigenvector corresponding to the smallest eigenvalue of \( H^{\text{poly}} \), checking to make sure its norm is one (so the coefficients are of order one). Plot the corresponding polynomial in the range \([0, 1]\): does it stay small everywhere in the interval? Plot it in a larger range \([-1, 2]\) to contrast its behavior inside and outside the fit interval.

This turns out to be a fundamental property that is shared with many other multiparameter fitting problems. Many different terms are used to describe this property. The fits are called ill-conditioned: the parameters \( \theta_n \) are not well constrained by the data. The inverse problem is challenging: one cannot practically extract the parameters from the behavior of the model. Or, as our group describes it, the fit is sloppy: only a few directions in parameter space (eigenvectors corresponding to the largest eigenvalues) are constrained by the data, and there is a huge space of models (polynomials) varying along sloppy directions that all serve well in describing the data.

At root, the problem with polynomial fits is that all monomials \( x^n \) have similar shapes on \([0, 1]\): they all start flat near zero and bend upward. Thus they can be traded for one

\(^{11}\)For fits to discrete data, at points \((x_0, \ldots, x_{N-1})\), the Hessian \( H^{\text{poly}} = V^T V \), where \( V_{i\alpha} = x_i^\alpha \) is the Vandermonde matrix, also famous for being ill-conditioned.
another; the coefficient of \(x^4\) can be lowered without changing the fit if the coefficients of \(x^3\) and \(x^5\) are suitably adjusted to compensate. Indeed, if we change basis from the coefficients \(\theta_n\) of the monomials \(x^n\) to the coefficients \(\ell_n\) of the orthogonal (shifted Legendre) polynomials, the situation completely changes. The Legendre polynomials are designed to be different in shape (orthogonal), and hence cannot be traded for one another. Their coefficients \(\ell_n\) are thus well determined by the data, and indeed the Hessian for the cost \(C^{\text{poly}}\) in terms of this new basis is the identity matrix.

N3.12 **Conservative differential equations: Accuracy and fidelity.** (Ordinary differential equations)

In this exercise, we will solve for the motion of a particle of mass \(m = 1\) in the potential

\[
V(y) = \frac{1}{8} y^2 (-4A^2 + \log^2(y^2))\,.
\]

That is,

\[
\frac{d^2 y}{dt^2} = -\frac{dV}{dy}
\]

\[
= -\left(\frac{1}{4}\right) y (-4A^2 + 2 \log(y^2) + \log^2(y^2))\,.
\]

We will start the particle at \(y_0 = 1\), \(v_0 = \frac{dy}{dt}|_{0} = -A\), and choose \(A = 6\).

(a) Show that the solution to this differential equation is\(^\text{12}\)

\[
F(t) = \exp(-6 \sin(t))\,.
\]

Note that the potential energy \(V(y)\) is zero at the five points \(y = 0, y = \pm \exp(\pm A)\).

(b) Plot the potential energy for \(-3/2 \exp(\pm A) < y < 3/2 \exp(\pm A)\) (both zoomed in near \(y = 0\) and zoomed out). The correct trajectory should oscillate in the potential well with \(y > 0\), turning at two points whose energy is equal to the initial total energy. What is this initial total energy for our the particle? How much of an error in the energy would be needed, for the particle to pass through the origin when it returns? Compare this error to the maximum kinetic energy of the particle (as it passes the bottom of the well). Small energy errors in our integration routine can thus cause significant changes in the trajectory.

(c) (Black-box) Using a professionally written black-box differential equation solver of your choice, solve for \(y(t)\) between zero and \(4\pi\), at high precision. Plot your answer along with \(F(t)\) from part (a). (About half of the solvers will get the answer qualitatively wrong, as expected from part (b).) Expand your plot to examine the region \(0 < y < 1\); is energy conserved? Finally, read the documentation for your black box routine, and describe the combination of algorithms being used. You may wish to implement and compare more than one algorithm, if your black-box has that option.

\(^{12}\)I worked backward to do this. I set the kinetic energy to \(1/2 (dF/dt)^2\), set the potential energy to minus the kinetic energy, and then substituted \(y\) for \(t\) by solving \(y = F(t)\).
Choosing an error tolerance for your differential equation limits the error in each time step. If small errors in early time steps lead to important changes later, your solution may be quite different from the correct one. Chaotic motion, for example, can never be accurately simulated on a computer. All one can hope for is a faithful simulation—one where the motion is qualitatively similar to the real solution. Here we find an important discrepancy—the energy of the numerical solution is drifting upward or downward, where energy should be exactly conserved in the true solution. Here we get a dramatic change in the trajectory for a small quantitative error, but any drift in the energy is qualitatively incorrect.

The Leapfrog algorithm is a primitive looking method for solving for the motion of a particle in a potential. It calculates the next position from the previous two:

\[ y(t + h) = 2y(t) - y(t - h) + h^2 f[y(t)] \quad (N3.21) \]

where \( f[y] = -dV/dy \). Given an initial position \( y(0) \) and an initial velocity \( v(0) \), one can initialize and finalize

\[
\begin{align*}
  y(h) &= y(0) + h(v(0) + (h/2)f[y(0)]) \\
  v(T) &= (y(T) - y(T - h))/h + (h/2)f[y(T)]
\end{align*}
\quad (N3.22)
\]

Leapfrog is one of a variety of Verlet algorithms. In a more general context, this is called \textit{Stoermer’s rule}, and can be extrapolated to zero step-size \( h \) as in the Bulirsch–Stoer algorithms. A completely equivalent algorithm, with more storage but less roundoff error, is given by computing the velocity \( v \) at the midpoints of each time step:

\[
\begin{align*}
  v(h/2) &= v(0) + (h/2)f[y(0)] \\
  y(h) &= y(0) + hv(h/2) \\
  v(t + h/2) &= v(t - h/2) + hf[y(t)] \\
  y(t + h) &= y(t) + hv(t + h/2)
\end{align*}
\quad (N3.23)
\]

where we may reconstruct \( v(t) \) at integer time steps with \( v(t) = v(t-h/2)+(h/2)f[y(t)] \).

(d) (Leapfrog and symplectic methods) \textit{Show that} \( y(t + h) \) \textit{and} \( y(h) \) \textit{from eqns N3.21 and N3.22 converge to the true solutions as} \( h \to 0 \)—\textit{that is, the time-step error compared to the solution of} \( d^2y/dt^2 = f[y] \) \textit{vanishes faster than} \( h \). \textit{To what order in} \( h \) \textit{are they accurate after one time step?} \textsuperscript{13} \textit{Implement Leapfrog, and apply it to solving eqn N3.19 in the range} \( 0 < x < 4\pi \), \textit{starting with} \( h = 0.01 \). \textit{How does the accuracy compare to your more sophisticated integrator, for times less than} \( t = 2 \)? \textit{Zoom in to the range} \( 0 < y < 1 \), \textit{and compare with your packaged integrator and the true solution. Which has the more accurate period (say, by measuring the time to the first minimum in} \( y \)? \textit{Which has the smaller energy drift (say, by measuring the change in depth between subsequent minima)?}

\textsuperscript{13} \textit{Warning: for subtle reasons, the errors in Leapfrog apparently build up quadratically as one increases the number of time steps, so if your error estimate after one time step is} \( h^n \) \textit{the error after} \( N = T/h \) \textit{time steps can’t be assumed to be} \( Nh^n \sim h^{n-1} \), \textit{but is actually} \( N^2h^n \sim h^{N-2} \).
Fidelity is often far more important than accuracy in numerical simulations. Having an algorithm that has a small time-step error but gets the behavior qualitatively wrong is less useful than a cruder answer that is faithful to the physics. Leapfrog here is capturing the oscillating behavior far better than vastly more sophisticated algorithms, even though it gets the period wrong.

How does Leapfrog do so well? Systems like particle motion in a potential are Hamiltonian systems. They not only conserve energy, but they also have many other striking properties like conserving phase-space volume (Liouville’s theorem, the basis for statistical mechanics). Leapfrog, in disguise, is also a Hamiltonian system. (Eqns N3.23 can be viewed as a composition of two canonical transformations—one advancing the velocities at fixed positions, and one advancing the positions at fixed velocities.) Hence it exactly conserves an approximation to the energy—and thus doesn’t suffer from energy drift, satisfies Liouville’s theorem, etc. Leapfrog and the related Verlet algorithm are called *symplectic* because they conserve the *symplectic form* that mathematicians use to characterize Hamiltonian systems.

It is often vastly preferable to do an exact simulation (apart from rounding errors) of an approximate system, rather than an approximate analysis of an exact system. That way, one can know that the results will be physically sensible (and, if they are not, that the bug is in your model or implementation, and not a feature of the approximation).
Chapter 4

Quantum

These exercises were designed for a graduate quantum mechanics course at Cornell.

N4.1 Quantum notation. (Notation) ②

For each entry in the first column, match the corresponding entries in the second and third column.

\begin{itemize}
  \item \textbf{Schrödinger}
    \begin{align*}
      \psi(x) \\
      \psi^*(x) \\
      |\psi(x)|^2 \\
      \int \phi^*(x)\psi(x)dx \\
      |\int \phi^*(x)\psi(x)dx|^2 \\
      \int |\psi(x)|^2dx \\
      (\hbar/2m)(\psi^*\nabla\psi - \psi\nabla\psi^*)
    \end{align*}
  \item \textbf{Dirac}
    \begin{enumerate}
      \item \textbf{A.} \langle \psi|x\rangle p/2m\langle x|\psi\rangle - \langle x|\psi\rangle p/2m\langle x|\psi\rangle
      \item \textbf{B.} \textit{Ket} \langle \psi | \rangle \textit{in position basis, or} \langle x | \psi \rangle
      \item \textbf{C.} \textit{Bra} \langle x | \psi \rangle \textit{in position basis, or} \langle \psi | x \rangle
      \item \textbf{D.} \langle \psi | \rangle \langle x | \psi \rangle
      \item \textbf{E.} \textit{Braket} \langle \phi | \psi \rangle
      \item \textbf{F.} \langle \psi | \phi \rangle
      \item \textbf{G.} \langle \psi | \phi \rangle \langle \phi | \psi \rangle
    \end{enumerate}
\end{itemize}
N4.2 Light proton atomic size. (Dimensional analysis) ③

In this exercise, we examine a parallel world where the proton and neutron masses are equal to the electron mass, instead of \( \sim 2,000 \) times larger.

In solving the hydrogen atom in your undergraduate quantum course, you may have noted that in going from the 6-dimensional electron-proton system into the three-dimensional center-of-mass coordinates, the effective mass gets shifted to a reduced mass \( m_{\text{red}} = \mu = 1/(1/m_e + 1/m_{\text{nucleus}}) \), and is otherwise the hydrogen potential with a fixed (infinite-mass) nucleus. Let us assume that the atomic sizes and the excitation energies are determined solely by this mass shift.

What is the reduced mass for the hydrogen atom in the parallel world of light protons, compared to the electron mass? How much larger will the atom be? How much will the binding energy of the atom change? (You may approximate \( M_p \sim \infty \) when appropriate.)

(Units hint: \( [\hbar] = ML^2/T, [ke^2] = \text{Energy} \times L = ML^3/T^2 \), and \( [m_e] = M \). Here \( k = 1 \) in CGS units, and \( k = 1/(4\pi\epsilon_0) \) in SI units.)

N4.3 Light proton tunneling. (Dimensional analysis) ③

![Fig. N4.1 Atom tunneling. A hydrogen atom tunnels a distance \( a_0 \), breaking a bond of strength \( E_{\text{bind}} \) equal to its ionization energy.](image)

In this exercise, we continue to examine a parallel world where the proton and neutron masses are equal to the electron mass, instead of \( \sim 2,000 \) times larger.

With everything two thousand times lighter, will atomic tunneling become important? Let’s make a rough estimate of the tunneling suppression (given by the approximate WKB formula \( \exp(-\sqrt{2MVQ/\hbar}) \)).

Imagine an atom hopping between two positions, breaking and reforming a chemical bond in the process—an electronic energy barrier, and an electronic-scale distance. The
distance will be some fraction of a Bohr radius $a_0$ and the barrier energy will be some fraction of a Rydberg, but the the atomic mass would be some multiple of the proton mass.

*In our world, what would the suppression factor be for an hydrogen atom of mass $\sim M_p$ tunneling through a barrier of height $V$ of one Rydberg $= \hbar^2/(2m_ea_0^2)$, and width $Q$ equal to the Bohr radius $a_0$? How would this change in the parallel world where $M_p \rightarrow m_e$? (Simplify your answer as much as possible.)* (Use the real-world\(^1\) $a_0$ and Rydberg for the parallel world, not your answers from a previous exercise. Also please use the simple formula above: don’t do the integral. Your answer should involve only two of the fundamental constants.)

**N.4.4 Light proton superfluid. (Quantum) \(^3\)**

In this exercise, we yet again examine a parallel world where the proton and neutron masses are equal to the electron mass, instead of $\sim 2,000$ times larger.

![Fig. N4.2 Atoms with imaginary box of size equal to average space per atom](image)

Atoms Bose condense when the number density $n$ is high and the temperature $T$ is low. One can view this condensation point as where temperature and the confinement energy become competitive. Up to a constant of order one, the thermal energy at the transition temperature $k_B T_c$ equals the confinement energy needed to put a helium atom in a box with impenetrable walls of length the mean nearest-neighbor spacing $L = n^{-1/3}$.

*Calculate the ratio of the confinement energy $E_{\text{He,ours}}$ of Helium in our world and water $E_{\text{H}_2\text{O,light}}$ in the parallel world. Helium goes superfluid at $\sim 2^\circ K$ in our universe. From that, estimate the superfluid transition temperature $T_c$ for water in the parallel universe. Will it be superfluid at room temperature?* (Assume the box sizes for He in our world and water in the parallel world are the same.\(^2\) Room temperature is about $300^\circ K$. Helium 4, the isotope that goes superfluid, has two neutrons and two protons, with roughly equal masses. Oxygen 16 has eight protons, eight neutrons, and eight electrons.)

---

\(^1\)The reduced mass effects you found in the earlier exercise will be much less important for larger atoms and molecules, so we shall not include them here.

\(^2\)The number of molecules of water per unit volume is comparable to the number of atoms of liquid helium per unit volume in the real world, and the water molecule will stay about the same size in the parallel world.
N4.5 **Aharonov-Bohm Wire.** (Parallel transport) ③

What happens to the electronic states in a thin metal loop as a magnetic flux $\Phi_B$ is threaded through it? This was a big topic in the mid-1980’s, with experiments suggesting that the loops would develop a spontaneous current, that depended on the flux $\Phi_B/\Phi_0$, with $\Phi_0 = \hbar c/e$ the quantum of flux familiar from the Bohm-Aharonov effect. In particular, Nandini Trivedi worked on the question while she was a graduate student here:

Nandini Trivedi and Dana Browne, “Mesoscopic ring in a magnetic field: Reactive and dissipative response”, *Phys. Rev. B* 38, 9581-9593 (1988);
she’s now a faculty member at Ohio State.

Some of the experiments clearly indicated that the periodicity in the current went as $\Phi_0/2 = \hbar c/2e$—half the period demanded by Bohm and Aharonov from fundamental principles. (This is OK; having a greater period would cause one to wonder about fractional charges.) Others found (noisier) periods of $\Phi_0$. Can we do a free-particle-on-a-ring calculation to see if for some reason we get half the period too?

Consider a thin wire of radius $R$ along $x^2 + y^2 = R^2$. Let a solenoid containing magnetic flux $\Phi_B$, thin compared to $R$, lie along the $\hat{z}$ axis. Let $\phi$ be the angle around the circle with respect to the positive $x$-axis. (Don’t confuse the flux $\Phi_B$ with the angle $\phi$!) We’ll assume the wire confines the electron to lie along the circle, so we’re solving a one-dimensional Schrödinger’s equation along the coordinate $s = R\phi$ around the circle. Assume the electrons experience a random potential $V(s)$ along the circumference $C = 2\pi R$ of the wire.

(a) Choose a gauge for $\vec{A}$ so that it points along $\hat{\phi}$. What is the one-dimensional time-independent Schrödinger equation giving the eigenenergies for electrons on this ring? What is the boundary conditions for the electron wavefunction $\psi$ at $s = 0$ and $s = C$? (Hint: the wire is a circle; nothing fancy yet. I’m not asking you to solve the equation—only to write it down.)

Deriving the Bohm-Aharonov effect using Schrödinger’s equation is easiest done using a singular gauge transformation.

(b) Consider the gauge transformation $\Lambda(r, \phi) = -\phi \Phi_B/(2\pi)$. Show that $\vec{A}' = \vec{A} + \nabla \Lambda$ is zero along the wire for $0 < s < C$, so that we are left with a zero-field Schrödinger equation. What happens at the endpoint $C$? What is the new boundary condition for the electron wave function $\psi'$ after this gauge transformation? Does the effect vanish for $\Phi_B = n\Phi_0$ for integer $n$, as the Bohm-Aharonov effect says it should?

Realistically, the electrons in a large, room-temperature wire get scattered by phonons or electron-hole pairs (effectively, a quantum measurement of sorts) long before they propagate around the whole wire, so these effects were only seen experimentally when the wires were cold (to reduce phonons and electron-hole pairs) and mesoscopic (tiny, so the scattering length is comparable to or larger than the circumference).
Finally, let’s assume free electrons, so $V(s) = 0$. What’s more, to make things simpler, let’s imagine that there is only one electron in the wire.

(c) Ignoring the energy needed to confine the electrons into the thin wire, solve the one-dimensional Schrödinger equation to give the ground state of the electron as a function of $\Phi_B$. Plot the current in the wire as a function of $\Phi_B$. Is it periodic with period $\Phi_0$, or periodic with period $\Phi_0/2$?

In the end, it was determined that there were two classes of experiments. Those that measured many rings at once (measuring an average current, an easier experiment) got periodicity of $hc/2e$, while those that attempted the challenge of measuring one mesoscopic ring at a time find $hc/e$.

N4.6 Anyons. (Statistics) ³


In quantum mechanics, identical particles are truly indistinguishable (Fig. N4.3). This means that the wavefunction for these particles must return to itself, up to an overall phase, when the particles are permuted:

$$\Psi(r_1, r_2, \ldots) = \exp(i\chi)\Psi(r_2, r_1, \ldots).$$  \hspace{1cm} (N4.1)

where $\cdots$ represents potentially many other identical particles.

We can illustrate this with a peek at an advanced topic mixing quantum field theory and relativity. Here is a scattering event of a photon off an electron, viewed in two reference frames; time is vertical, a spatial coordinate is horizontal. On the left we see two “different” electrons, one which is created along with an anti-electron or positron $e^+$, and the other which later annihilates the positron. On the right we see the same event viewed in a different reference frame; here there is only one electron, which scatters two photons. (The electron is *virtual*, moving faster than light, between the collisions; this is allowed in intermediate states for quantum transitions.) The two electrons on the left are not only indistinguishable, they are the *same particle*! The antiparticle is also the electron, traveling backward in time.$^3$

³This idea is due to Feynman’s thesis advisor, John Archibald Wheeler. As Feynman quotes in his Nobel lecture, *I received a telephone call one day at the graduate college at Princeton from Professor Wheeler, in which he said, “Feynman, I know why all electrons have the same charge and the same mass.” “Why?” “Because, they are all the same electron!” And, then he explained on the telephone, “suppose that the world lines which we were ordinarily considering before in time and space - instead of only going up in time were a tremendous knot, and then, when we cut through the knot, by the plane corresponding to a fixed time, we would see many, many world lines and that would represent many electrons, except for one thing. If in one section this is an ordinary electron world line, in the section in which it reversed itself and is coming back from the future we have the wrong sign to the proper time - to the proper four velocities - and that’s equivalent to changing the sign of the charge, and, therefore, that part of a path would act like a positron.”*
Fig. N4.3 Feynman diagram: identical particles.

In three dimensions, $\chi$ must be either zero or $\pi$, corresponding to bosons and fermions. In two dimensions, however, $\chi$ can be anything: *anyons* are possible! Let’s see how this is possible.

In a two-dimensional system, consider changing from coordinates $r_1, r_2$ to the center-of-mass vector $R = (r_1 + r_2)/2$, the distance between the particles $r = |r_2 - r_1|$, and the angle $\phi$ of the vector between the particles with respect to the $\hat{x}$ axis. Now consider permuting the two particles counter-clockwise around one another, by increasing $\phi$ at fixed $r$. When $\phi = 180^\circ \equiv \pi$, the particles have exchanged positions, leading to a boundary condition on the wavefunction

$$\Psi(R, r, \phi, \cdots) = \exp(i\chi)\Psi(R, r, \phi + \pi, \cdots). \quad (N4.2)$$

Permuting them counter-clockwise (backward along the same path) must then\(^4\) give $\Psi(R, r, \phi, \cdots) = \exp(-i\chi)\Psi(R, r, \phi - \pi, \cdots)$. This in general makes for a many-valued wavefunction (similar to Riemann sheets for complex analytic functions).

Why can’t we get a general $\chi$ in three dimensions?

(a) *Show, in three dimensions, that $\exp(i\chi) = \pm 1$, by arguing that a counter-clockwise rotation and a clockwise rotation must give the same phase.* (Hint: The phase change between $\phi$ and $\phi + \pi$ cannot change as we wiggle the path taken to swap the particles, unless the particles hit one another during the path. Try rotating the counter-clockwise path into the third dimension: can you smoothly change it to clockwise? What does that imply about $\exp(i\chi)$?)

Fig. N4.4 Braiding of paths in two dimensions. In two dimensions, one can distinguish swapping clockwise from counter-clockwise. Particle statistics are determined by representations of the *Braid group*, rather than the permutation group.

Figure N4.4 illustrates how in two dimensions rotations by $\pi$ and $-\pi$ are distinguishable; the trajectories form *braids* that wrap around one another in different ways. You can’t change from a counter-clockwise braid to a clockwise braid without the braids crossing (and hence the particles colliding).

\(^4\)The phase of the wave-function doesn’t have to be the same for the swapped particles, but the gradient of the phase of the wavefunction is a physical quantity, so it must be minus for the counter-clockwise path what it was for the clockwise path.
An angular boundary condition multiplying by a phase should seem familiar: it is quite similar to that of the Bohm-Aharonov effect we studied in exercise 2.4. Indeed, we can implement fractional statistics by producing composite particles, by threading a magnetic flux tube of strength $\Phi$ through the center of each 2D boson, pointing out of the plane.

(b) *Remind yourself of the Bohm-Aharonov phase incurred by a particle of charge $e$ encircling counter-clockwise a tube of magnetic flux $\Phi$. If a composite particle of charge $e$ and flux $\Phi$ encircles another identical composite particle, what will the net Bohm-Aharonov phase be?* (Hint: You can view the moving particle as being in a fixed magnetic field of all the other particles. The moving particle doesn’t feel its own flux.)

(c) *Argue that the phase change $\exp(i\chi)$ upon swapping two particles is exactly half that found when one particle encircles the other. How much flux is needed to turn a boson into an anyon with phase $\exp(i\chi)$?* (Hint: The phase change can’t depend upon the precise path, so long as it braids the same way. It’s homotopically invariant, see Chapter 9 of “Entropy, Order Parameters, and Complexity”.)

Anyons are important in the quantum Hall effect. What is the quantum Hall effect? At low temperatures, a two-dimensional electron gas in a perpendicular magnetic field exhibits a Hall conductance that is quantized, when the filling fraction $\nu$ (electrons per unit flux in units of $\Phi_0$) passes near integer and rational values.

Approximate the quantum Hall system as a bunch of composite particles made up of electrons bound to flux tubes of strength $\Phi_0/\nu$. As a perturbation, we can imagine later relaxin the binding and allow the field to spread uniformly.\(^5\)

(d) **Composite bosons and the integer quantum Hall effect.** At filling fraction $\nu = 1$ (the integer quantum Hall state), what are the effective statistics of the composite particle? Does it make sense that the (ordinary) resistance in the quantum Hall state goes to zero?

- The excitations in the fractional quantum Hall effect are anyons with fractional charge. (The $\nu = 1/3$ state has excitations of charge $e/3$, like quarks, and their wavefunctions gain a phase $\exp(i\pi/3)$ when excitations are swapped.)
- It is conjectured that, at some filling fractions, the quasiparticles in the fractional quantum Hall effect have nonabelian statistics, which could become useful for quantum computation.
- The composite particle picture is a central tool both conceptually and in calculations for this field.

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\(^5\)This is not nearly as crazy as modeling metals and semiconductors as noninteracting electrons, and adding the electron interactions later. We do that all the time—electrons and holes in solid-state physics, $1s$, $2s$, $2p$ electrons in multi-electron atoms, all have obvious meanings only if we ignore the interactions. Both the composite particles and the noninteracting electron model are examples of how we use adiabatic continuity—you find a simple model you can solve, that can be related to the true model by turning on an interaction.
Consider the following cooperative game played by Alice and Bob: Alice receives a bit $x$ and Bob receives a bit $y$, with both bits uniformly random and independent. The players win if Alice outputs a bit $a$ and Bob outputs a bit $b$, such that $(a + b = xy \mod 2)$. They can agree on a strategy in advance of receiving $x$ and $y$, but no subsequent communication between them is allowed.

(a) **Give a deterministic strategy by which Alice and Bob can win this game with $3/4$ probability.**

(b) **Show that no deterministic strategy lets them win with more than $3/4$ probability.** (Note that Alice has four possible deterministic strategies $[0, 1, x, \sim x]$, and Bob has four $[0, 1, y, \sim y]$, so there’s a total of 16 possible joint deterministic strategies.)

(c) **Show that no probabilistic strategy lets them win with more than $3/4$ probability.** (In a probabilistic strategy, Alice plays her possible strategies with some fixed probabilities $p_0, p_1, p_x, p_{\sim x}$, and similarly Bob plays his with probabilities $q_0, q_1, q_y, q_{\sim y}$.)

The upper bound of $\leq 75\%$ of the time that Alice and Bob can win this game provides, in modern terms, an instance of the Bell inequality, where their prior cooperation encompasses the use of any local hidden variable.

Let’s see how they can beat this bound of $3/4$, by measuring respective halves of an entangled state, thus quantum mechanically violating the Bell inequality.

Suppose Alice and Bob share the entangled state $\frac{1}{\sqrt{2}}(|\uparrow_\ell\rangle|\uparrow_r\rangle + |\downarrow_\ell\rangle|\downarrow_r\rangle)$, with Alice holding the left Qbit and Bob holding the right Qbit. Suppose they use the following strategy: if $x = 1$, Alice applies the unitary matrix $R_{\pi/6} = \begin{pmatrix} \cos \frac{\pi}{6} & -\sin \frac{\pi}{6} \\ \sin \frac{\pi}{6} & \cos \frac{\pi}{6} \end{pmatrix}$ to her Qbit, otherwise doesn’t, then measures in the standard basis and outputs the result as $a$. If $y = 1$, Bob applies the unitary matrix $R_{-\pi/6} = \begin{pmatrix} \cos \frac{\pi}{6} & \sin \frac{\pi}{6} \\ -\sin \frac{\pi}{6} & \cos \frac{\pi}{6} \end{pmatrix}$ to his Qbit, otherwise doesn’t, then measures in the standard basis and outputs the result as $b$. (Note that if the Qbits were encoded in photon polarization states, this would be equivalent to Alice and Bob rotating measurement devices by $\pi/6$ in inverse directions before measuring.)

(d) **Using this strategy:** (i) **Show that if $x = y = 0$, then Alice and Bob win the game with probability 1.**

(ii) **Show that if $x = 1$ and $y = 0$ (or vice versa), then Alice and Bob win with probability $3/4$.**

---

6 This exercise was developed by Paul Ginsparg, based on an example by Bell ‘64 with simplifications by Clauser, Horne, Shimony, & Holt (’69).

7 There’s another version for GHZ state, where three people have to get $a+b+c \mod 2 = x$ or $y$ or $z$. Again one can achieve only $75\%$ success classically, but they can win every time sharing the right quantum state.
(iii) Show that if \( x = y = 1 \), then Alice and Bob win with probability \( 3/4 \).

(iv) Combining parts (i)-(iii), conclude that Alice and Bob win with greater overall probability than would be possible in a classical universe.

This proves an instance of the CHSH/Bell Inequality, establishing that “spooky action at a distance” cannot be removed from quantum mechanics. Alice and Bob’s ability to win the above game more than \( 3/4 \) of the time using quantum entanglement was experimentally confirmed in the 1980s (A. Aspect et al.).\(^8\)

(e) (Bonus) Consider a slightly different strategy, in which before measuring her half of the entangled pair Alice does nothing or applies \( R_{\pi/4} \), according to whether \( x \) is 0 or 1, and Bob applies \( R_{\pi/8} \) or \( R_{-\pi/8} \), according to whether \( y \) is 0 or 1. Show that this strategy does even better than the one analyzed in a–c, with an overall probability of winning equal to \( \cos^2 \pi/8 = (1 + \sqrt{1/2})/2 \approx 0.854 \).

(Extra bonus) Show this latter strategy is optimal within the general class of strategies in which before measuring Alice applies \( R_{\alpha_0} \) or \( R_{\alpha_1} \), according to whether \( x \) is 0 or 1, and Bob applies \( R_{\beta_0} \) or \( R_{\beta_1} \), according to whether \( y \) is 0 or 1.

This will demonstrate that no local hidden variable theory can reproduce all predictions of quantum mechanics for entangled states of two particles.

N4.8 Parallel Transport, Frustration, and the Blue Phase. (Liquid crystals) \(^3\)


“Frustration and Curvature: the Orange Peel Carpet”, http://www.lassp.cornell.edu/sethna/FrustrationCurvature/


(Optional: for those wanting a challenge.) Both the Aharonov Bohm effect and Berry’s phase (later) are generalizations of the idea of parallel transport. Parallel transport, from differential geometry, tells one how to drag tangent vectors around on smooth surfaces. Just as we discovered that dragging the phase of a wavefunction around a closed loop in space gave it a net rotation due to the magnetic field (Aharonov-Bohm),

\(^8\)Ordinarily, an illustration of these inequalities would appear in the physics literature not as a game but as a hypothetical experiment. The game formulation is more natural for computer scientists, who like to think about different parties optimizing their performance in various abstract settings. As mentioned, for physicists the notion of a classical strategy is the notion of a hidden variable theory, and the quantum strategy involves setting up an experiment whose statistical results could not be predicted by a hidden variable theory.
the phase can rotate also when the Hamiltonian is changed around a closed curve in Hamiltonian space (Berry’s phase). Here we discuss how vectors rotate as one drags them around closed loops, leading us to the curvature tensor.

(a) Parallel transport on the sphere. Imagine you’re in St. Louis (longitude 90° W, latitude ~ 40° N), pointing north. You walk around a triangle, first to the North Pole, then take a right-angle turn, walk down through Greenwich, England and Madrid, Spain (longitude ~ 0° W) down to the equator, turn right, walk back to longitude 90° W along the equator, turn north, and walk back to St. Louis. All during the walk, you keep pointing your finger in the same direction as far as feasible (i.e., straight ahead on the first leg, off to the left on the second, and so on). What angle does the vector formed by your final pointing finger make with respect to your original finger? If you turned west in Madrid and walked along that latitude (~ 40° N) to St. Louis (yes, it is just as far north), would the angle be the same? (Hint: what about transport around a tiny quarter circle centered at the north pole?)

Lightning Intro to Differential Geometry. Parallel transport of vectors on surfaces is described using a covariant derivative \((D_i v)^j = \partial_i v^j + \Gamma^j_{ik} v^k\) involving the Christofel symbol \(\Gamma^\mu_{iv}\). The amount a vector \(v^\nu\) changes when taken around a tiny loop \(\Delta s \Delta t (-\Delta s)(-\Delta t)\) is given by the Riemannian curvature tensor and the area of the loop

\[
v^n - v^i = R^{i}_{jk\ell} v^j k \Delta s \Delta t. \tag{N4.3}
\]

The algebra gets messy on the sphere, though (spherical coordinates are pretty ugly).

Instead, we’ll work with a twisty kind of parallel transport, that my advisor and I figured out describe how molecules of the Blue Phase like to locally align. These long, thin molecules locally line up with their axes in a direction \(\mathbf{n}(\mathbf{r})\), and are happiest when that axis twists so as to make zero the covariant derivative

\[
(D_i \mathbf{n})^j = \partial_i n^j - q \epsilon_{ijk} n_k, \tag{N4.4}
\]

where \(\epsilon_{ijk}\) is the totally antisymmetric tensor\(^9\) with three indices (Fig. N4.5 and N4.6). Since the Blue Phases are in flat space, you don’t need to worry about the difference between upper and lower indices.

\(^9\)Hence \(\Gamma^\mu_{ik} = -q \epsilon_{ijk}\). Warning: this connection has torsion: it is not symmetric under interchange of the two bottom indices. This is what allows it to have a nonzero curvature even when the liquid crystal lives in flat space. But it will make some of the traditional differential geometry formulas incorrect, which usually presume zero torsion.
**Fig. N4.5 Blue Phase Molecules.** long thin molecules aligned along an axis $\mathbf{n}(r)$, like to sit at a slight angle with respect to their neighbors. To align the threads and grooves on the touching surface between the molecules demands a slight twist. This happens because, like threaded bolts or screws, these molecules are *chiral*.

**Fig. N4.6 Twisted Parallel Transport.** The natural parallel transport, $D_i n_j = \partial_i n_j - q\epsilon_{ijk} n_k$, twists perpendicular to the long axis, but doesn’t twist when moving along the axis.

(b) What direction will a molecule with $\mathbf{n} = \hat{x}$ like to twist as one moves along the $x$-direction? The $y$-direction? The $z$-direction? Is the local low energy structure more like that of Fig. N4.7(a) (the low-temperature state of this system), or that of Fig. N4.7(b)?
So, what’s the curvature tensor (eqn N4.3) for the blue phase? I figured out at the time that it came out to be

$$R_{ijk\ell} = q^2 (\delta_{i\ell} \delta_{kj} - \delta_{ik} \delta_{\ell j}).$$ \hspace{1cm} (N4.5)

This curvature means that the blue phases are frustrated; you can’t fill space everywhere with material that’s “happy”, with $Dn = 0$.

(c) If $n$ starts at $\hat{x}$, and we transport along the tiny loop $\{\Delta x, \Delta y, -\Delta x, -\Delta y\}$, calculate the vector $n'$ as it returns to the origin using the curvature tensor (eqns N4.3 and N4.5). According to eqn N4.3. Is the shift in $n$ along the same direction we observe for the larger loop in Fig. N4.8?
**Fig. N4.8 Parallel transport frustration.** We can check your answer to part (c) qualitatively by thinking about a larger loop (Fig. N4.8). Consider starting from two places separated $\Delta x = d$ apart along the axis of a double-twisting region (the center of the tube in Fig. N4.7(b)). If we move a distance $\Delta y = \pi/(2q)$ radially outward, the orientation of both will now point along $\hat{z}$. Transporting them each inward by $d/2$ then will cause a further twist.

We can use the curvature tensor of the Blue Phase to calculate a scalar curvature $R_{ijij} = -6q^2$. Thus the blue phases are negatively curved, even though they live in flat space. We looked to see what would happen if we put the blue phases into a positively curved space. Picking the sphere in four dimensions with the correct radius, we could make the curvature (and the frustration) go away, and find the ideal template for the blue phases. We think of the real blue phase as pieces of this ideal template, cut, flattened, and sewn together to fill space, like an orange-peel carpet (Fig. N4.9).
N4.9 **Crystal field theory: d-orbitals.** (Group reps) ③

The vector space of functions $f(x, y, z)$ on the unit sphere transforms into itself under rotations $f(x) \to_R f(R^{-1}x)$. These transformations are linear $(af(x) + g(x) \to_R af(R^{-1}x) + g(R^{-1}x))$, and obey the group composition rule, and thus form a representation of the rotation group.

(a) Argue that the homogeneous polynomials of degree $\ell$,

$$f(x, y, z) = \sum_{m=0}^{\ell} \sum_{n=0}^{\ell-m} f_{\ell mn} x^m y^n z^{\ell-m-n}$$

(N4.6)

form a subspace that is invariant under rotations.

Thus the irreducible representations are contained in these invariant subspaces. Sakurai indeed mentions in his section 3.11 on tensor operators that $Y_{1^0} = \sqrt{3/\pi z/r}$ and $Y_{1^\pm1} = \sqrt{3/2\pi(x \pm iy)/r}$, and also gives a formula for $Y_{2^\pm2}$; since $r = 1$ on our unit sphere these are homogeneous polynomials of degree one.

(b) Look up the $\ell = 2$ spherical harmonics (e.g. in Sakurai’s appendix B) and write them as quadratic polynomials in $x$, $y$, and $z$.

The $\ell = 2$ spherical harmonics are the angular parts of the wavefunctions for electrons in $d$ orbitals (e.g. of transition metal atoms). ⑩ Electrons in $d$-orbitals are much more

---

⑩Here we use the common independent-electron language, where the complex many-body wavefunction of an atom, molecule, or solid is viewed as filling single-electron states, even though the electron-electron repulsion is almost as strong as the electron-nuclear attraction. This idea can be dignified in three rather different ways. First, one can view each electron as feeling an effective potential given by the nucleus plus the average density of electrons. This leads to mean-field Hartree-Fock theory. Second, one can show that the ground state energy can be written as an unknown functional of the electron density (the Hohenberg-Kohn theorem, and then calculate the kinetic energy terms as an effective single-body Schrödinger equation in the resulting effective potential due to the net electron density (the Kohn-Sham equations). Third, one can start with independent electrons (or Hartree-Fock electrons) and slowly “turn on” the electron-electron repulsion. The independent-electron excited eigenstates develop lifetimes and become resonances. For atoms these lifetimes represent Auger decay rates. For crystals these resonances are called quasiparticles and the theory is called Landau Fermi liquid theory. Landau Fermi-liquid theory is usually derived using Green’s functions and Feynman diagrams, but it has recently been re-cast as a renormalization-group flow.
tightly contained near the nucleus than $p$ and $s$ orbitals. In molecules and solids, the $s$ and $p$ orbitals usually hybridize (superimpose) into chemical bonds and broad electron bands, where the original orbitals are strongly distorted. In contrast, $d$-electrons rarely participate in chemical bonds, and their electron bands are narrow—almost undistorted orbitals with small hopping rates. The energy levels of the five $d$-orbitals are, however, shifted from one another by their environments. (For crystals, these shifts are called crystal field splittings.)

We can use group representation theory to understand how the $d$-orbitals are affected by their molecular or crystalline environment.

First, we need to calculate the character $\chi(R) = \chi(\hat{n}, \phi)$ of the $\ell = 2$ representation. Remember that the character is the trace of the (here $5 \times 5$) matrix corresponding to the rotation. Remember that this trace depends only on the conjugacy class of $R$—that is, if $S$ is some other group element then $\chi(S^{-1}RS) = \chi(R)$. Remember that any two rotations by the same angle $\phi$ are conjugate to one another.\(^1\)

In class, we found that the character of the $\ell = 1$ representation by using the Cartesian $x, y, z$ basis, where $R_{\hat{z}}(\phi) = \begin{pmatrix} \cos(\phi) & -\sin(\phi) & 0 \\ \sin(\phi) & \cos(\phi) & 0 \\ 0 & 0 & 1 \end{pmatrix}$. Hence $\chi^{(1)}(\phi) = 1 + 2\cos(\phi)$. We can do this same calculation in the $m_z$ basis of the spherical harmonics, where $Y_{01}$ is unchanged under rotations and $Y_{1\pm1} \rightarrow \exp(\pm i\phi)Y_{1\pm1}$. Here $R_{\hat{z}}(\phi) = \begin{pmatrix} \exp(i\phi) & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & \exp(-i\phi) \end{pmatrix}$, and again $\chi^{(1)}(\phi) = 1 + 2\cos(\phi)$.

(c) Calculate $\chi^{(2)}(\phi)$. Give the characters for rotations $C_n$ by $2\pi/n$, for $n = 1, 2, 3,$ and $4$ (the important rotations for crystalline symmetry groups.)

The most common symmetry groups for $d$-electron atoms in crystals is $O$, the octahedral group (the symmetry group of a cube). Look up the character tables for the irreducible representations of this finite group. (To simplify the calculation, we’ll assume that inversion symmetry is broken; otherwise we should use $O_h$, which has twice the number of group elements.)

(d) Use the orthogonality relations of the characters of irreducible representations for $O$, decompose the $\ell = 2$ representation above into irreducible representations of the octahedral group. How will the energies of the single-particle $d$-orbitals of a transition metal atom split in an octahedral environment? What will the degeneracies and symmetries ($A_1$, $A_2$, $E$, …) be of the different levels? (Hint: If you are doing it right, the dot product of the characters should equal a multiple of the number of octahedral group elements o($O$) = 24, and the dimensions of the sub-representations should add up to five.)

The five $d$-orbitals are often denoted $d_{xy}$, $d_{xz}$, $d_{yz}$, $d_{z^2}$ and $d_{x^2-y^2}$. This is a bit of a cheat; really $d_z$ should be written $d_{2z^2-x^2-y^2}$ or something like that.

(e) Figure out which of these orbitals are in each of the two representations you found

\(^1\)If the two rotations have axes $\hat{n}$ and $\hat{n}'$, choose $S$ to rotate $\hat{n}'$ into $\hat{n}$.
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in part (d). (Hint: Check how these five orbitals transform under the octahedral symmetries that permute $x$, $y$, and $z$ among themselves.)

N4.10 Entangled Spins. (Spins) 3

In class, we studied the entanglement of the singlet spin state $|S\rangle = (1/\sqrt{2})(|\uparrow\rangle_L|\downarrow\rangle_R - |\downarrow\rangle_L|\uparrow\rangle_R)$ of electrons of a diatomic molecule as the atoms $L$ and $R$ are separated;\(^{12}\) the spins on the two atoms are in opposite directions, but the system is in a superposition of the two choices. Here we discuss another such superposition, but with a different relative phase for the two choices:

$$|\chi\rangle = (1/\sqrt{2})(|\uparrow\rangle_L|\downarrow\rangle_R + |\downarrow\rangle_L|\uparrow\rangle_R)$$  \hspace{1cm} (N4.7)

You should know from angular momentum addition rules that the space of product wavefunctions of two spin $\frac{1}{2}$ states can be decomposed into a spin 1 and a spin 0 piece: $\frac{1}{2} \otimes \frac{1}{2} = 1 \oplus 0$. So there are two orthogonal eigenstates of $S_z$ with eigenvalue zero: one of total spin zero and one of total spin one.

(a) Total spin. Which state is which? (If you don’t know from previous work, calculate!) Why do we call $|S\rangle$ a singlet?

Now, is the spin wavefunction compatible with what we know about electron wavefunctions?

(b) Symmetry. When the two spins are exchanged, how does $|\chi\rangle$ change? If the total wavefunction $\Psi(x_L, s_L, x_R, s_R)$ is a product of this spin wavefunction $\chi(s_L, s_R)$ and and a two-particle spatial wavefunction $\psi(x_L, x_R)$, what symmetry must $\psi$ have under interchange of electrons?

We noted in class that two other spin-1 product states, $|\uparrow\rangle_L|\uparrow\rangle_R$ and $|\downarrow\rangle_L|\downarrow\rangle_R$ do not form entangled states when $L$ and $R$ separate. Is $|\chi\rangle$ like these spin-1 states, or is it entangled like $|S\rangle$ is?

(c) Entanglement. Give the Schmidt decomposition of $|\chi\rangle$. What are the singular values? What is the entanglement entropy? (Hint: The steps should be very familiar from class.)

(d) Singular value decomposition (SVD). Let $M$ be the matrix which gives $|\chi\rangle$ as a product of left and right spin states:

$$|\chi\rangle = (|\uparrow\rangle_L \mid\downarrow\rangle_R) M \left( \begin{array}{c} |\uparrow\rangle_R \\ |\downarrow\rangle_R \end{array} \right).$$  \hspace{1cm} (N4.8)

What is $M$? Give an explicit singular value decomposition\(^{13}\) $M = U\Sigma V^T$ of the matrix $M$. Explain how the SVD gives us the Schmidt decomposition of part (c).

---

\(^{12}\)We assumed that, when separated, one electron is localized basically on each of the two atoms, and the spin kets are labeled based on the primary locale of the corresponding spatial wavefunction for that electron.

\(^{13}\)Remember that the SVD guarantees that $U$ and $V$ have orthonormal columns, and $\Sigma$ is a diagonal matrix whose diagonal elements $\sigma_i$ are all positive and decreasing (so $\sigma_i \geq \sigma_{i+1} \geq 0$). There is some flexibility in the singular vectors (i.e., matched pairs can both have their signs changed), but the singular values are unique and hence a property of the matrix.
N4.11 **Square well ground state.**\(^1\) (Quantum numeric) ②

Consider a particle of mass \(M\) in one dimension, confined in a potential that vanishes for \(-a \leq x \leq a\) and becomes infinite at \(x = \pm a\), so the wavefunction must vanish at \(x = \pm a\).

Find the normalized ground state \(\psi_0(x)\). Suppose that the particle is placed in a state \(\psi\) with a wave function proportional to \(a^2 - x^2\), which vanished properly at the edges. If the energy of the particle is measured, what is the probability that the particle will be found in the state of lowest energy? Use a computational method (symbolic or numerical) to avoid tedious evaluation of integrals. Evaluate your answer numerically (it should not depend on \(a\)). Also, plot the ground state and the trial state, using \(a = 3\); they should look rather similar. This exercise may take much longer than it seems, since you’ll likely spend some time setting up your computational environment—installing software, getting plots to work, and so forth. The Hints files should help once the software is set up.

N4.12 **Evolving Schrödinger: Free particles and uncertainty.** (Computation) ③

Several of our computational exercises will involve solving the time-dependent Schrödinger equation for one-dimensional quantum systems. In this first of these exercises, we shall evolve the free particle Hamiltonian

\[
\frac{i\hbar}{\partial t} \psi(t) = -\frac{\hbar^2}{2m} \frac{\partial^2 \psi}{\partial x^2} = H_{\text{kin}} \psi
\]

\[
\psi(x, t) = U(t) \psi(0) = e^{-i H_{\text{kin}} t / \hbar} \psi(x, 0)
\]  (N4.9)

In a later exercise, we shall solve for the behavior of a hydrogenic harmonic oscillator with angular frequency \(\omega = 10^{12}\) radians/sec. We’ll use the ground state of this harmonic oscillator as our initial condition for the free-particle evolution:

\[
\psi_0(x) = \left( \frac{m\omega}{\pi \hbar} \right)^{1/4} e^{-m\omega x^2 / 2\hbar} = \left( \frac{1}{2\pi a_0^2} \right)^{1/4} e^{-x^2 / 4a_0^2}.
\]  (N4.10)

Here \(a_0 = \sqrt{\hbar/2m\omega}\) is the RMS width of the Gaussian probability distribution \(|\psi(x)|^2\).

With no potential energy, we can solve for the motion of a free particle, \(\widetilde{\psi}(k, t) = \mathcal{U}_{\text{kin}}(t) \psi(k, t = 0)\) using Fourier transforms.

\[
\widetilde{\psi}(k, t) = e^{-i(k^2/2m)t} \widetilde{\psi}(k, t = 0)
\]  (N4.11)

\[
\widetilde{\psi}(x, t) = \text{IFFT}[e^{-i(k^2/2m)t} \text{FFT}[\widetilde{\psi}(x, t = 0)]](\text{N4.12})
\]

where FFT takes a Fourier transform of the wavepacket, and IFFT takes an inverse Fourier transform. (FFT stands for Fast Fourier Transform.)

---

\(^1\)Adapted from Weinberg, *Lectures on Quantum Mechanics*, ex. 1.1. Hints files available on the course website for Mathematica, Python, and Octave/Matlab.
Let’s numerically solve for the evolution of \( \psi_0(x) \) from eqn N4.10. We’ll evaluate it at a discrete set of \( N_p = 200 \) points spanning a distance \( L = 80a_0 \). With \( dx = L/N_p \), the points will be at \(-L/2, L/2 + dx, \ldots, L/2 - dx\). In CGS units, the hydrogen mass is about the proton mass \( 1.672610^{-24} \) gm, \( \hbar = 1.0545710^{-27} \) erg sec, and we decided \( \omega = 10^{12} \) radians/sec.

(a) Define \( \psi_0 \) on this grid. Plot your \( |\psi_0|^2 \), and check that it roughly has width \( a_0 \).

The Fast Fourier Transform of \( \psi \) returns \( \tilde{\psi}(k) \) evaluated at points \( k = (0, dk, 2dk, \ldots, -2dk, -dk) \), \( (N4.13) \) with \( dk = 2\pi/L \). (These correspond to the plane waves with period \( L \); FFTs assume periodic boundary conditions.) The maximum value \( k_{\text{max}} = \pi/dx \) happens in the center of the FFT. To do our time evolution, we need to define an array \( k^2 \) evaluated at these points, which should rise quadratically, come to a cusp, and then fall back to zero.

(b) Define \( k^2 \) on this grid. Plot it.

Now we can evolve \( \psi(x, t) = U_{\text{kin}}(t)\psi_0(x) \).

(c) Create a routine that calculates \( \psi(x, t) \) using eqn N4.12. Plot the real and imaginary parts of \( \psi(x, t) \) at \( t = P/4, P, \) and \( 2P \) on separate plots, where \( P = 2\pi/\omega \) is the period of the harmonic oscillator in the later exercise. What happens to the width of the wave packet?

The Heisenberg uncertainty principle tells us that \( \Delta x \Delta p \geq \hbar/2 \). The Gaussian wavepacket we use is the form that minimizes this inequality, so we expect the packet width to grow like \( vt \) with \( v \) given by the momentum uncertainty \( v \sim \Delta p/m \).

(d) Give the formula for \( v \) for our packet. Calculate \( \Delta x = \sqrt{\langle x^2 \rangle} \) as a function of time, for points \( 0 \leq t \leq 2P \). Plot \( \Delta x \) versus time and \( vt \) versus time on the same plot. Why do they not agree at short times?

N4.13 Rotating Fermions. (Group theory) 

In this exercise, we’ll explore the geometry of the space of rotations.

Spin 1/2 fermions transform upon rotations under \( SU(2) \), the unitary \( 2 \times 2 \) matrices with determinant one. Vectors transform under \( SO(3) \), the ordinary \( 3 \times 3 \) rotation matrices you know of.

Sakurai argues that a general \( SU(2) \) matrix \( U = ( \begin{array}{cc} a & b \\ -b^* & a^* \end{array} ) \) with \( |a|^2 + |b|^2 = 1 \). Viewing \( \{\text{Re}(a), \text{Im}(a), \text{Re}(b), \text{Im}(b)\} \) as a vector in four dimensions, \( SU(2) \) then geometrically is the unit sphere \( S^3 \) in \( \mathbb{R}^4 \).

Remember that for every matrix \( R \) in \( SO(3) \), there are two unitary matrices \( U \) and \(-U \) corresponding to the same physical rotation. The matrix \(-U \) has coordinates \((-a, -b)\)—it is the antipodal point on \( S^3 \), exactly on the opposite side of the sphere. So \( SO(3) \) is geometrically the unit sphere with antipodal points identified. This is called (for obscure reasons) the projective plane, \( \mathbb{R}P^3 \).
Feynman’s plate (in Feynman’s plate trick) as it rotates 360° travels in rotation space from one orientation to its antipode. While I'm not sure anyone has figured out whether arms, shoulders, and elbows duplicate the properties of fermions under rotations, the plate motion illustrates the possibility of a minus sign.

But we can calculate this trajectory rather neatly by mapping the rotations not to the unit sphere, but to the space $\mathbb{R}^3$ of three-dimensional vectors. (Just as the 2-sphere $S^2$ can be projected onto the plane, with the north pole going to infinity, so can the 3-sphere $S^3$ be projected onto $\mathbb{R}^3$.) Remember the axis-angle variables, where a rotation of angle $\phi$ about an axis $\hat{n}$ is given by

$$\exp(-iS \cdot \hat{n}\phi/\hbar) = \exp(-i\sigma \cdot \hat{n}\phi/2) = \exp(-iJ \cdot \hat{n}\phi/\hbar) \quad (N4.14)$$

where the middle formula works for $SU(2)$ (where $S = \hbar\sigma/2$, because the particles have spin 1/2) and the last formula is appropriate for $SO(3)$.

We figure $\mathbf{n}$ will be the direction of the vector in $\mathbb{R}^3$ corresponding to the rotation, but how will the length depend on $\phi$? Since all rotations by 360° are the same, it makes sense to make the length of the vector go to infinity as $\phi \to 360°$. We thus define the Modified Rodrigues coordinates for a rotation to be the vector $\mathbf{p} = \hat{n}\tan(\phi/4)$.

(a) Fermions, when rotated by 360°, develop a phase change of $\exp(i\pi) = -1$ (as discussed in Sakurai & Napolitano p. 165, and as we illustrated with Feynman’s plate trick). Give the trajectory of the modified Rodrigues coordinate for the fermion’s rotation as the plate is rotated 720° about the axis $\hat{n} = \hat{z}$. (We want the continuous trajectory on the sphere $S^3$, perhaps which passes through the point at $\infty$. Hint: The trajectory is already defined by the modified Rodrigues coordinate: just describe it.)

(b) For a general Rodrigues point $\mathbf{p}$ parameterizing a rotation in $SO(3)$, what antipodal point $\mathbf{p}'$ corresponds to the same rotation? (Hint: A rotation by $\phi$ and a rotation by $\phi + 2\pi$ should be identified.)

### N4.14 Lithium ground state symmetry. (Quantum)

A simple model for heavier atoms, that’s surprisingly useful, is to ignore the interactions between electrons (the independent electron approximation).

$$\mathcal{H}^Z = \sum_{i=1}^{Z} \frac{p_i^2}{2m} - k_ze^2/r_i \quad (N4.15)$$

Remember that the eigenstates of a single electron bound to a nucleus with charge $Z$ are the hydrogen levels ($\psi_n^Z = \psi_{1s}^Z, \psi_{2s}^Z, \psi_{2p}^Z, \ldots$), except shrunken and shifted upward

15The factor of two is there for $SU(2)$ because the spin is 1/2. For $SO(3)$, infinitesimal generators are antisymmetric matrices, so $S_{i}^{(1)} = J_{i}^{(1)} = i\hbar\epsilon_{ijk}$ in the $xyz$ basis; in the usual quantum basis $m_z = (-1, 0, 1)$ the formula will be different.

16Here $k_z = 1$ in CGS units, and $k_z = 1/(4\pi\varepsilon_0)$ in SI units. We are ignoring the slight shift in effective masses due to the motion of the nucleus.
in binding energy ($E^Z$ more negative):

\[ \mathcal{H}^Z \psi^Z_n = E^Z_n \psi^Z_n \]

\[ \psi^Z_n(r) = \psi^H_n(\lambda_r r) \]

\[ E^Z = \lambda_E E^H \quad (N4.16) \]

(a) By what factor $\lambda_r$ do the wavefunctions shrink? By what factor $\lambda_E$ do the energies grow? (Hint: Dimensional arguments are preferred over looking up the formulas.)

In the independent electron approximation, the many-body electron eigenstates are created from products of single-electron eigenstates. The Pauli exclusion principle (which appears only useful in this independent electron approximation) says that exactly two electrons can fill each of the single-particle states.

(b) Ignoring identical particle statistics, show that a product wavefunction

\[ \Psi(r_1, r_2, r_3, \ldots) = \psi^Z_{n_1}(r_1)\psi^Z_{n_2}(r_2)\psi^Z_{n_3}(r_3) \ldots \quad (N4.17) \]

has energy $E = \sum_i E^Z_{n_i}$.

The effect of the electron-electron repulsion in principle completely destroys this product structure. But for ground-state and excited-state quantum numbers, the language of filling independent electron orbitals is quite useful. However, the energies of these states are strongly corrected by the interactions between the other electrons.

(c) Consider the 2s and 2p states of an atom with a filled 1s shell (one electron of each spin in 1s states). Which state feels a stronger Coulomb attraction from the nucleus? Argue heuristically that the 2s state will generally have lower (more negative) energy and fill first.

Let’s check something I asserted, somewhat tentatively, in lecture. There I said that, for atoms with little spin-orbit coupling, the ground state wavefunction can be factored into a spatial and a spin piece:

\[ \Psi(r_1, s_1; r_2, s_2; r_3, s_3; \ldots) \equiv \psi(r_1, r_2, r_3 \ldots) \chi(s_1, s_2, s_3 \ldots) \quad (N4.18) \]

We’ll check this in the first nontrivial case—the lithium atom ground state, in the independent electron approximation. From part (c), we know that two electrons should occupy the 1s orbital, and one electron should occupy the 2s orbital. The two spins in the 1s orbital must be antiparallel; let us assume the third spin is pointing up $\uparrow_3$:

\[ \Psi^0(r_1, s_1; r_2, s_2; r_3, s_3) = \psi^L_{1s}(r_1)\psi^L_{1s}(r_2)\psi^L_{2s}(r_3) \uparrow_1 \downarrow_2 \uparrow_3. \quad (N4.19) \]

\(^{17}\)The excited states of an atom aren’t energy eigenstates, they are resonances, with a finite lifetime. If you think of starting with the independent electron eigenstates and gradually turning on the Coulomb interaction and the interaction with photons, the true ground state and the resonances are adiabatic continuations of the single-particle product eigenstates—inheriting their quantum numbers.
But this combination is not antisymmetric under permutations of the electrons.

(d) Antisymmetrize \( \Psi^0 \) with respect to electrons 1 and 2. Show that the resulting state is a singlet with respect to these two electrons. Antisymmetrize \( \Psi^0 \) with respect to all three electrons (a sum of six terms). Does it go to zero (in some obvious way)? Can it be written as a product as in eqn N4.18?

**N4.15 Exponentials of matrices.** (Math) 

In quantum mechanics, one often takes exponentials of operators. The exponential of a matrix \( \exp(M) \) can be computed using several different equivalent relations.

First, one can compute it as a power series:

\[
\exp(M) = \sum_{n=0}^{\infty} M^n / n!
\]  \hspace{1cm} \text{(N4.20)}

Let’s take the exponential \( \exp(-i\phi \sigma_2/2) \), where \( \sigma_2 = \begin{pmatrix} 0 & -i \\ i & 0 \end{pmatrix} \) is the second Pauli matrix (also known as \( \sigma_y \)). This is the definition of how spin \( \frac{1}{2} \) particles transform under rotations.

(a) Note that \( \sigma_2^2 = 1 \). Separate the infinite series into even and odd terms, and express \( \exp(-i\phi \sigma_2/2) \) as a linear combination of the identity matrix \( \mathbb{1} \) and the matrix \( \sigma_2 \). In your answer, note that a 360° rotation is not equal to the identity, but to minus the identity!

Secondly, one can compute it as an infinite product of infinitesimal transformations:

\[
\exp(M) = \lim_{n \to \infty} \exp(M/n)^n \\
= \lim_{n \to \infty} (\mathbb{1} + M/n)^n.
\]

This will be the basic trick we use to generate the path-integral formulation of quantum mechanics. It is also the way we generate symmetry operations (like rotations) from infinitesimal generators (like angular momentum). For example, in two dimensions the angular momentum operator is \( J = i\hbar \begin{pmatrix} 0 & 1 \\ -1 & 0 \end{pmatrix} \).

(b) Show that a \( 2 \times 2 \) rotation matrix by an angle \( \theta/n \), in the limit \( n \to \infty \), can be written as \( \mathbb{1} + (C\theta/n)J \). What is the constant \( C \) ? Argue, without calculation, that the product in eqn ?? must generate the finite-angle rotations.

Finally, many matrices which arise in quantum mechanics (symmetric matrices, Hermitian matrices, and the more general category of normal matrices) can be diagonalized by a unitary change of basis: \( D = \begin{pmatrix} \lambda_1 & 0 & \cdots \\ 0 & \lambda_2 & \cdots \\ \vdots & \vdots & \ddots \end{pmatrix} = U\dagger MU \), with \( U\dagger = (U^T)^* = U^{-1} \). Thus \( M^n = (UDU\dagger)^n = UD(U\dagger U)D\cdots DU\dagger = UD^nU\dagger \). For these matrices, we can

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18 Continuous groups like the rotations are called Lie groups. The corresponding infinitesimal generators, and their commutation relations, are called the Lie algebra for the group.
compute the exponential of a matrix by doing a coordinate change to the basis that diagonalizes it:

\[
\exp(M) = \sum_{n=0}^{\infty} M^n / n! = \sum_{n=0}^{\infty} UD^n U^\dagger / n!
\]

\[
= U \left( \sum_{n=0}^{\infty} D^n / n! \right) U^\dagger
\]

\[
= U \left( \begin{array}{ccc}
  e^{\lambda_1} & 0 & \ldots \\
  0 & e^{\lambda_2} & \ldots \\
  0 & 0 & \ldots
\end{array} \right) U^\dagger
\]

(N4.21)

Let’s apply this to the time evolution operator \(\exp(-iHt/\hbar)\) for the Hamiltonian we studied in the Eigen exercise (3.1): \(H = \begin{pmatrix} 0 & -4 \\ -4 & 6 \end{pmatrix}\).

(c) Apply the relation eqn N4.21 to calculate the 2×2 time evolution operator \(\exp(-iHt/\hbar)\) for our Hamiltonian. Apply the resulting time evolution operator to the state \(\psi(0) = \begin{pmatrix} 1 \\ 0 \end{pmatrix}\) to calculate \(\psi(t)\). Also write the time evolved state as \(\sum_n \exp(-iE_n t/\hbar) |n\rangle \langle n| \psi\rangle\), where \(|n\rangle\) are the eigenstates of \(H\). Do the two methods agree?

N4.16 Baker-Campbell-Hausdorff identity. (Expanded upon from Gottfried & Yan exercise 2.13.)

For operators \(A\) and \(B\), we know \(e^{A+B} \neq e^A e^B\) unless \(A\) and \(B\) commute, so \(C = [A, B] = 0\). The BCH theorem tells us that, for any two linear operators \(A\) and \(B\), that

\[
e^A e^B = \exp(A + B + \frac{1}{2}[A, B] + \frac{1}{12}([A, [A, B]] - [B, [A, B]]) \ldots)
\]

(N4.22)

where \(\ldots\) alludes to multiple commutators of even higher order.

We start with the special case where \(A\) and \(B\) do not commute with each other but which both commute with \([A, B]\). In that case, they satisfy

\[
e^{A+B} = e^A e^B e^{-\frac{1}{2}[A,B]}
\]

(N4.23)

(a) Examine the power series of both sides. Show that they agree at low orders.

(b) To prove this, first show that \([B, e^{xA}] = e^{xA}[B, A] x, \) where \(x\) is a scalar (a number, which commutes with everything). Next, define \(G(x) = e^{xA} e^{xB}\) and show that

\[
\frac{dG}{dx} = (A + B + [A, B]x) G.
\]

(N4.24)

Integrate this to obtain the desired result.

(c) Let \(K\) and \(P\) be two noncommuting operators (e.g., the kinetic and potential energy parts of the Hamiltonian \(H = K + P\)). Let \(\epsilon\) be small (e.g., \(i\hbar / \hbar\) in a small time-step \(U(dt) = \exp(-iHdt/\hbar)\). Show that

\[
e^{\epsilon(K+P)} = e^{\epsilon K} e^{\epsilon P} + O(\epsilon^2)
\]

(N4.25)

by explicitly expanding out the power series. What order is the error in the approximation

\[
e^{\epsilon(K+P)} \approx e^{\epsilon K/2} e^{\epsilon P} e^{\epsilon K/2}?
\]

(N4.26)

(Hint: We have a good reason to do the extra work of adding the third step.)
Harmonic oscillators and symbolic manipulation. (Computation) ③

In this exercise, we shall use symbolic manipulation environments (Mathematica or SymPy) to explore the raising and lowering operators \( a \) and \( a^\dagger \). We’ll use them to generate the position-space eigenstates \( \psi_n(x) \), along with their associated Hermite polynomials. We’ll distinguish between analytical calculations (paper and pencil) and symbolic calculations (using the symbolic manipulation package on the computer).

Remember that the Hamiltonian for a simple harmonic oscillator\(^{19}\) is

\[
H = \frac{p^2}{2m} + \frac{1}{2}m\omega^2 x^2. \tag{N4.27}
\]

with

\[
p = -i\hbar \frac{\partial}{\partial x} \tag{N4.28}
\]

The ground state probability distribution is a Gaussian of width \( a_0 = \sqrt{\hbar/2m\omega} \), so

\[
\psi_0(x) = \left(\frac{m\omega}{\pi\hbar}\right)^{1/4} e^{-m\omega x^2/2\hbar}. \tag{N4.29}
\]

For plots, we’ll take constants from McEuen’s bouncing buckyballs (Park et al., “Nanomechanical oscillations in a single-C\(_{60}\) transistor”, Nature 407, 57 (2000)). Thus \( m \approx 60 \times 12\text{amu} \), with an amu = 1.66054e-24 gm, \( \hbar \approx 1.0545716 \times 10^{-27} \text{erg sec} \), and \( \omega = 1.2\text{THz} \).

(a) Do a symbolic integration to check if \( \psi_0 \) is properly normalized. Plot \( \psi_0 \) from \(-4a_0\) to \(4a_0\) using McEuen’s constants. How do the zero-point fluctuations for McEuen’s buckyball compare to the size of an atom?

(b) Define an operator \( H(\psi) \) that symbolically takes a function \( \psi(x) = |\psi\rangle \) and returns another function \( H|\psi\rangle \). Symbolically calculate the ground state energy \( E_0 = H(\psi_0)/\psi_0 \).

(Hint: it should be independent of \( x \).)

Remember that the ladder operators are written in terms of the position and momentum:

\[
a = \sqrt{m\omega/2\hbar}(x + ip/m\omega) \tag{N4.30}
\]

\[
a^\dagger = \sqrt{m\omega/2\hbar}(x - ip/m\omega) \tag{N4.31}
\]

Remember the number operator \( N = a^\dagger a \).

(c) Using the commutation relation \([x,p] = i\hbar\), analytically (paper and pencil) show that \((N + \frac{1}{2})\hbar\omega = H\), \([a,a^\dagger] = 1\), \([N,a^\dagger] = a^\dagger\), and \([N,a] = -a\).

(d) Using the commutation relations from above, analytically show that \( H(a^\dagger)^n\psi_0 = (n + \frac{1}{2})\hbar\omega(a^\dagger)^n\psi_0 \), so \( \psi_n \propto (a^\dagger)^n\psi_0 \).

(e) As in parts (b), symbolically define the operator \( p \) from eqn N4.28. Using it, define the operator \( a^\dagger(\psi) \). Is it normalized? Symbolically calculate \( E_1 = H(a^\dagger\psi_0)/a^\dagger\psi_0 \).

---

\(^{19}\)The spring constant \( K = m\omega^2 \); this gives \( \omega = \sqrt{K/m} \), which may be more familiar.
(f) Symbolically, is \(a^i a^j \psi_0\) normalized? Calculate symbolically the norm of higher powers of \((a^i)^n \psi_0\) until you figure out what we need to divide it by to normalize it.

(g) Analytically calculate the norm of \((a^i)^n \psi_0\), using the commutation relations of part (c). Does it agree with your conclusion of part (f)?

(h) Symbolically define \(\psi_n\) recursively in terms of \(\psi_{n-1}\), using the proper normalization from parts (f) and (g). Evaluate it symbolically for \(n = 1, 2, 3, 4\).

(i) Using McEuen’s buckyball constants, plot \(\psi_n(x)\) for \(n = 1, 2, 3, 4\) for \(-5a_0 < x < 5a_0\).

N4.18 Matrices, wavefunctions, and group representations. (Group reps) (3)

In this exercise, we shall explore the tensor product of two vector spaces, and how they transform under rotations. We’ll draw analogies between two examples: vectors \(\rightarrow\) matrices and single-particle-states \(\rightarrow\) two-particle-wavefunctions.

The tensor product between two vectors is \((v \otimes w)_{ij} = v_i w_j\). The tensor product between two single-particle wavefunctions \(\zeta(x)\) for particle \(A\) and \(\phi(y)\) for particle \(B\) is the product wavefunction \(\Psi(x, y) = \zeta(x)\phi(y)\). If \(H(A)\) and \(H(B)\) are the Hilbert spaces for particles \(A\) and \(B\), the tensor product space \(H(AB) = H(A) \otimes H(B)\) is the space of all linear combinations of tensor products \(\Psi(x, y) = \zeta(x)\phi(y)\) of states in \(H(A)\) and \(H(B)\). Two-particle wavefunctions live in \(H(AB)\).

Let \(\{\hat{e}_i\} = \{\hat{x}, \hat{y}, \hat{z}\}\) be an orthonormal basis for \(\mathbb{R}^3\), and let \(\{\zeta_i\}\) and \(\{\phi_j\}\) be orthonormal bases for the Hilbert spaces \(H(A)\) and \(H(B)\) for particles \(A\) and \(B\).

(a) Show that the tensor products \(\Psi_{ij}(x, y) = \zeta_i(x)\phi_j(y)\) are orthonormal. (The dot product is the usual \(\int dx dy \Psi^* \Psi\).) With some more work, it is possible to show that they are also complete, forming an orthonormal basis of \(H(AB)\).

Suppose the two particles are both in states with total angular momentum \(L_A = L_B = 1\), and are then coupled with a small interaction. Angular momentum addition rules then say that the two-particle state can have angular momentum \(L_{(AB)}\) equal to 2 or 1 or 0: \(1 \otimes 1 = 2 \oplus 1 \oplus 0\). In group representation theory, this decomposition corresponds to finding three subspaces that are invariant under rotations.

The tensor product space \(\mathbb{R}^3 \otimes \mathbb{R}^3\) are normally written as \(3 \times 3\) matrices \(M_{ij}\), where \(M = \sum_{i=1}^{3} \sum_{j=1}^{3} M_{ij} \hat{e}_i \hat{e}_j\). Vectors transform under \(L = 1\), so we would expect matrices to decompose into \(L = 2, 1,\) and 0.

(b) Show that the antisymmetric matrices, the multiples of the identity matrix, and the traceless symmetric matrices are all invariant under rotation (i.e., \(R^{-1} MR\) is in the same subspace as \(M\) for any rotation \(R\)). Which subspace corresponds to which angular momentum?

(c) Consider the \(L = 1\) subspace of matrices \(M\). Provide the (standard) formula taking this space into vectors in \(\mathbb{R}^3\). Why are these called pseudovectors?

I always found torque \(\tau = r \times F\) quite mysterious. (Its direction depends on whether you are right- or left-handed!) Fundamentally, we see now that this is because torque is not a vector—it is an antisymmetric \(3 \times 3\) matrix.
How does this relate back to quantum wavefunctions? Suppose our two $L = 1$ particles are identical, with spins in the same state.

(d) Which angular momentum states are allowed for spin-aligned fermions? For spin-aligned or spinless bosons?

Many physical properties are described by symmetric matrices: the dielectric constant in electromagnetism, the stress and strain tensors in elastic theory, and so on.

N4.19 Molecular rotations. (Quantum) ③

In class, we estimated the frequency of atomic vibrations, by generating a simple model of an atom of mass $AM_P$ in a harmonic potential whose length and energy scales were set by electron physics (a Bohr radius and a fraction of a Rydberg). In the end, we distilled the answer that atomic vibrations were lower in frequency than those of electrons by a factor $\sqrt{M_P/m_e}$, times constants of order one.

Here we consider the frequencies of molecular rotations.

(a) By a similar argument, derive the dependence of molecular rotation energy splittings on the mass ratio $M_P/m_e$.

(b) Find some molecular rotation energy splittings in the literature. Are they in the range you expect from your estimates of part (a)?
in $\Delta t$ for the free particle and the simple harmonic oscillator? For simplicity, let’s ignore the prefactors (coming from the normalizations), and focus on the terms inside the exponentials.

Taking $t = t_0 + \Delta t$ and $x' = x_0 + \dot{x}\Delta t$, expand to first order in $\Delta t$ the terms in the exponential for the free particle propagator (eqn N4.32) and the simple harmonic oscillator (eqn N4.33). Do they agree with Feynman’s formula? (Hint: For the simple harmonic oscillator, the first term is proportional to $1/\Delta t$, so you’ll need to keep the second term to second order in $\Delta t$.)

N4.21 Three particles in a box. (Quantum) ③

(Adapted from Sakurai, p. 4.1)

Consider free, noninteracting particles of mass $m$ in a one-dimensional box of length $L$ with infinitely high walls.

(a) What are the lowest three energies of the single-particle energy eigenstates?

If the particles are assumed noninteracting, the quantum eigenstates can be written as suitably symmetrized or antisymmetrized single-particle eigenstates. One can use a level diagram, such as in Fig. N4.10, to denote the fillings of the single particle states for each many-electron eigenstate.

\[ \text{Fermions} \]
\[ \text{Distinguished} \]
\[ \text{Bosons} \]

Fig. N4.10 Level diagram, showing one of the ground states for each of the three cases.

(b) If three distinguishable spin-1/2 particles of the same mass are added to the box, what is the energy of the three-particle ground state? What is the degeneracy of the ground state? What is the first three-particle energy eigenvalue above the ground state? Its degeneracy? The degeneracy and energy of the second excited state? Draw a level diagram for one of the first excited states, and one of the second excited states (the ground state being shown on the left in Fig. N4.10).

(c) The same as part (b), but for three identical spin-1/2 fermions.

(d) The same as part (b), but for three identical spin-zero bosons.
N4.22  **Rotation matrices.** (Mathematics) 

A rotation matrix $R$ takes an orthonormal basis $\hat{x}, \hat{y}, \hat{z}$ into another orthonormal triad $\hat{u}, \hat{v}, \hat{w}$, with $\hat{u} = R\hat{x}$, $\hat{v} = R\hat{y}$, and $\hat{w} = R\hat{z}$.

(a) Which is another way to write the matrix $R$?

I. $R = (u_1v_1 + v_1w_1 + w_1u_1 \ \cdots )$

II. $R = (\begin{pmatrix} \hat{u} \\ \hat{v} \\ \hat{w} \end{pmatrix})$

III. $R = (\begin{pmatrix} \hat{u} \\ \hat{v} \\ \hat{w} \end{pmatrix})$

IV. $R = \hat{u} \otimes \hat{v} + \hat{v} \otimes \hat{w} + \hat{w} \otimes \hat{u}$

Rotation matrices are to real vectors what unitary transformations (common in quantum mechanics) are to complex vectors. A unitary transformation satisfies $U^\dagger U = 1$, where the dagger gives the complex conjugate of the transpose, $U^\dagger = (U^T)^*$. Since $R$ is real, $R^\dagger = R^T$.

(b) Argue that $R^T R = 1$.

Thus $R$ is an **orthogonal** matrix, with transpose equal to its inverse.

(c) In addition to (b), what other condition do we need to know that $R$ is a proper rotation (i.e., in $SO(3)$), and not a rotation-and-reflection with determinant -1?

I. $\hat{u}, \hat{v}, \hat{w}$ must form a right-handed triad (presuming as usual that $\hat{x}, \hat{y}, \hat{z}$ are right-handed),

II. $\hat{u} \cdot \hat{v} \times \hat{w} = 1$

III. $\hat{w} \cdot \hat{u} \times \hat{v} = 1$

IV. All of the above

One of the most useful tricks in quantum mechanics is multiplying by one. The operator $|k\rangle\langle k|$ can be viewed as a projection operator: $|k\rangle\langle k|\psi\rangle$ is the part of $|\psi\rangle$ that lies along direction $|k\rangle$. If $k$ labels a complete set of orthogonal states (say, the eigenstates of the Hamiltonian), then the original state can be reconstructed by adding up the components along the different directions: $|\psi\rangle = \sum_k |k\rangle\langle k|\psi\rangle$. Hence the identity operator $\mathbb{1} = \sum_k |k\rangle\langle k|$. We’ll use this to derive the path-integral formulation of quantum mechanics, for example. Let’s use it here to derive the standard formula for rotating matrices.

Under a change of basis $R$, a matrix $A$ transforms to $R^T A R$. We are changing from the basis $\hat{x}_1, \hat{x}_2, \hat{x}_3 = |x_i\rangle$ to the basis $|u_j\rangle$, with $|u_n\rangle = R|x_n\rangle$. Since $|u_j\rangle = R|x_j\rangle$, we know $\langle x_i|u_j\rangle = \langle x_i|R|x_j\rangle = R_{ij}$, and similarly $\langle u_i|x_j\rangle = R^T_{ij}$. Let the original components of the operator $A$ be $A_{k\ell} = \langle x_k|A|x_\ell\rangle$ and the new coordinates be $A'_{ij} = \langle u_i|A|u_j\rangle$.

(d) Multiplying by one twice into the formula for $A'$: $A'_{ij} = \langle u_i|\mathbb{1}A\mathbb{1}|u_j\rangle$ and expanding the first and second identities in terms of $x_k$ and $x_\ell$, derive the matrix transformation formula $A'_{ij} = R^T_{ik}A_{k\ell}R_{\ell j} = R^T AR$, where we use the Einstein summation convention over repeated indices.
N4.23 **Trace.** (Mathematics) (2)

The trace of a matrix $A$ is $\text{Tr}(A) = \sum_i A_{ii} = A_{ii}$ where the last form makes use of the Einstein summation convention.

(a) Show the trace has a cyclic invariance: $\text{Tr}(ABC) = \text{Tr}(BCA)$. (Hint: write it out as a sum over components. Matrices don’t commute, but products of components of matrices are just numbers, and do commute.) Is $\text{Tr}(ABC) = \text{Tr}(ACB)$ in general?

Remember from exercise N4.22(b) that a rotation matrix $R$ has its inverse equal to its transpose, so $R^T R = 1$, and that a matrix $A$ transforms into $R^T AR$ under rotations.

(b) Using the cyclic invariance of the trace, show that the trace is invariant under rotations.

Rotation invariance is the primary reason that the trace is so important in mathematics and physics.

N4.24 **Complex exponentials.** (Mathematics) (2)

You can prove the double angle formulas using complex exponentials: just take the real and imaginary parts of the identity $\cos(A + B) + i \sin(A + B) = e^{i(A+B)} = e^{iA}e^{iB} = (\cos A + i \sin A)(\cos B + i \sin B) = \cos A \cos B - \sin A \sin B + i(\cos A \sin B + \sin A \cos B)$.

In a similar way, use complex exponentials and $(x + y)^3 = x^3 + 3x^2y + 3xy^2 + y^3$ to derive the triple angle formulas. Which is true?

(A) $\cos(3\theta) = \cos^3(\theta) - 3 \cos(\theta) \sin^2(\theta)$, $\sin(3\theta) = \sin^3(\theta) - 3 \cos^2(\theta) \sin(\theta)$;

(B) $\sin(3\theta) = \cos^3(\theta) - 3 \cos(\theta) \sin^2(\theta)$, $\cos(3\theta) = 3 \cos^2(\theta) \sin(\theta) - \sin^3(\theta)$;

(C) $\cos(3\theta) = \cos^3(\theta) - 3 \cos(\theta) \sin^2(\theta)$, $\sin(3\theta) = 3 \cos^2(\theta) \sin(\theta) - \sin^3(\theta)$;

(D) $\sin(3\theta) = 3 \cos^2(\theta) \sin(\theta) + \sin^3(\theta)$; $\cos(3\theta) = \cos^3(\theta) + 3 \cos(\theta) \sin^2(\theta)$;

(E) $\cos(3\theta) = \cos^3(\theta) - 3i \cos(\theta) \sin^2(\theta)$, $\sin(3\theta) = 3i \cos^2(\theta) \sin(\theta) - \sin^3(\theta)$.

N4.25 **Dirac $\delta$-functions.** (Mathematics) (3)

Quantum bound-state wavefunctions are unit vectors in a complex Hilbert space. If there are $N$ particles in 3 dimensions, the Hilbert space is the space of complex-valued functions $\psi(x)$ with $x \in \mathbb{R}^{3N}$ whose absolute squares are integrable: $\int dx |\psi(x)|^2 < \infty$.

But what about unbound states? For example, the propagating plane-wave states $\psi(x) = |k\rangle \propto \exp(-ikx)$ for a free particle in one dimension? Because unbound states are spread out over an infinite volume, their probability density at any given point is zero—but we surely don’t want to normalize $|k\rangle$ by multiplying it by zero.

Mathematicians incorporate continuum states by extending the space into a rigged Hilbert space. The trick is that the unbound states form a continuum, rather than a discrete spectrum—so instead of summing over states to decompose wavefunctions $|\phi\rangle = 1\phi = \sum_n |n\rangle \langle n|\phi\rangle$ we integrate over states $\phi = 1\phi = \int dk |k\rangle \langle k|\phi\rangle$. This tells us how we must normalize our continuum wavefunctions: instead of the Kronecker-$\delta$ function $\langle m|n\rangle = \delta_{mn}$ enforcing orthonormal states, we demand $|k\rangle = 1|k\rangle = \int dk |k\rangle \langle k|k\rangle = |k\rangle = \int dk |k\rangle \delta(k - k')$ telling us that $\langle k|k\rangle = \delta(k - k')$ is needed to ensure the useful decomposition $1 = \int dk |k\rangle \langle k|$. 
Let’s work out how this works as physicists, by starting with the particles in a box, and then taking the box size to infinity. For convenience, let us work in a one dimensional box $0 \leq x < L$, and use periodic boundary conditions, so $\psi(0) = \psi(L)$ and $\psi'(0) = \psi'(L)$. This choice allows us to continue to work with plane-wave states $|n\rangle \propto \exp(-ik_n x)$ in the box. (We could have used a square well with infinite sides, but then we’d need to fiddle with wave-functions $\propto \sin(kx)$.)

(a) What values of $k_n$ are allowed by the periodic boundary conditions? What is the separation $\Delta k$ between successive wavevectors? Does it go to zero as $L \to \infty$, leading to a continuum of states?

To figure out how to normalize our continuum wavefunctions, we now start with the relation $\langle m | n \rangle = \delta_{mn}$ and take the continuum limit. We want the normalization $N_k$ of the continuum wavefunctions to give $\int_{-\infty}^{\infty} N_k N_{k'} \exp(-i(k' - k)x)dx = \delta(k' - k)$.  

(b) What is the normalization $\langle x | n \rangle = N_n \exp(ik_n x)$ for the discrete wave-functions in the periodic box, to make $\langle n | n' \rangle = \int_0^L N_n N_m \exp(-i(k'_n - k_n)x)dx = \delta_{mn}$? Write $\mathbb{1} = \sum_n |n\rangle\langle n|$, and change the sum to an integral in the usual way ($\int dk |k\rangle\langle k| \approx \sum_n |n\rangle\langle n|\Delta k$). Show that the normalization of the continuum wavefunctions must be $N_k = 1/\sqrt{2\pi}$, so $\psi_k(x) = \langle x | k \rangle = \exp(ikx)/\sqrt{2\pi}$. (Hint: If working with operators is confusing, ensure that $\langle x | 1 | x' \rangle$ for $0 < x, x' < L$ is the same for $\mathbb{1} = \sum_n |n\rangle\langle n|$ (valid in the periodic box) and for $\mathbb{1} = \int dk |k\rangle\langle k|$ (valid for all $x$).

Notice some interesting ramifications:

I. The fact that our continuum plane waves have normalization $1/\sqrt{2\pi}$ incidentally tells us one form of the $\delta$ function:

$$\delta(k' - k) = \langle k' | k \rangle = \frac{1}{2\pi} \int_{-\infty}^{\infty} e^{-ik'(k-k)x}dx. \quad (N4.35)$$

Also, $\delta(x' - x) = 1/(2\pi) \int_{-\infty}^{\infty} dk \exp(ik(x' - x))$.

II. The same normalization is used for position eigenstates $|x\rangle$, so $\langle x' | x \rangle = \delta(x' - x)$ and $\mathbb{1} = \int dx |x\rangle\langle x|$.

III. The Fourier transform can be viewed as a change of variables from the basis $|x\rangle$ to the basis $|k\rangle$:

$$\tilde{\phi}(k) = \langle k | \phi \rangle = \langle k | \mathbb{1} | \phi \rangle = \int dx \langle k | x \rangle \langle x | \phi \rangle = 1/\sqrt{2\pi} \int dx \exp(-ikx)\phi(x) \quad (N4.36)$$

Note that this definition is different from that I used in the appendix of my book (Statistical Mechanics: Entropy, Order Parameters, and Complexity, http://pages.physics.cornell.edu/~sethna/StatMech/EntropyOrderParametersComplexity.pdf); there the $2\pi$
is placed entirely on the inverse Fourier transform, which here it is split symmetrically between the two, so the inverse Fourier transform is
\[
\phi(x) = \langle x | \phi \rangle = \int dk \langle x | k \rangle \langle k | \phi \rangle = 1/\sqrt{2\pi} \int dk \exp(ikx) \tilde{\phi}(k).
\]

IV. The Dirac \(\delta\)-function can be written in many different ways. It is basically the limit\(^{20}\) as \(\epsilon \to 0\) of sharply-peaked, integral-one functions of width \(\epsilon\) and height \(1/\epsilon\) centered at zero. Let’s use this to derive the useful relation:
\[
\lim_{\epsilon \to 0^+} \frac{1}{x - i \epsilon} = p.v. \frac{1}{x} + i \pi \delta(x).
\]

Here all these expressions are meant to be inside integrals, and p.v. is the Cauchy principal value of the integral:\(^{21}\)
\[
p.v. \int_{-\infty}^{\infty} = \lim_{\epsilon \to 0^+} \int_{-\infty}^{-\epsilon} + \int_{\epsilon}^{\infty}.
\]

(c) Note that \(\epsilon/(x^2 + \epsilon^2)\) has half-width \(\epsilon\) at half-maximum, height \(\epsilon\), and integrates to \(\pi\), so basically (i.e., in the weak limit) \(\lim_{\epsilon \to 0} \epsilon/(x^2 + \epsilon^2) = \pi \delta(x)\). Argue that \(\lim_{\epsilon \to 0} \int f(x)/(x - i \epsilon) dx = p.v. \int f(x)/x dx + i \pi f(0)\). (Hints: The integral of \(1/(1+y^2)\) is \(\arctan(y)\), which becomes \(\pm \pi/2\) at \(y = \pm \infty\). Multiply numerator and denominator by \(x + i \epsilon\).)

N4.26 Eigen Stuff. (Mathematics) \(^{2}\)

Consider an operator for a two-state system \(O = \begin{pmatrix} 0 & -4 \\ 4 & 6 \end{pmatrix}\) Its eigenvectors are \(|e_1\rangle = \frac{1}{\sqrt{5}} (\frac{2}{1})\) and \(|e_2\rangle = \frac{1}{\sqrt{5}} (\frac{-1}{2})\)

(a) What are the associated eigenvalues \(o_1\) and \(o_2\)?

(b) Use \(|e_1\rangle\) and \(|e_2\rangle\) to construct a rotation matrix \(R\) that diagonalizes \(O\), so \(R^T OR = \begin{pmatrix} o_1 & 0 \\ 0 & o_2 \end{pmatrix}\). (Hint: See problem N4.22(a). We want \(R\) to rotate the axes into \(\hat{u} = |e_1\rangle\) and \(\hat{v} = |e_2\rangle\).) What angle does \(R\) rotate by?

(c) Assume that the system is in a state \(|L\rangle = \begin{pmatrix} 1 \\ 0 \end{pmatrix}\) Decompose \(|L\rangle\) into the eigenvectors of \(O\). (Hint: As in Exercise N4.22(d), multiplying \(|L\rangle\) by one is useful.) If the observable corresponding to the operator \(O\) is measured for state \(|L\rangle\), what is the probability of finding the value \(o_1\)? Does the probability of finding either \(o_1\) or \(o_2\) sum to one?

---

\(^{20}\)Clearly this is not a limit in the ordinary sense: the difference between functions does not go to zero as \(\epsilon\) goes to zero, but rather (within \(\epsilon\) of the origin) has large positive and negative values that cancel. It is a weak limit—when integrated against any smooth functions, the differences go to zero.

\(^{21}\)If \(f(x)\) is positive at zero, \(\int f(x)/x dx\) is the sum of minus infinity for \(x < 0\) and positive infinity for \(x > 0\); taking the principal value tells us to find the sum of these two canceling infinities by chopping them symmetrically about zero.
N4.27 Fine and hyperfine structure: Hydrogen and angular momentum addition.

(Angular Momentum) 3

Symmetries have powerful implications for energy eigenstates of composite systems. They are ordinarily the only cause for degenerate states, for example. Here we use rotational symmetry, and the corresponding angular-momentum addition laws, to derive the degeneracies of the hydrogen \( n = 2 \) states.

Including the spin \( \frac{1}{2} \) of the electron and the spin \( \frac{1}{2} \) of the proton, and the four \( n = 2 \) states of hydrogen, there are sixteen degenerate energy eigenstates in Schrödinger’s solution for hydrogen with \( n = 2 \). In this exercise, we shall follow how these energy eigenstates split up when we include the fine splitting and hyperfine splitting. We shall not need to do any calculations with Hamiltonians; we shall just use the rotational symmetry of the Hamiltonian and angular momentum addition rules.

(a) **What is the energy of the \( n = 2 \) state of hydrogen, ignoring spin, relativity, and the nuclear spin?** (Include the fact that the proton and electron have spin \( \frac{1}{2} \) in the degeneracy calculation, but ignore their effects on the energy for now.)

The 2s and 2p states in hydrogen both have \( n = 2 \), and are degenerate to this order. This degeneracy is not due to a straightforward symmetry of the Hamiltonian.\(^{22}\) It is split by terms of order \( \alpha^2 \), where \( \alpha = e^2/\hbar c \approx 1/137 \) is the fine structure constant, representing the importance of relativity.

The relativistic correction to the kinetic energy splits the 2s and 2p states, but does not couple to the electron or proton spin.

(b) **Including these kinetic energy terms, how do the sixteen original states split up in energy?**

The spin-orbit coupling, also of order \( \alpha^2 \), is proportional to \( \mathbf{L} \cdot \mathbf{S} \), where \( \mathbf{L} \) is the angular momentum of the electron and \( \mathbf{S} \) is the spin of the electron. Because it is a dot product, it maintains rotational symmetry.

(c) **Using angular momentum addition rules, discuss what happens to the twelve 2p orbitals after incorporating the spin-orbit coupling. What values of \( j \) are allowed, where \( J = L + S \)? What are the degeneracies of the coupled states?** (Hint: The different energy eigenstates with the same \( J \) are related by rotations. You should not need the form of the interaction to solve this part or the next.)

The splitting due to the spin-orbit interaction is called fine structure, and also arises in heavier atoms. For example, the yellow light from sodium vapor lamps is comprised of two nearby spectral lines, split by the spin-orbit interaction.\(^{23}\)

For states with \( L > 0 \) the coupling to the nuclear spin \( I \) is approximately given by \( \hat{A} \mathbf{I} \cdot \mathbf{J} \). This is called the hyperfine splitting; it is smaller than the fine structure

\(^{22}\)It’s peculiar to the \( 1/r \) potential energy law, and an associated conserved Lenz’s vector. The hydrogen problem can be mapped in an obscure way to the four-dimensional harmonic oscillator: see Sk&N sect. 4.1.

\(^{23}\)Wikipedia also calls the 2s-2p splitting in hydrogen a fine structure effect, but I’m not sure that’s standard. In heavier atoms, the energies of these orbitals (quasiparticle resonance energies, not eigenstates) are shifted primarily not due to relativity, but due to the effects of the other electrons.
splittings because the nucleus is heavy compared to the electron. Again, this interaction maintains rotational symmetry (as it must).

(d) For each of your degenerate families of 2p states in part (c) ignoring the hyperfine splitting, what are the allowed values of $F = I + J$? What degeneracies in the final eigenvalues do you expect?

N4.28 Mystery: Properties of the group character table. (Group Reps)

In the week following this assignment, we shall learn about representations of finite groups. Group representation theory involves new conceptual ideas, new mathematical theorems, and some new calculational methods. Even knowing the ideas and the theorems, I find the calculational methods seem mysterious, almost magical. Let’s try to introduce these tools first, to motivate the lectures to come. I am not pretending to introduce why we do these manipulations—this is an experiment, giving you the mechanics of the calculation before we explain the context in order to motivate your interest.

Consider the following table. It is an expanded version of the character table for the group representations of $C_3v$, the symmetry group of a triangle. But just treat it as a list of row vectors $A_1$, $A_2$, and $E$, along the six ‘directions’ labeled by the six symmetry group elements $g = e, r, r^2, v, rv, \text{ and } r^2v$ in the group $G$.

<table>
<thead>
<tr>
<th>$C_3v$</th>
<th>e</th>
<th>r</th>
<th>r$^2$</th>
<th>v</th>
<th>rv</th>
<th>r$^2$v</th>
</tr>
</thead>
<tbody>
<tr>
<td>$A_1$</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>$A_2$</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>-1</td>
<td>-1</td>
<td>-1</td>
</tr>
<tr>
<td>$E$</td>
<td>2</td>
<td>-1</td>
<td>-1</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Table N4.1: Expanded character table for $C_3v$. The group elements $g = e, r, \ldots$ label the columns; the representations $R = A_1, A_2, \ldots$ label the rows, and the entries are the characters $\chi_R(g)$.

(a) Orthogonality Show that the three character row vectors are orthogonal to one another. Show that the naive “dot product” of a row vector with itself is equal to the number of group elements (called $O(G)$).

Thus the three representations $A_1$, $A_2$, and $E$ are orthonormal using the inner product given by the naive dot product divided by the order of the group:

$$\chi_1 \ast \chi_2 = (1/O(g)) \sum_{g \in G} \chi_1(g)\chi_2(g). \quad (N4.41)$$

Group representations give one matrix $R(g)$ for each abstract symmetry operation $g$. So rotation matrices form a representation of the rotation group. (Mathematicians carefully distinguish between the abstract multiplication table $G$ for a group, and the implementation $R(g)$ of that group in matrix form.) The characters of the group $\chi(g)$
are the traces of these matrices. (Much more about groups and characters will in
lecture.)

For example, we can write a representation of the triangle symmetry group $C_{3v}$ by
thinking of how each symmetry operation permutes the three vertices of the triangle.
Label the three vertices of the triangle by the three unit vectors. Let $R(g)_{ij}$ be one if
vertex $j$ shifts to vertex $i$ under the symmetry operation $g$.

What triangle symmetry corresponds to the six group elements $e, r, \ldots$? We always
use $e$ to represent the ‘do-nothing’ symmetry, so $R(e) = \begin{pmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{pmatrix}$. The matrix $r$ rotates
the triangle $180^\circ$, so $R(r) = \begin{pmatrix} 0 & 1 & 0 \\ 0 & 0 & 1 \\ 1 & 0 & 0 \end{pmatrix}$. The matrix $v$ flips the triangle around the first
vertex, so $R(v) = \begin{pmatrix} 1 & 0 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & 0 \end{pmatrix}$.

(b) What are the characters $\chi(e)$, $\chi(r)$, and $\chi(v)$?

We define the product of two symmetries (say $rv$) as performing the symmetry opera-
tions from left to right (so, flipping by $v$ and then rotating by $r$).

(c) What is the matrix $R(rv)$? Do this two ways. Figure out how the symmetry $rv$
permutes the vertices. Or use the property $R(rv) = R(r)R(v)$; the matrices have the
same multiplication table as the group. What is $\chi(rv)$? Is it the same as that of $e$, $r$, or $v$?

Notice in Table N4.1 that the column vectors labeled by $r$ and $r^2$ are the same, while
$v$, $rv$, and $r^2v$ also share the same characters. This is generally true: the characters of
two elements in the same conjugacy class are always the same. Use this to check your
character for $rv$ in section (c). [We will explain why this is true in lecture.] We put
the two rotations $r$ and $r^2$ into the conjugacy class $C_3$, and we put the three reflections
$v$, $rv$, and $r^2v$ into the conjugacy class $\sigma_v$; the identity $e$ is put into the one-element
class $E$. This allows us to make a more efficient character table (Table N4.2), where the
number of elements in multiply occupied conjugacy classes is included in the column
heading (hence $3\sigma_v$, because there are three $\sigma_v$ rotations). Now, to find the character of
a representation, you only need to compute the trace of one element of each conjugacy
class, and to take the inner product of two characters (eqn N4.41) one can sum over
conjugacy classes but multiply by the multiplicity (number of elements in the class).

<table>
<thead>
<tr>
<th>$C_{3v}$</th>
<th>$E$</th>
<th>$2C_3$</th>
<th>$3\sigma_v$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$A_1$</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>$A_2$</td>
<td>1</td>
<td>1</td>
<td>-1</td>
</tr>
<tr>
<td>$E$</td>
<td>2</td>
<td>-1</td>
<td>0</td>
</tr>
</tbody>
</table>

Table N4.2: Traditional character table for $C_{3v}$

One of the main uses for character tables is for finding decompositions of representations
into irreducible representations. This turns out to be related to Fourier transforms, to
angular momentum addition rules, and to many other standard problems in math-
ematics and quantum mechanics. We shall leave what this means mysterious until
lecture, but let us perform a decomposition of the representation $R$ we have described in parts (b) and (c).

(d) What would the character row for our representation $R$ look like in Table N4.2? Show that the inner product (eqn N4.41) of the representation with itself is an integer, but not one. Irreducible representations have norm one. Take the inner product of $\chi_R$ with the three irreducible representations, and show that they are integers. Any reducible representation can be decomposed into integer numbers of the irreducible representations.

I always find it surprising when my naive dot products work out to be multiples of the size of the group. In more complicated cases, it seems magical.

N4.29 F-electrons and graphene. (Quantum) 3

In this exercise, we shall explore how seven degenerate $f$-electron states of an atom split under a weak perturbation which breaks the rotational symmetry.

Atoms often sit atop surfaces with weak interactions without strong bonding; we describe them as adsorbed. Consider a light atom in an electronic $f$-state (i.e., with $\ell = 3$), adsorbed on a monolayer of graphene (Fig. N4.11). Assume the atom is positioned above a point of hexagonal symmetry, so the symmetry group for the atom is broken from $SO(3)$ to $C_{6v}$.

How do we know this? Why is the symmetry group not just $C_6$? Why is our system not symmetric under $D_{6h}$, the symmetry group of graphene?

(a) What symmetry is exhibited by our adsorbed atom that is not in $C_6$? What symmetry in $D_{6h}$ is not a symmetry of our adsorbed atom?

The character of a spin-$\ell$ representation for $SO(3)$ for a rotation by angle $\theta$ is $\chi^{(\ell)}(\theta) = \sin[(\ell + \frac{1}{2})\theta]/\sin[\frac{1}{2}\theta]$. (Check this for the $\ell = 1$ representation, where you know $\chi^{(1)}$ in terms of $\cos[\theta]$. You’ll need to use L’Hôpital’s rule to evaluate $\chi^{(\ell)}(0)$.)

Six of the symmetry operations in $C_{6v}$ (conjugacy classes $\sigma_v$ and $\sigma'_{v}$) are reflections—in $O(3)$ but not in $SO(3)$. The characters for representations of $O(3)$ are not so commonly studied. Let’s figure them out for the special case of reflections.

Fig. N4.11 Atom adsorbed on graphene.

24The atom is light so that we may ignore the spins of the electrons. A heavy atom would have significant spin-orbit interactions.
Every reflection $\Sigma(\hat{n})$ in $O(3)$ takes the mirror plane into itself, and the perpendicular $\hat{n}$ of the mirror plane to $-\hat{n}$. Thus $\Sigma(\hat{y})$ is a reflection in the $x-z$ mirror plane. Let $R_\hat{n}$ be a rotation that takes the coordinate axis $\hat{y}$ to $\hat{n}$.

(b) Using $R_\hat{n}$, show that all reflections in $O(3)$ are conjugate to $\Sigma(\hat{y})$.

Since the trace is invariant under rotations, and conjugacy in $SO(3)$ is a rotation, and the character is a trace, this means that all reflections will have the same character under representations of $O(3)$. Consider the angular momentum $\ell$ representation of $O(3)$ generated by the rotations of the spherical harmonics $Y^m_\ell(\theta, \phi)$. Remember that $\theta$ is the angle from the $\hat{z}$ axis, and $\phi$ is measured from the $\hat{x}$ axis.

(c) How does $Y^m_\ell$ transform under the reflection $\Sigma(\hat{y})$ in the $x-z$ plane? In the $(2\ell+1)$-dimensional space of $Y^m_\ell$ for fixed $\ell$, what are the elements of the $(2\ell+1) \times (2\ell+1)$ matrix $D_{mm'}$ representing $\Sigma(\hat{y})$? Show that the trace $\chi^{(\ell)}(\Sigma(\hat{y})) = 1$, and hence that the character for all reflections is one in all (integer) representations of $O(3)$, independent of $\ell$.

Table N4.3 gives the character table for $C_{6v}$.

(d) When the f-electron eigenstates are split by the hexagonal crystal field from the graphene, what irreducible representations and degeneracies will be represented? (Hint: Use the orthogonality of the representations to decompose the $\ell = 3$ representation. Also, check that the total number of states equals the number of f-states.) For example, your answer might be “Two nondegenerate eigenstates with reps $A_1$ and $B_2$, and three doublet eigenstates, two with reps $E_2$ and one with rep $E_1$."

N4.30 Juggling buckyballs. (Path Integrals) 3

Paul McEuen in Physics and Jiwoong Park in Chemistry here discovered in 2000 that buckyballs ($C_{60}$ molecules) bounce inside their transistors.\textsuperscript{25} Here use path integrals to discuss how buckyballs evolve under juggling. (We’ll focus on juggling one buckyball, by throwing it straight up into the air and waiting for it to fall down.) The Lagrangian for the buckyball is

$$\mathcal{L} = \frac{1}{2}m\dot{y}^2 - mgy. \quad (N4.42)$$

(a) In classical mechanics, if the buckyball starts and ends at \( y = 0 \) and travels for a time \( 2\Delta t \), how high \( y_{\text{peak}} \) must its trajectory reach at the midpoint?  (Hint: Nothing tricky yet.)

Feynman tells us that the propagator for a particle starting at \((y = y_i, t = t_i)\) and ending at \((y = y_f, t = t_f)\) is a path integral over all trajectories \( y(t) \):

\[
\langle y_f | U(t_f - t_i) | y_i \rangle = \int \int \int_{y_i, t_i}^{y_f, t_f} \mathcal{D}[y(t)] \exp \left( i/\hbar S[y(t)] \right) \tag{N4.43}
\]

where the three integral signs represent a suitably normalized integral over all paths \( y(t) \). We, like Feynman, will make a discrete “trapezoidal rule” approximation to the propagator. As a rough example, we’ll do two segments and only one intermediate point \( y_2 \):

\[
S[y(t)] \approx \left[ \frac{1}{2} m \left( \frac{y_3 - y_2}{\Delta t} \right)^2 - \frac{1}{2} mg(y_1 + y_2) - \frac{1}{2} mg(y_2 + y_3) + \frac{1}{2} m \left( \frac{y_2 - y_1}{\Delta t} \right)^2 \right] \Delta t. \tag{N4.44}
\]

(b) What intermediate point \( y_2^* \) minimizes the trapezoidal action (eqn N4.44), for general \( y_1 \) and \( y_3 \)? For the symmetric path \( y_1 = y_3 = 0 \), how does this compare to the peak of the trajectory in part (a)? What is the action \( S^* = S[y_2^*] \) for this symmetric minimum action trajectory?  (Note: we’re doing an approximation; the heights need not be the same. Hint: Check units of \( S^* \). Also, does it have the right sign?)

(c) What is our one-point trapezoidal approximation to the propagator

\[
\langle y = 0 | U(2\Delta t) | y = 0 \rangle \tag{N4.45}
\]

(Request: Please write your answer factoring out the contribution from the minimum action part \( S^* \). Hints: Don’t forget the “weight factor” from Sakurai. You can check that you’ve included the right number of weight factors by checking the units of your propagator: at \( t_f = t_i \), for example, \( \langle y_f | U(0) | y_i \rangle = \delta(y_f - y_i) \) has units of inverse length. Also, \( \int_{-\infty}^{\infty} dx \exp(-iAx^2) = \sqrt{\pi/|A|} \).)

N4.31 Solving Schrödinger: Nuclear decays and resonances.  (Computation) ③
N4.32 **Resonances: α-decay.** (Quantum) \(^3\)

Fig. N4.12 **One-dimensional nuclear potential.**

In this exercise, we solve a one-dimensional model of radioactive α-decay, where a nucleus ejects a particle formed by two protons and two neutrons (a Helium-4 nucleus).

We assume that the strong force minus the Coulomb repulsion provides a constant potential for the α particle inside a nucleus of radius \(R\), which for simplicity we shall assume is zero. At the edge of the nucleus in the real world, the (short-range) strong interaction drops rapidly to zero, but the Coulomb repulsion decays slowly with distance, leading to a tunneling barrier. We model this barrier with a \(δ\)-function of strength \(U > 0\) \(^{26}\) (see Fig. N4.12). Both inside and outside the nucleus, the potential is zero:

\[
V(x) = Uδ(x ± R)
\]

(The attractive case \(U < 0\) is a model for the hydrogen molecule, and is discussed for example in Wikipedia’s *Double Delta Potential* article, http://en.wikipedia.org/wiki/Delta_potential#Double_Delta_Potential.)

Parts (a)-(c) of this exercise solve analytically for the energy eigenstates, but getting them correct is important for the later parts.\(^27\)

Our Hamiltonian has a symmetry which allows us to choose energy eigenstates that are even (\(ψ_E\)) or odd (\(φ_E\)).

(a) **What symmetry of the Hamiltonian is this? Given an energy eigenstate \(ζ_E(x)\) with mixed symmetry (in particular, \(ζ_E\) is not odd), construct an even eigenstate of the same energy (ignoring the overall normalization).**

In this exercise, we will be interested in the family of even eigenstates \(ψ_E\) which can be nonzero at \(x = 0\), and for which \(ψ'_E(0) = \partial ψ_E/\partial x|_{x=0} = 0\). To solve for these even energy eigenstates, there are three steps.

\(^{26}\) In one-dimensional quantum mechanics, the first derivative of the wave-function jumps where the potential has a \(δ\)-function. Find details in a textbook or on the Web.

\(^{27}\) Feel free to check your answers by solving Schrödinger’s equation numerically, approximating \(δ(x - R) = (1/\sqrt{2πσ^2}) \exp(-x^2/(2σ^2))\) for \(σ\) as small as is numerically convenient.
First, we deduce the form of the wavefunction. Note that, away from the δ-function, the wavefunction has wave-vector \( k(E) = \sqrt{2mE}/\hbar \); it is convenient to label the wavefunctions by \( k(E) \) instead of \( E \). Using the boundary condition at zero, we write the wavefunction for \( |x| < R \) as \( \psi_k^{\text{in}} = A_k \cos(kx) \), with an overall amplitude \( A_k \). For \( x > R \), we write the wavefunction as a standing sine wave instead of \( \psi_k^{\text{out}} = B \sin(kx + \Delta_k) \). Note that there is a continuum of \( \psi_k \) eigenstates, so it is proper for us to use the δ-function normalization \( \langle \psi_k | \psi_{k'} \rangle = \delta(k - k') \).

(b) Show that \( B = 1/\sqrt{\pi} \) for our continuum wavefunction to be properly normalized. (Hints: Since we’re studying only even eigenstates, \( k \geq 0 \). Also, because the region \( |x| < R \) is finite, we can ignore it for the normalization in an infinite box.)

Second, we impose the conditions induced by the δ-potential at the edge of the nucleus.

(c) Write the condition on \( A_k \) and \( \Delta_k \) given by imposing continuity of \( \psi_k(x) \) at \( x = R \). Write the conditions on \( A_k \) and \( \Delta_k \) given by the discontinuity of \( \psi_k' \) imposed by the δ-function potential (see note 26). For convenience, write your answers from here on in terms of the unitless ratio \( \bar{U} = 2mRU/\hbar^2 \).

Third, we solve for the eigenstates of our Hamiltonian that are nonzero at \( x = 0 \).

(d) Use the conditions of part (c), solve for \( A_k^2 \). (Trick: Arrange the two equations of part (c) to be \( \sin(kR + \Delta_k) = \cdots \) and \( \cos(kR + \Delta_k) = \cdots \), where \( \cdots \) is independent of \( \Delta_k \). Sum the squares of the right-hand sides: what must the sum be equal to?)

We now consider the decay of an α-particle injected into this potential at \( x = 0 \). That is, consider an initial wavefunction \( \Psi(x) = \delta(x) \).29

(e) What is the probability \( P(k) \) of being in eigenstate \( \psi_k \)? (Write your answer abstractly in terms of \( \psi_k(x) \). This you can do without solving parts (a-d).)

(f) Plot the probabilities \( P(k) \) versus \( kR \) with \( \bar{U} = 30 \) and for \( 0 < kR < 10 \).

In the limit \( U \to \infty \), the nucleus should approximate a particle in a box of size \( 2R \). In that limit, the injection of an α-particle can only occur at certain discrete energies—the nuclear eigenstates \( \mathcal{E}_m^\infty \) of a free particle in a box of size \( 2R \).

(g) Compare the peaks you found in part (f) to the wavevectors for the particle-in-a-box states. Why are you missing half of the peaks?

(h) (Extra credit) Change variables from \( P(k) \) to \( P(E) \) by using \( dE/dk \).31 Using the FWHM of the peaks in \( P(E) \), estimate the lifetimes of the first three even resonances

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28 For \( x < -R \), we use the even symmetry of \( \psi_E \) to set \( \psi_k = \psi_k^{\text{out}}(-x) = B \sin(-kx + \Delta_k) \). Note that we are solving for standing waves in this problem. For other purposes, scattering waves or outgoing waves might be preferable.

29 This is a nuclear version of tunneling from an STM tip; \( P(E) = P(k(E)) (dk/dE) \) measures the local density of states for the α particle at the center of the nucleus.

30 The position eigenstate \( \Psi(x) = |x = 0\rangle \) is δ-function normalized, with \( \langle x|x' \rangle = \delta(x-x') \). Hence the “probability” \( P(k) \) integrates to infinity, and not to one. You can alternatively think of this calculation as the first step in evaluating the Green’s function \( G(x',t';0,0) \) from \( x = t = 0 \), which evolves an initial packet \( \delta(x) \) from the origin.

31 The overall normalization of these densities of states may be off by a factor with dimensions of length.
of our nucleus (either numerically or analytically). Calculate the integrated probability for being in each of these three resonances. Do they go to the particle-in-a-box values as $U \to \infty$?

N4.33 Solving Schrödinger: Alpha decay, Green’s functions, and resonances. (Computation) ³

(a)

N4.34 Harmonic Fermi sea. (Quantum, fermions) ³

$N$ identical spin $\frac{1}{2}$ Fermions are subject to a three-dimensional simple harmonic-oscillator potential. Ignore any mutual interactions between the particles.

(a) Show that the change in the ground state energy when adding an additional particle jumps at certain “magic” $N$. These are analogous to (but not the same as) the atomic numbers of the noble gases in atomic physics.³² Give the first four of these magic numbers.

The effective potential for nuclear matter is smooth near the center of the nucleus, so the magic numbers for the number of protons $Z$ and neutrons $N$ in the nucleus are similar to those found in the harmonic oscillator potential.

(b) Show, if $N$ is large, that the Fermi energy is approximately $E_F = \hbar \omega \sqrt{3N}$, and the ground state energy is $E_B = (3/4)\hbar \omega \sqrt{3N}^4$.

The number of neutrons in heavy nuclei is larger than the number of protons, because it costs extra Coulomb energy to push protons into the nucleus. (Protons can turn into neutrons by emitting positrons in a $\beta^+$ decay.) But when there are more neutrons, they become more costly because their Fermi energy is higher (the Pauli exclusion principle forces them into a high-energy, unoccupied harmonic oscillator state). This is incorporated, in the high-$N$ limit (ignoring the shell structure) into a “Pauli term” in the semi-empirical mass formula for estimating nuclear binding energies.

(c) If there are $Z = A/2 - \Delta/2$ protons and $N = A/2 + \Delta/2$ neutrons in a 3D harmonic oscillator of frequency $\hbar \omega$, what is the change in the ground state energy $E_B(Z,N) - E_B(A/2,A/2)$ for small $\Delta$, to second order in $\Delta$?

This does not quite give the form used in the semi-empirical mass formula (Pauli energy $= -a_A(A-2Z)^2/A$).³³ It is usual to use the free Fermi gas in a confining sphere instead of a 3D harmonic oscillator to model the nuclear potential; the latter gives a different denominator.

³²Noble gases are particularly stable; the binding energy of electrons filling the closed shell is unusually high, and the binding energy for the electron for the next alkali metal is unusually low. This corresponds to a jump in the energy per particle at the atomic number of the noble gas.

³³Note that the semi-empirical mass formula gives the binding energy, which is a constant minus the ground state energy—hence the minus sign.
In this exercise, we examine the periodic table to gain insight into the amazing but mysterious utility of thinking of electrons in atoms as filling independent orbitals.

The electron-electron repulsion in atoms, molecules, and solids is almost as large as the electron-nuclear attraction. However, in physics and chemistry we discuss many-electron systems in the language of noninteracting electrons. Thus metals and semiconductors have electron and hole excitations, atoms have 1s electrons near the nucleus and 3d electrons in the transition metals, and diamond has tetrahedral coordination because of $sp^3$ hybridization of the 2s/2p and 3s/3p orbitals. All of these labels would be valid if the electrons did not interact—but the true electron wavefunctions for an $N$-electron atom is a complex function of $3N$ variables that in no way factors into orbitals.

First, let us pretend that electrons did not interact with one another: the Coulomb repulsion between electrons is set to zero. Remember that the first four angular momentum states $L = 0, 1, 2, 3$ are called s, p, d, and f for obscure historical reasons.\(^{34}\) Remember also that the hydrogen spectrum has energy levels $E_n = -(1/n^2)13.6$eV, with the $n^{th}$ energy level including states with $L = 0, \ldots, n - 1$.

(a) How many noninteracting electrons can fit into the s, p, d, and f states? How many total electrons can fit into each level $E_n$? If we assume the elements in this noninteracting world kept the same names for each atomic number, which elements would be noble gases?

These atomic levels are not just a fiction. One can use a high-energy electron or X-ray to eject an electron from an atom (creating a 'core hole' in, say, the 1s state). An electron from the 2s state may then transition into the hole, simultaneously ejecting a third 2p electron from the nucleus. This emitted particle is called an Auger electron. The kinetic energies of the Auger electrons will be given, to a good approximation, by the energy difference $-(E_{1s} - E_{2s} - E_{2p})$, just as one would expect if the electrons did not interact.\(^{35}\) Auger transitions are often used to identify chemical species.

(b) In our noninteracting world, we can have multi-electron atoms with a hydrogen nucleus. What would the energy be of the emitted Auger electron in the above transition?

The problem here, however, is if the electrons did not interact the 1s core hole would be an eigenstate that would not decay.\(^{36}\) This last point is the key to the puzzle. The eigenstates of the noninteracting electrons become resonances when interactions are turned on. Their energies are complex, with imaginary parts that correspond to their decay widths.

---

\(^{34}\)There is yet a different notation for these states in the X-ray community ($K$, $L_1$, $\ldots$, $M_5$), involving also the total angular momentum $J = L + S$ of the electrons.

\(^{35}\)Corrections for the interaction energy between the 2s and 2p holes in the final state, and corrections for the electronic screening energies, improve the accuracy of this rough estimate.

\(^{36}\)The 2s-1s transition could happen by two-photon emission, but would not eject the other electron. Of course, allowing photon emission also makes the electronic excited states into resonances, not eigenstates. Photon emission is small because the fine structure constant $\alpha = e^2/\hbar c \sim 1/137$ is small. The electron-electron interaction is not small, which makes the question more subtle.
lifetimes—here due to their interaction with other electrons. Electrons and holes in metals and semiconductors become *quasiparticles*—quasielectrons and quasiholes that carry around a screening cloud or atomic polarization cloud, and decay eventually into lower-energy excitations.

The resonance energy levels of a 1s core excitation of an atom with nuclear charge $Z$ should be roughly given by the 1s ionization energy of the corresponding helium-like two-electron ion; the outer electrons are mostly farther away from the nucleus and thus will not lower the interaction energy. But the 2s electron resonance energy will roughly be given by the 2s state for an atom with charge $Z - 2$, since the 1s electrons will partially screen the nucleus.\(^{37}\) The different orbitals (1s, 2s, 2p, 3s, 3p, 3d, \ldots) will get shifted in energy away from their hydrogenic values because of this screening. Various approximate quantum methods for incorporating this screening energy can be developed (e.g., Hartree-Fock).

We can gain some understanding of the power of this picture, without getting buried in arcane Auger tables, by examining the periodic table. Find a periodic table that conveniently shows the fillings of the different subshells (1s, 2s, 2p, 3s, \ldots).

(c) *Find an ordering of the energies of the subshells that mostly explains the ground state level filling of the different atoms, up to Radon (i.e., ignoring the late radioactive ones). What are the exceptions to your rule?*

(d) *Are the noble metals closed-shell like the noble gases?*

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**N4.36 Nuclear Shell Model.** (nuclear) \(^{3}\)

Nuclear physics is challenging. Unlike atomic physics, where the interaction of electromagnetism with matter is weak (of order $\alpha = 1/137$), the interaction between quarks (and hence nucleons) is strong—so we cannot use perturbation theory. Unlike condensed matter physics, where we can assume many particles and hence describe liquids and crystals with continuum theories, the number of protons and neutrons in a nucleus is relatively small.

Nuclear physics is thus a field where creative use of simple models is widespread. We have explored earlier the use of *random matrix theory* to describe excitations of nuclei. In this exercise, we shall introduce both the *nuclear shell model* and the nuclear *semi-empirical mass formula*.

We shall use real data, downloaded directly from the Web. There is a table of atomic masses of various isotopes at the Atomic Mass Data Center, [http://amdc.in2p3.fr/masstables/Ame2011int/mass.mas114](http://amdc.in2p3.fr/masstables/Ame2011int/mass.mas114).

\(^{37}\) Actually, in electronic structure calculations one often treats the core electrons with a *pseudopotential*. Instead of treating the 2s excitation of a $Z$ atom as a 2s excitation of a $Z - 2$ atom, one treats it as a 1s excitation of a $Z - 2$ atom, but with a potential that smoothly blurs out the nucleus inside the radius of the 1s shell. They choose this smooth potential to match the scattering amplitudes of the original core-electron ion.
(a) Download the mass table directly into your computational environment. After reading them, drop the first 39 lines of header. For each \( N \) and \( Z \) in the table with experimental data\(^{38} \) make a table of the nuclear names (e.g., 56Fe) and of the “mass excess” column (converted to floats). The mass excess is the atomic mass minus one amu (atomic mass unit) per nucleon, where an amu = 931.494061 MeV is one-twelfth the energy of \(^{12}\)C. The semi-empirical mass formula estimates the nuclear mass, which means we need to add back \( A \times \text{amu} \) and subtract the \( Z \) electron masses \( m_e \). Store the binding energy indexed by \( Z \) and \( N \):

\[
\text{nuclear mass} = \text{mass excess} + A \text{amu} - Z m_e. \tag{N4.46}
\]

\[
\text{binding energy} = Z m_p + N m_n - \text{nuclear mass}. \tag{N4.47}
\]

Create a matrix with these entries (zero where no experimental data), and make a plot of the nonzero entries.\(^{39} \)

(Hint: the main feature of this plot will be that larger \( A \) have larger nuclear masses.)

The semi-empirical mass formula treats the nucleus primarily as a drop of liquid, with a “condensation energy” \( a_V A \), where \( A = N + Z \) is the number of nucleons, and a surface tension energy \( a_S A^{2/3} \). (If the nucleus is a liquid of nucleons of roughly constant density, then its radius \( R \sim A^{1/3} \) and hence the surface area \( \sim R^2 \sim V^{2/3} \).) Packing \( Z \) protons into the nucleus costs a Coulomb energy (as in \( (Z q)^2 / R \)) of \(-a_C Z^2 / A^{1/3}\); this Coulomb energy is why there are more neutrons than protons in heavy nuclei.

In addition, there are two quantum terms. The first is the Pauli term, which is related to Exercise N4.34; if the number of neutrons \( N \) is different than the number of protons \( Z \), the neutron Fermi energy will be different than the proton Fermi energy, and there will be an energy cost that grows as the difference. Since either positive or negative differences will reduce the binding energy,\(^{40} \) it is natural to approximate this difference with the square \((N - Z)^2\). It turns out that the magnitude of this effect grows weaker as \( A \) grows, so using \( A = N + Z \) we approximate the Pauli energy \(-a_A(N - Z)^2 / A = -a_A(A - 2Z)^2 / A. \)

The second quantum term is a pairing energy. If the number of protons (or neutrons) is even, the energy is lower than if it is odd, by a pairing energy that goes roughly as \( \delta = a_P / A^{1/2} \). (Some sources say this is due to the fact that the second nucleon can “go into the same orbital” as the first; others say this even-odd term is due to spin-orbit interactions; others blame it on the formation of superconducting Cooper pairs of protons or neutrons.) The pairing energy is considered zero for even-odd nuclei, positive (binding) for even-even nuclei, and negative (destabilizing) for odd-odd nuclei. (It is interesting to note that there are only four stable nuclei with an odd number of protons and an odd number of neutrons.)

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\(^{38}\) Warning: data with # instead of decimal points are theoretical extrapolations. Drop those points.

\(^{39}\) You can use a masked array to do this in Python; in Mathematica you can set ColorRules \( \rightarrow 0.0 \rightarrow \text{White}. \)

\(^{40}\) Usually there will be more neutrons
We use Rohlf’s values (Rohlf: Modern Physics from a to Z, James William Rohlf, Wiley, 1994, section 11.3, quoted from Wikipedia), who give \( a_V = 15.75 \text{ MeV} \), \( a_S = 17.8 \text{ MeV} \), \( a_C = 0.711 \text{ MeV} \), \( a_A = 23.7 \text{ MeV} \), and \( a_P = 11.18 \text{ MeV} \).

(b) Create a function \( \text{SemiEmpiricalMassFormula}(Z, N) \) that evaluates this formula. Check against the actual value for \( ^{56}\text{Fe} \) from your calculation in part (a). \(^{41}\) (Hint: Iron has \( Z = 26 \) and hence \( ^{56}\text{Fe} \) has \( N = 30 \). Your answer should be within less than a percent of the experimental value. This can also help debug part (a). For assistance in checking for typos, I got volumeTerm=882.0 MeV, surfaceTerm=-260.54 MeV, coulombTerm=-125.628 MeV, PauliTerm=-6.77 MeV, \( \delta = 1.49 \text{ MeV} \).)

One major piece of physics that the semi-empirical mass formula misses are the magic numbers. These are explained by assuming that, like atoms, the nucleons approximately fill orbitals that have shells (like the 1s, 2s, 2p, . . . for atoms). Just as noble gases arise when a shell fills and the next orbital has a big jump in energy, so nuclei have especially stable states. Notice, though, that there is only one type of electron, while there are two types of nucleons. So we expect especially stable nuclei when either the proton or the neutron number is magic: horizontal or vertical bands in a color plot of mass versus \( Z \) and \( N \).

(c) Make a 2D color plot of the difference between the semi-empirical mass formula and the actual binding energy, as a grid of \( Z \) versus \( N \). Don’t plot colors (or plot white) where experimental data is not available. \(^{42}\) Note the horizontal and vertical bands where the theory underestimates the binding energy. Estimate these ‘magic numbers’. Do your estimates agree with the Harmonic Fermi sea estimates of an earlier exercise? Do they agree with Wikipedia’s list 2, 8, 20, 28, 50, 82, 126?

N4.37 Mirror path integrals. (Path Integrals) \(^3\)

![Fig. N4.13 Qbit weirdness](image)

A photon, passing through a pair of half-silvered mirrors \( H_1 \) and \( H_2 \), undergoes quantum interference between different paths.

One of the most compelling examples of Qbits and their weirdness is provided by the example of photons and half-silvered mirrors. Fig. N4.13 shows a photon\(^{43}\) coming

\(^{41}\)Iron \( ^{56} \) is one of the most stable nuclei. It dominates the endpoint of fusion reactions in stars. Lower mass nuclei tend to fuse; higher masses tend to fission. I hear that \( ^{62}\text{Ni} \) is even more stable, but is not accessible easily in nuclear reactions.

\(^{42}\)You can use a masked array to do this in Python; in Mathematica you can set ColorRules \( \rightarrow 0.0 \rightarrow \text{White} \).

\(^{43}\)We assume, as before, that the polarization of the photon lies perpendicular to the plane of the paper, so that it remains unchanged and hence unimportant to the interference.
from the left in a superposition $|\gamma\rangle$, through a set of mirrors, to two detectors named Alice (A) and Bob (B). We work in the basis $|1\rangle = (\frac{1}{\sqrt{2}})$ representing the upper of the two beams at a given position, and $|0\rangle = (\frac{\sqrt{2}}{1})$ representing the lower of two beams. As discussed in Schumacher & Westmoreland, the half-silvered mirror $H_2$ approximately acts as a unitary transformation $H_2 = \frac{1}{\sqrt{2}} \left[ \begin{array}{cc} 1 & 1 \\ 1 & -1 \end{array} \right]$. The first mirror $H_1$, with its mirrored side on the top, changes the sign of the beam reflecting from its top, hence $\frac{1}{\sqrt{2}} \left[ \begin{array}{cc} -1 & 1 \\ 1 & 1 \end{array} \right]$. The product $G = H_2H_1$ is analogous to the propagator, or Green’s function, for this system.\(^{44}\)

(a) What is $G$? Given the impinging wave $|\gamma\rangle$ from the left, what are the probabilities $P_A$ and $P_B$ that Alice and Bob will see the photon? (Hint: Remember Bob did not see anything when the initial photon came from below.)

We can develop a kind of discrete path integral representation of the propagator $G$ by writing

$$G = \mathbb{1}H_2\mathbb{1}H_1\mathbb{1} \quad \text{(N4.48)}$$

$$= \sum_{x_i=0}^{1} \sum_{x_m=0}^{1} \sum_{x_f=0}^{1} |x_f\rangle\langle x_f| H_2 |x_m\rangle\langle x_m| H_1 |x_i\rangle\langle x_i|. \quad \text{(N4.49)}$$

Here $i$, $m$, $f$ representing the initial states, the states in the middle, and the final (detected) states, and $\mathbb{1} = |1\rangle\langle 1| + |0\rangle\langle 0|$. If we assume the initial photon is coming from the bottom left ($x_i = 0$), there are four remaining paths in this sum.

(b) Give the four amplitudes contributed by these four paths. Which ones contribute to $\langle 1|G|0\rangle = G_{10}$? Which ones contribute to $\langle 0|G|0\rangle = G_{00}$? Which go to Bob? Do the sum of the amplitudes going to Bob equal zero (as they should)?

\(^{44}\)Note two confusing things in our notation. First, the photon moves from left to right (hitting $H_1$, then $H_2$), but the matrices describing the evolution propagate from right to left ($H_2H_1$). Second, $|1\rangle = (\frac{1}{\sqrt{2}})$ is a vector whose first element is one [zeroth in Python/C], and $|0\rangle = (\frac{\sqrt{2}}{1})$ is a vector whose second element is one [first in Python/C].
Imagine an electron traversing an electron-mirror array, impinging from below. The mirrors $H_1$ and $H_2$ have the same effect on the amplitudes as the former half-silvered ones did for the photon. Here, though, we thread a solenoid between the upper and lower paths in the middle region, enclosing a net magnetic flux $\Phi$ pointing upward out of the page. The field is zero outside the solenoid, and you may ignore the electron’s spin.

(c) As a function of $\Phi$, what is the probability that an initial electron will be seen by Alice? What values of $\Phi$, in multiples of the elementary flux quantum $\Phi_0 = \hbar c/e = 2\pi\hbar c/e$, prevent Alice from seeing any electrons? (Remember, the initial electron comes from below. Hints: $\oint_C \mathbf{A} \cdot d\ell = \Phi$ if the path $C$ encircles the solenoid counterclockwise once. The path-integrand amplitude for $\mathbf{x}(t)$ in a field $\mathbf{A}$ gains a phase $\zeta = \int (q/c) \mathbf{A}(x) \cdot dx/\hbar$. The charge on an electron is $q = -e$.)

N4.38 Harmonic oscillator spectrum: The propagator. (Path integrals) ③

(a) Show that the trace of the propagator can be written in terms of the energy eigenvalues:

$$\int_{-\infty}^{\infty} K(x,t_2;x,t_1)dx = \sum_n \exp(-iE_n(t_2-t_1)/\hbar).$$

(N4.50)

(Hint: write

$$K = \langle x_2 | U(t_2-t_1) | x_1 \rangle = \langle x_2 | e^{-iH(t_2-t_1)/\hbar} | x_1 \rangle = \langle x_2 | 1 | e^{-iH(t_2-t_1)/\hbar} | x_1 \rangle$$

and insert a complete set of energy eigenstates for $\mathbf{1}$. )
(b) Sum the geometrical series in eqn N4.50 for a one-dimensional harmonic oscillator of frequency $\omega$.

The propagator for the harmonic oscillator is

$$K_{HO}(x_2, t_2; x_1, t_1) = \sqrt{\frac{m\omega}{2\pi i \hbar \sin(\omega(t_2 - t_1))}} \exp \left[ \frac{im\omega}{2\hbar \sin(\omega(t_2 - t_1))} \left\{ (x_2^2 + x_1^2) \cos(\omega(t_2 - t_1)) - 2x_2x_1 \right\} \right].$$

(There is a typo in Sakurai’s formula 2.6.18).

(c) Write $K_{HO}(x, t_2, x, t_1) = f(t) \exp(-iA(t)x^2)$. Evaluate the trace. Show that you get the same answer as part (b). (Hint: Use the Gaussian integral formula $\int_{-\infty}^{\infty} \exp(-iA(t)x^2) = \sqrt{\pi/(iA)}$. You may want to use the half-angle formula $\sin(a/2) = \sqrt{1/2(1 - \cos(a))}$.)

N4.39 Evolving Schrödinger: Coherent states. (Computation) (3)

In this exercise, we shall build upon the numerical work of exercise 3.6 (free particle evolution), exercise 4.2 (Baker-Campbell-Hausdorff identity), and exercise 4.3 (Coherent States). We shall solve the time-dependent Schrödinger equation for a harmonic oscillator in its ground state, and after the ground state is translated to the side by a distance $x_0$.

The time-dependent Schrödinger equation for our one-dimensional quantum system is:

$$i\hbar \frac{\partial \psi}{\partial t} = -\frac{\hbar^2}{2m} \frac{\partial^2 \psi}{\partial x^2} + V(x) \psi = H\psi = H_{\text{kin}}\psi + H_{\text{pot}}\psi$$

$$\psi(t) = U(t)\psi(0) = e^{-iH_{\text{kin}}t/\hbar}e^{-iH_{\text{pot}}t/\hbar}\psi(0)$$

As in the last exercise, we use the constants for McEuen’s bouncing buckyballs, with $m = 60m_C \sim 12 * 60m_p$ and the frequency to $\omega = 10^{12}$ radians/sec, and will evaluate it at $N_p = 200$ points spanning $L = 30a_0$.

In the free particle example (no potential energy), we advanced time by $dt$ by multiplying the Fourier transform by $U_{\text{kin}}(k, dt) = \exp[i(h^2k^2/2m)dt/\hbar]$. If, on the other hand, there were no kinetic energy (infinite mass), we could solve for the time evolution $\psi(x, t) = U_{\text{pot}}(t)\psi(x, t = 0)$ by multiplying $\psi(x)$ in real space by a time-dependent phase depending on position:

$$\psi(x, t + dt) = U_{\text{pot}}(dt)\psi(x, t) = e^{-iH_{\text{pot}}dt/\hbar}\psi(x, t) = e^{-iV(x)dt/\hbar}\psi(x, t).$$

To approximately solve Schrödinger’s time evolution, we alternate advancing the wave function in real space and Fourier space, using the Baker–Campbell–Hausdorff formula of exercise 4.2(b):

$$\psi(t + dt) = e^{-iH_{\text{kin}}t/h - iH_{\text{pot}}t/2\hbar}\psi(0) \approx e^{-iH_{\text{pot}}t/2\hbar}e^{-iH_{\text{kin}}t/h}e^{-iH_{\text{pot}}t/2\hbar}\psi(0) = U_{\text{pot}}$$
(a) Define the two arrays \text{UkinTildeDt} and \text{UpotDtOver2}. Define the initial wavefunction \( \psi[0](x) \). (Hint: If your implementation stores \( \psi[n][x] \) as a two-dimensional complex array, you may want to allocate it and initialize \( \psi[0][x] \) as part of that array.)

(b) Evolve the wavefunction to a time equal to twice the period \( P \) of the oscillator, in steps of \( dt = P/100 \), storing your answer after each step. Plot \( \psi(x, P/5) \), showing the real part, the imaginary part, and the absolute value all on the same graph. (Why don’t we plot \( |\psi^2(x)| \) on this graph?) If possible, animate these three curves; otherwise, plot several snapshots until you see the evolution. What happens to the probability density? Why? What happens to the real and imaginary parts? Why?

(c) Now shift the wavefunction \( \psi(x, t = 0) = \psi_0(x - x_0) \), with \( x_0 = 10a_0 \), where \( a_0 \) is the RMS width of the ground state wavefunction (see exercise 3.6). Time evolve as in part (b). How does the evolution compare to a classical particle in the harmonic well?

(d) Using your answer to exercise 4.3(d), write the initial wavefunction for part (c) in terms of a coherent state. What is \( \lambda \)?

For photons and phonons and other harmonic systems, the coherent states evolve just as classical particles would.

N4.40 Coherent State Evolution. (Operator algebra) ③

Consider the annihilation operator \( a \) for a simple harmonic oscillator, transformed into the time-dependent Heisenberg-representation operator \( a_H(t) \):

\[
a_H(t) = e^{iHt/\hbar}ae^{-iHt/\hbar} = U^\dagger(t)aU(t). \quad \text{(N4.54)}
\]

The time evolution for an operator in the Heisenberg representation is given by the commutator with the Hamiltonian, so

\[
\frac{da_H}{dt} = \frac{iH}{\hbar}a_H - a_H \frac{iH}{\hbar} = -i/\hbar[a_H, H]. \quad \text{(N4.55)}
\]

You may use the fact that the Hamiltonian for the harmonic oscillator in the Schrödinger representation is \( H = \hbar\omega(a^\dagger a + \frac{1}{2}) \), and that \([a, a^\dagger] = 1\).

(a) Calculate \([a_H, H]\), and write it in terms of \( a_H \). What is \( da_H/dt \)? (Simplify your answers until they only involve \( a_H \) and constants, not \( H \) or \( a \).)

(b) Show that \( a_H(t) = \exp(-i\omega t)a_H(0) = \exp(-i\omega t)a \) is the solution to the time evolution you found in part (a). (Hint: This can also be a check for part (a).)

We discovered in a computational exercise that the probability density for a displaced harmonic oscillator ground state oscillates like a classical particle with the oscillator frequency \( \omega \). In another exercise, we showed that a displaced harmonic oscillator ground state is one example of a \textit{coherent state}, an eigenstate of the annihilation operator \( a \):

\[
a|\lambda\rangle = \lambda|\lambda\rangle, \quad \text{(N4.56)}
\]

which is also normalized \( \langle \lambda|\lambda\rangle = 1 \). Here \( \lambda \in \mathbb{C} \) can be any complex number.
(c) In the Schrödinger representation show that a coherent state \( |\lambda\rangle \) evolves after a time \( t \) to a state \( |\xi(t)\rangle = U(t)|\lambda\rangle \) which is also an eigenstate of the annihilation operator \( a \). What is its eigenvalue \( \tilde{\lambda} \)? (Hints: Multiply \( a|\xi(t)\rangle = aU(t)|\lambda\rangle \) on the left by \( 1 = U(t)U(-t) \) and use part (b). You don’t need to compute \( U(t)|\lambda\rangle \), you just need to show it is an eigenstate of \( a \).)

Since there is only one coherent state with eigenvalue \( \tilde{\lambda} \), our evolved state \( U(t)|\lambda\rangle = C|\tilde{\lambda}\rangle \) for some constant \( C \). Since time evolution conserves probability (and hence \( U(t) \) is unitary), \( \langle\lambda|U^{\dagger}(t)U(t)|\lambda\rangle = |C|^2 = 1 \), so \( C \) is a pure phase.

The phase \( C \) depends on time and \( \lambda \). Let us solve for the case of the oscillator in the ground state.

(d) Calculate \( C(t) \) for the special case \( \lambda = 0 \). (Hint: the coherent state with \( \lambda = 0 \) is the ground state of the harmonic oscillator. You don’t need to know the solutions of previous sections to solve this.)

N4.41 Decoherence. (Density Matrices)

In this exercise, we will explore the effects of decoherence on a quantum system using density matrices and the Bloch sphere. We will study the dynamics of spin-1/2 particles in a magnetic field, with and without decoherence. We will work in the \( z \)-spin basis, and denote the spins pointing parallel to and anti-parallel to the \( z \)-direction by \( |\uparrow_z\rangle \) and \( |\downarrow_z\rangle \). The spins are subjected to a magnetic field \( \vec{B} = B\hat{x} \) in the \( x \)-direction. Convince yourself that the Hamiltonian modeling this is \( \mathcal{H} = -\mu_0 B(|\uparrow_x\rangle\langle\uparrow_x| - |\downarrow_x\rangle\langle\downarrow_x|) \).

(a) Write this Hamiltonian in the \( z \)-spin basis.

(b) Suppose the initial wavefunction is \( |\psi(t = 0)\rangle = |\uparrow_z\rangle \). Solve the Schrödinger equation to find \( |\psi(t)\rangle \). Do you observe that the spin oscillates between \( |\uparrow_z\rangle \) and \( |\downarrow_z\rangle \)? What is the frequency \( \omega \) of the oscillation?

Recall from problem 11.1 that any \( 2 \times 2 \) density matrix can be written as \( \rho = \frac{1}{2}(1 + \vec{n} \cdot \vec{\sigma}) \).

The vector \( \vec{n} \) is called the Bloch vector, and always has norm \( |\vec{n}| \leq 1 \), forming the solid Bloch sphere. (Remember \( \sigma_x = (0 \ 1 \ 0 \ 0 \ 1 \ 0 \ 0 \ 0 \ 1) \), \( \sigma_y = (0 \ -i \ 0 \ 0 \ i \ 0 \ 0 \ 0 \ -i) \), and \( \sigma_z = (1 \ 0 \ 0 \ 0 \ 0 \ 1 \ 0 \ 0 \ 0) \).)

(c) Calculate \( \rho(t) \) for \( |\psi(t)\rangle \) from your calculation in part (b), in terms of \( \omega \). Calculate \( \vec{n}(t) \), and use the double angle formulas to simplify your answer. Geometrically, what is the trajectory of \( \vec{n}(t) \)? Show that this agrees with your solution to exercise 11.1, \( d\vec{n}/dt \propto -\vec{B} \times \vec{n} \).

(d) Show that the eigenvalues of a general \( 2 \times 2 \) density matrix \( \rho = \frac{1}{2}(1 + \vec{n} \cdot \vec{\sigma}) \) are \( \frac{1}{2}(1 \pm |\vec{n}|) \). What is the entropy \( S = -k_B \text{Tr}(\rho \log \rho) \) of a general density matrix in terms of \( \vec{n} \)? (Hint: use the basis in which \( \rho \) is diagonal, and your eigenvalues.) Show that the zero-entropy pure states are those on the surface \( |\vec{n}| = 1 \) of the Bloch sphere. (Hint: \( x \log x \) is negative for \( 0 < x < 1 \), and equal to 0 at the end points \( x = 0 \) and \( x = 1 \).)

Does your solution \( \vec{n}(t) \) from part (c) stay a zero entropy pure state, as it should?

---

45. As opposed to the Heisenberg representation of part (b).
46. Developed in collaboration with Bhuvanesh Sundar.
47. That exercise had funny units, but the form of the equation and the sign should agree.
Decoherence arises in a system due to interaction with a large environment. Essentially, the universe is constantly looking at our system, and as a result of interaction with the rest of the universe, our spins get entangled with the universe. Since we observe only the spins and do not observe the infinitely many degrees of freedom in the rest of the universe, it appears to us that the spins lose information about any coherences they may have developed.

(e) How does a general density matrix \( \rho = (\rho_{11} \rho_{12} \rho_{21} \rho_{22}) \) written in the \( z \) basis change when its \( s_z \) component is measured? Show that the effect of a measurement in the \( z \)-basis is to project \( \vec{n} \) onto the \( z \)-axis.

For the remainder of the exercise, we consider the evolution under the specific Hamiltonian \( \mathcal{H} \) you studied in parts (a) through (c). We shall model decoherence as a measurement being done on the spins with small probability \( \Gamma \) per unit time. In a small time interval \( \delta t \), the spin is measured in the \( z \)-basis with a probability \( \Gamma \delta t \), and not measured with a probability \( 1 - \Gamma \delta t \), and then the system evolves for a time \( \delta t \).

(f) What is \( \vec{n}(t + \delta t) \) in terms of \( \vec{n}(t) \), including first the possibility of observation and then the time evolution from \( \mathcal{H} \)? Write a differential equation for the components of \( \vec{n}(t) \) by taking \( \delta t \to 0 \).

(g) Show that \( n_z \) obeys the second-order differential equation for a damped harmonic oscillator, \( a^2 n_z / dt^2 + \eta dn_z / dt + \omega^2_0 n_z = 0 \). What are \( \eta \) and \( \omega_0 \) in terms of \( \Gamma \) and \( \omega \)?

(h) What is the long-time limit for \( \vec{n} \)? For \( \rho \)? For the entropy?

N4.42 Quantum Algorithms.\(^48\) (Quantum Information Processing) \(^4\)

Are quantum computers faster than our standard classical computers?

Clearly, we need to define our terms here—since factoring 143 (the current quantum-computing world record) doesn’t take long on a classical computer. The key question is how the computer time would scale for large problems. Factoring \( M \)-digit numbers on a quantum computer takes no more than \( O(M^3) \) time (that is, some constant times \( M^3 \), using Shor’s algorithm), while the most efficient known method for factoring on classical computers takes \( O(e^{1.9M^{1/3} (log M)^{2/3}}) \). For large numbers of digits, quantum computers win (if they can be built). There are a few other problems where classical computers are known to be much slower than quantum computers: Grover’s algorithm for searching an unsorted database, Simon’s algorithm, …

Here we explore a somewhat artificial problem, solved in the quantum case using the Deutsch-Josza algorithm.\(^49\) This will also introduce the reversibility of quantum computing, and the use of gates—unitary operators that transform Qbits to execute the quantum computer program.

The Deutsch-Josza algorithm considers functions that map \( n \) bits to one bit. Let us denote the \( n \) bits as \( x_0, x_1, \ldots, x_{n-1} \), where \( x_0, x_1, \ldots, x_{n-1} \) are all 0 or 1. Let

48 Developed in collaboration with Bhuvanesh Sundar, based on an exercise by Paul Ginsparg.
49 There are many discussions of the Deutsch-Josza algorithm in the literature—feel free to consult them. If you find one that is particularly pertinent, reference it properly in your writeup.
\[ x = \sum_{m=0}^{n-1} 2^m x_m \] denote the integer represented by the bit sequence \( x_0, x_1, \ldots, x_{n-1} \); \( x_0 \) is the least significant bit and \( x_{n-1} \) is the most significant bit. We’ll also denote \( |x_0⟩|x_1⟩ \cdots |x_{n-1}⟩ \) as \( |x⟩ \).

Let \( f \) be a function that maps the \( n \) bits \( x_0, x_1, \ldots, x_{n-1} \) to one bit (that is, either True or False, one or zero). For example, \( f \) could be a function Prime that returns True if the integer \( x \) is prime, or Even that returns True if the integer is divisible by two, or Big that returns True if the integer is greater than or equal to \( 2^n \). Our algorithm is not concerned with implementing \( f(x) \), but with testing properties of an unknown \( f \) by sampling its output. For example, testing whether \( f \) is a constant function (either True for all possible \( x \), or False for all arguments) is a challenge for classical computers.

(An experiment may find a thousand Trues in a row, but to be sure that the function always returns True one must test all \( 2^n \) choices of \( x \).) We define a balanced function to be one which returns True for exactly half of the possible inputs. Thus Even and Big above are balanced, but Prime is neither balanced nor constant.

(a) Write the four possible functions \( f(x_0) \) for \( n = 1 \) (two possible inputs, two possible outputs). Which are constant? Which are balanced? For larger \( n \), most possible functions are neither balanced nor constant.

Deutsch and Josza considered the artificial problem of distinguishing between balanced and constant functions. Let us define DJ functions to be those functions guaranteed to be either balanced or constant. Given that \( f \) is a DJ function, can a quantum computer probing \( f \) distinguish between the two cases faster than a classical computer? Let us first consider how a classical computer would solve this.

(b) Argue that in the worst case, the \( n \)-bit DJ function \( f \) would have to be called \( 2^{n-1} + 1 \) times in order to determine for certain whether it is balanced or constant.

Our challenge is to use a quantum computer program to do this calculation with one operation of the operator \( f \). How do we set this up?

A quantum computer performs unitary operations on Qbits to execute the program. Unitary operations are reversible,\(^{50}\) indeed, the only irreversible step in a perfect quantum computer is the macroscopic observer reading the answer. This means that no quantum computer can perform the classical AND operation, for example—since \( AND(x_0, x_1) \) is False for three different values of \( x_0 \) and \( x_1 \), it would throw out information that could not be retrieved. The workaround is to encode the answer in a final Qbit \( y \). So if \( n = 2 \) and \( f(x_0, x_1) = AND(x_0, x_1) \) (a function that is neither balanced nor constant), we could implement \( f \) on a quantum computer by writing a program that took \( |x_0⟩|x_1⟩ |y⟩ \) and returned \( U_f(|x_0⟩|x_1⟩ |y⟩) = |x_0⟩|x_1⟩ |y \oplus AND(x_0, x_1)⟩ \) where \( \oplus \) is addition modulo 2. If you input \( |x_0⟩|x_1⟩ |y = 0⟩ \), the output value of \(|y⟩ \) gives \( f(x) \). If you input \( |x_0⟩|x_1⟩ |y = 1⟩ \), the output value of \(|y⟩ \) gives \( 1 \oplus f(x) = NOT(f(x)) \)—this feature is important to keep \( U_f \) reversible. \( U_f \) is also linear: for example, \( U_f((\alpha|0⟩ + \beta|1⟩)|0⟩ |y⟩) = \alpha|0⟩ |0⟩ |y \oplus f(0, 0)⟩ + \beta|1⟩ |0⟩ |y \oplus f(1, 0)⟩ \).

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\(^{50}\)The reverse operation is \( U^\dagger = U^{-1} \).
(c) Show that $U_f$ is reversible for the case where $f = \text{AND}$ by giving an explicit method for reconstructing $x_0$, $x_1$, and $y$ from $x_0$, $x_1$, and $y \oplus \text{AND}(x_0, x_1)$. Then show in general that $U_f$ is its own inverse for any $n$-bit function $f$.

We are now given a quantum computer operation that evaluates an unknown DJ function $f$: $U_f(|x_0\rangle|x_1\rangle \cdots |y\rangle) = |x_0\rangle|x_1\rangle \cdots |y \oplus f(x)\rangle$.

Just as a classical computer can be made of AND gates, NOT gates, OR gates, etc., so a quantum computer is composed of gates that manipulate one or two Qbits by application of unitary operators. The single-Qbit gates are thus $2 \times 2$ unitary matrices.

(d) In the basis $|0\rangle = (\frac{1}{\sqrt{2}}, \frac{1}{\sqrt{2}})$ and $|1\rangle = (\frac{1}{\sqrt{2}}, -\frac{1}{\sqrt{2}})$, write the single-Qbit gate $\text{NOT}$ as a $2 \times 2$ matrix. Show that the Hadamard gate, written $H = 1/\sqrt{2}(1 1 \ 1 -1)$, is unitary. We can implement both the NOT gate and the $H$ gate on electron Qbits, for example, by exposing them in a magnetic field with a suitable direction and orientation.

Our strategy will be to apply $U_f$ not to a Qbit product that corresponds to a classical bit sequence $|x_0\rangle|x_1\rangle \cdots |x_{n-1}\rangle$, but rather to a Qbit string that represents a quantum superposition of all possible classical bit sequences. Let us first consider the case $n = 1$. Our strategy is to use the Hadamard gate to create a superposition of bit sequences and then apply $U_f$, and then re-apply the Hadamard gate to find out whether our function is constant or balanced. We shall abuse notation to use $H^{n+1}|x\rangle|y\rangle$ to mean $(H|x\rangle)(H|x_1\rangle) \cdots (H|x_{n-1}\rangle)(H|y\rangle)$.

(e) Starting with the case of $n = 1$ Qbit plus $|y\rangle$, initialize our two Qbits to $|\Psi_0\rangle = |x_0\rangle|y\rangle = |0\rangle|1\rangle$. Apply the Hadamard operation on both Qbits (exposing them both to the same magnetic field). What is the resulting superposition? Apply $U_f$ for the four cases of $f$ you found in part (a), and then apply the Hadamard transformation on both the Qbits again. What is the measured final value of $x_0$ for the constant functions with $n = 1$? For the balanced functions?

You should have found that you could conclusively say if $f$ were constant or balanced with just one call to $U_f$.

Now that you have worked out the $n = 1$ case, let us generalize to arbitrary $n$. The algorithm proceeds in the same way. We initialize each of the $n$ Qbits $|x_0\rangle, |x_1\rangle, \ldots, |x_{n-1}\rangle$ to $|0\rangle$, and $|y\rangle$ to $|1\rangle$, so $|\Psi_0\rangle = |0\rangle^n|1\rangle$. We perform the Hadamard operation on all the Qbits, so $|\Psi_1\rangle = H^{n+1}|\Psi_0\rangle = (H|0\rangle)^n(H|1\rangle)$. We pass them through $U_f$, so $|\Psi_2\rangle = U_f|\Psi_1\rangle$. We perform another Hadamard operation on all the Qbits, $|\Psi_3\rangle = H^{n+1}|\Psi_2\rangle$. Finally, we measure the overlap with the initial state, $|\langle \Psi_0|\Psi_3\rangle|^2$, measuring the probability that $x_0 = 0$, $x_1 = 0$, $\ldots$, $x_{n-1} = 0$, $y = 1$.

Let us do this step by step. The initial state of the Qbits is $|\Psi_0\rangle = |0\rangle^n|1\rangle$. A Hadamard operation is then applied on all of them. The state of the Qbits after this operation is $|\Psi_1\rangle = H^{n+1}|\Psi_0\rangle = (H|0\rangle)^n(H|1\rangle)$.

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51 I apologize for the shift in notation: I used the in Exercise N4.43, $|1\rangle = (\frac{1}{\sqrt{2}}, \frac{1}{\sqrt{2}})$ and $|0\rangle = (\frac{1}{\sqrt{2}}, -\frac{1}{\sqrt{2}})$.

52 The special case $n = 1$ of the Deutsch-Josza algorithm is called the Deutsch’s algorithm.
(f) Write \(|\Psi_1\rangle\) as a superposition of \(|x\rangle|y\rangle\) for all possible \(n\)-bit binary numbers \(x\) and both values of \(y\). Show that the probabilities of being in these states are all equal (but the amplitudes may have different signs).

Now the Qbits are passed through \(U_f\). The state of the Qbits after passing through \(U_f\) is \(|\Psi_2\rangle = U_f|\Psi_1\rangle\). When \(f\) is a constant function, \(U_f\) changes the Qbit \(y\) in the same way for all arguments \(x\).

(g) If \(f\) is a constant function, show that \(|\Psi_2\rangle\) is a constant times \(|\Psi_1\rangle\). What is this constant if \(f \equiv 0\)? If \(f \equiv 1\)? Show that the measured values of \(x_0, x_1, \ldots, x_{n-1}\) in \(|\Psi_3\rangle\) after the final Hadamard operation are all 0, so in particular that \(|\langle 0|\Psi_3\rangle|^2 = 1\).

(Hint: Do not try to apply \(H^{n+1}\) on \(|\Psi_2\rangle\) written as a superposition of several terms. Instead, decompose \(|\Psi_2\rangle\) as a product of a state for each Qbit (\(|\Psi_2\rangle = \Pi_{0\leq i < n}|\phi_i\rangle\) where \(|\phi_i\rangle\) is a superposition of \(|0\rangle\) and \(|1\rangle\)), and use the fact that \(H^2 = 1\).

Hence for a constant function, the result of our quantum computation always has unit probability of returning the state with all \(x_i = 0\) and \(y = 1\).

When \(f\) is a balanced function, the value in the Qbit \(y\) is changed differently for different arguments \(x\); for half of those \(2^n\) arguments, \(y\) is left unchanged, and for half of those arguments, \(y\) is flipped (from 0 to 1 or vice versa). Let us illustrate this with an example: let us consider the function Even, which returns True if the integer \(x\) is divisible by two.

(h) What is the least significant bit of an integer if it was even? If the integer was odd? Argue that whether the Qbit \(y\) is flipped by \(U_f\) or not is determined solely by the least significant bit \(x_0\) in \(|\Psi_1\rangle\). We know that the Qbits were in a product state (a product of single Qbits) \(|\Psi_1\rangle = (H|0\rangle)^n(H|1\rangle)\) before passing through \(U_f\). Show that the Qbits are in a product state after passing through \(U_f\) as well (i.e. \(|\Psi_2\rangle = \Pi_{0\leq i < n}|\phi_i\rangle\), and write this product explicitly. Is \(|\Psi_2\rangle\) different from \(|\Psi_1\rangle\)? What are the measured values of the Qbits \(x_0, \ldots, x_{n-1}\) after the final Hadamard operation? (Hint: Perform the Hadamard operation on each term in the product above, and use the fact that \(H^2 = 1\).

You should have found that \(x_0, \ldots, x_{n-1}\) are measured to be something other than all zeros. The Deutsch-Josza algorithm states that for any balanced function \(f\), the probability of measuring \(x_0, \ldots, x_{n-1}\) to be all zeros is 0. We’ll prove this in the following way.

(i) Show that for an arbitrary balanced function \(f\),

\[
|\Psi_2\rangle = U_f|\Psi_1\rangle = \frac{1}{2^{n+1}} \sum_{0 \leq x < 2^n} (-1)^{f(x)}|x\rangle(|0\rangle - |1\rangle) \tag{N4.57}
\]

\[
= \frac{1}{2^{n+1}} \left( \sum_{x : f(x) = 0} |x\rangle - \sum_{x : f(x) = 1} |x\rangle \right) (|0\rangle - |1\rangle).
\]

After the final Hadamard operation, show that the probability of measuring \(x_0 = 0\), \(x_1 = 0\), \ldots, \(x_{n-1} = 0\), \(y = 1\) is zero, i.e. \(|\langle 0|(H^{n+1}|\Psi_2\rangle) = 0\). (Hint: Rather than cal-
culating \(\langle \Psi_0 | \Psi_3 \rangle\), calculate the same quantity in the form \(\langle \Psi_1 | \Psi_2 \rangle = \langle \Psi_0 | H^{n+1} | \Psi_2 \rangle = \langle \Psi_0 | \Psi_3 \rangle\), and use eqn N4.57.)

Hence with one application of the function \(f\), with 100% certainty a constant function returns the initial state and a balanced function with 100% certainty will never return the initial state.

It is amazing that we could determine whether a DJ function \(f\) is constant or balanced in just one evaluation of \(U_f\). The Deutsch-Josza algorithm achieves an exponential speedup over its classical counterpart. The problems considered in the above (Deutsch and Deutsch-Josza) algorithms may seem far removed from applications to real world problems, but these algorithms paved the way for more complicated and powerful algorithms.

Why are we still factoring 143? The great challenge in building quantum computer is decoherence, the tendency of Qbits to interact with the environment and go from quantum superpositions into mixtures.

N4.43 **Supersymmetric harmonic oscillator.**\(^{53}\) (Quantum, Supersymmetry) \(^{3}\)

One of the main predictions of supersymmetry\(^{54}\) is that each particle comes with a supersymmetric partner with the same mass but with opposite statistics.\(^{55}\) For example, the fermionic electron is paired with the bosonic selectron. Supersymmetry is also a potential symmetry of nature, with an unusual connection to the translational symmetries in space and time (the Poincaré group). Finally, supersymmetry allows one to calculate remarkable things about certain Hamiltonians. In this exercise, we shall explore a “zero-dimensional”\(^{56}\) example of a supersymmetric Hamiltonian, and try to illustrate each of these features of supersymmetry.\(^{57}\)

Remember the commutation relations for creation and annihilation operators suitable for bosons

\[
[a, a^\dagger] = 1 \quad [a, a] = [a^\dagger, a^\dagger] = 0, \quad (N4.58)
\]

and fermions

\[
\{b, b^\dagger\} = 1 \quad \{b, b\} = \{b^\dagger, b^\dagger\} = 0. \quad (N4.59)
\]

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\(^{53}\)Developed in collaboration with John Stout, Fall 2013.

\(^{54}\)The footnotes in this problem are meant as inspiration—tying it to fundamental ideas in theoretical physics. *None of the footnotes are necessary or useful for solving the problem*—ignore them if you wish.

\(^{55}\)Supersymmetric partners have the same mass as long as supersymmetry is unbroken. We expect supersymmetry to be *spontaneously broken* at low energy scales, given that we have not yet detected any supersymmetric partners of the Standard Model particles.

\(^{56}\)We often talk about quantum field theories in \(d\) spatial dimensions and one time dimension as \(d+1\)-dimensional field theories: our space-time is thus 3+1 dimensional. We can view nonrelativistic quantum mechanics as a \(d = 0\) quantum field theory, and it is in this regard that we consider the supersymmetric Hamiltonians described here as “zero-dimensional” or 0+1-dimensional.

\(^{57}\)There are a number of discussions of the supersymmetric harmonic oscillator and zero-dimensional supersymmetry in the literature and on the Web. Feel free to consult these. If you find one particularly useful, reference it properly in your writeup.
where \([A, B] = AB - BA\) is the commutator and \(\{A, B\} = AB + BA\) is the anticommutator.\(^{56}\) For this simple example, we take our bosons and fermions to be noninteracting, so their creation and annihilation operators commute,

\[
[a, b] = [a, b^\dagger] = [a^\dagger, b] = [a^\dagger, b^\dagger] = 0.
\]  
(N4.60)

In one dimension, the Hamiltonian of the simple harmonic oscillator of frequency \(\omega\) can be written either in terms of \(x\) and \(p\):

\[
H_B = \frac{p^2}{2m} + \frac{1}{2}m\omega^2 x^2
\]  
(N4.61)

or in terms of the creation and annihilation operators

\[
H_B = \hbar\omega(a^\dagger a + \frac{1}{2}).
\]  
(N4.62)

Here \(\frac{1}{2}\hbar\omega\) is the ground state energy of the harmonic oscillator—the zero-dimensional analogue of the *vacuum energy* in field theory.

The harmonic oscillator Hamiltonian can be written in a more symmetric way by using the anticommutator.

(a) Show that \(H_B = \frac{1}{2}\hbar\omega \{a^\dagger, a\}\). Is the vacuum energy still \(\frac{1}{2}\hbar\omega\)?

Note that we’re now calling the ladder operators \(a\) and \(a^\dagger\) creation and annihilation operators. In this new language, the \(n^{th}\) excited state of the harmonic oscillator can be viewed as a state with \(n\) bosons.

Define a “fermionic harmonic oscillator” in analogy to the bosonic one, \(H_F = \frac{1}{2}\hbar\omega [b^\dagger, b]\).

Again, we can view the \(n^{th}\) excited state as a state of \(n\) fermions.

(b) What is the ground state energy of \(H_F\)? How many fermions are in the ground state, in this new language? What is the energy of the state with one fermion?

(c) If we write the zero-fermion state as \(|0\rangle = (\begin{smallmatrix} 0 \\ 1 \end{smallmatrix})\) and the one-fermion state as \(|1\rangle = (\begin{smallmatrix} 1 \\ 0 \end{smallmatrix})\), then write \(b, b^\dagger\), and \(H_F\) in terms of the three Pauli matrices \(\sigma_x, \sigma_y,\) and \(\sigma_z\). Check that your form for \(b, b^\dagger\) satisfy the anticommutation relations of eqn N4.59 (Remember \(\sigma_x = (\begin{smallmatrix} 0 & 1 \\ 1 & 0 \end{smallmatrix})\), \(\sigma_y = (\begin{smallmatrix} 0 & -i \\ i & 0 \end{smallmatrix})\), and \(\sigma_z = (\begin{smallmatrix} 1 & 0 \\ 0 & -1 \end{smallmatrix})\).)

We can write our first supersymmetric Hamiltonian by adding the boson and fermion harmonic oscillators:

\[
H_S = H_B + H_F = \frac{1}{2}\hbar\omega \left( \{a^\dagger, a\} + [b^\dagger, b]\right).
\]  
(N4.63)

\(^{56}\)Be sure to avoid getting confused by our multiple uses of the terms boson and fermion in this exercise. There are really three different ways we use the terms, each extremely useful and compelling. They are:

N4.1 the objects which vibrate or have spins, that produce harmonic oscillators or two-state systems,

N4.2 the 'primitive' bosons (and fermions) which are excitations within a harmonic oscillator (e.g., \(N\) bosons = \(N\)th excited state inside the vibrating object)

N4.3 the composite objects inside the supersymmetric Hamiltonian that merge zero or more 'primitive' bosons and fermions.

\(^{59}\)Here we use the convention \(|1\rangle = (\begin{smallmatrix} 1 \\ 0 \end{smallmatrix})\) and \(|0\rangle = (\begin{smallmatrix} 0 \\ 1 \end{smallmatrix})\), instead of the notation used in quantum computing.
Note that the ground state energy for this Hamiltonian is zero.\textsuperscript{60}

This supersymmetric Hamiltonian is not particularly difficult to solve. Because there is no interaction between the bosonic and fermionic parts of the Hamiltonian, the solution separates and the eigenstates are just products $\psi(x)\chi(s)$, and the energy of the eigenstate is the sum of the Fermi and Bose energies.

Remember that a composite particle with an odd number of primitive fermions is a fermion—so half of our eigenstates represent composite bosons, and half represent composite fermions.

(d) Solve for the energies for the eigenstates of $\mathcal{H}_S$. Which eigenstates represent composite fermions? Which composite bosons? Draw the level diagram for $\mathcal{H}_S$, with the first few composite boson eigenenergies as a column of horizontal lines on the left, and the first few composite fermion eigenenergies on the right. On each line, write the number of primitive bosons and fermions making up the composite. Is there a composite fermion state for each composite boson state? What state is the exception? We shall hitherto drop the “composite” label. If we interpret the energy of a state as the mass of a particle,\textsuperscript{61}, supersymmetry gives us for every fermion a boson with the same mass.

The fact that our Hamiltonian has (almost) one fermion state for each boson state is a result of an unusual symmetry of the Hamiltonian. To see this, let’s define an operator, called the supercharge,

\[ Q = b \left( \frac{p}{\sqrt{m}} + i \sqrt{m} \omega x \right) = i \sqrt{2} \hbar \omega ba^\dagger. \tag{N4.64} \]

(Remember that $x = \sqrt{\hbar/2m}(a^\dagger + a)$ and $p = i \sqrt{m} \omega \hbar/2(a^\dagger - a)$.)

(e) Show that $[\mathcal{H}_S, Q] = 0$. (Hence $Q$ is a symmetry of the Hamiltonian.) Show that $Q$ acting on a fermion state gives a constant times a boson state of the same energy, and that $Q^\dagger$ acting on a boson state almost always gives a constant times a fermion state of the same energy. Which of the ground states is the exception to this rule? Show that this ground state is an eigenfunction of $Q$ and $Q^\dagger$ with eigenvalue zero.\textsuperscript{62}

Supersymmetry has been shown (by Haag, Lopuszanski, and Sohnius\textsuperscript{63}) to be the only way to consistently extend the symmetries of spacetime. Spacetime has a spatial

\textsuperscript{60} That is a hint for part (b).

\textsuperscript{61} We can motivate this by remembering that we are dealing with a theory with zero spatial dimensions, and so the usual relativistic energy of a particle (which should correspond to a eigenstate of our Hamiltonian) $E = \sqrt{p^2c^2 + m^2c^4}$ reduces to $E = mc^2$. We often interpret the mass of a particle as being the energy required to create a “copy” of the particle at rest, and it is analogous to the band gap energy in semiconductors.

\textsuperscript{62} $Q\Psi = 0$ gives us a first-order differential equation which can be directly integrated to obtain this ground state wave function! This trick extends to field theory applications too—yet another way in which supersymmetry simplifies theorists lives.

\textsuperscript{63} The story starts with the Coleman-Mandula no-go theorem in 1967. (According to n-Lab, a no-go theorem is “any theorem...that shows that an idea is not possible even though it may appear as if it should be.” Thus Bell’s theorem is a no-go theorem dictating the impossibility of local, hidden variable theories that reproduce the predictions of quantum mechanics.) The Coleman-Mandula theorem tells us that in a realistic quantum field theory, space-time symmetries (like the Lorentz group) can only be combined with internal
translational symmetry (with an associated conserved momentum), a time-translational symmetry (associated with the conserved energy, with the Hamiltonian giving the infinitesimal time-translation operator), and other symmetries (rotations and relativistic boosts). Combining these symmetries gives us the Poincaré group.

In our “zero-dimensional” harmonic oscillator, only the time-translational symmetry remains from the Poincaré group. How does supersymmetry extend time-translation invariance? Can we somehow create a time translation by supersymmetrically transforming it?

(f) Show that \( H_S = \frac{1}{2} \{Q, Q^\dagger\} \). We see that a combination of two supercharges generates a time translation!

The supersymmetric harmonic oscillator we looked at above may seem pretty trivial: how hard is it to get degenerate states when all states have the same energy splitting? However, we can generate lots of interacting supersymmetric Hamiltonians by specifying a supercharge

\[
Q_W = b \left( \frac{p}{\sqrt{m}} + i\sqrt{m} W'(x) \right)
\]

where \( W'(x) = \frac{dW}{dx} \), and requiring that \( \mathcal{H}_W = \frac{1}{2} \{Q_W, Q_W^\dagger\} \), where the real function \( W(x) \) is called the superpotential.

Our Hamiltonian \( \mathcal{H}_S \) can be viewed as the special case of \( W(x) = \frac{1}{2} \omega x^2 \). Note that our superpotential need not have units of energy.

(g) Show that \( Q_W \) and \( \mathcal{H}_W \) are 2\times2 matrices

\[
\mathcal{H}_W = \begin{pmatrix} \mathcal{H}_1 & 0 \\ 0 & \mathcal{H}_2 \end{pmatrix}
\]

\[
Q_W = \begin{pmatrix} 0 & 0 \\ A & 0 \end{pmatrix}
\]

where the elements of the matrices are functions of \( p \) and \( x \). (Hint: Remember \( p = -i\hbar \partial/\partial x \). You might check this against the Web, which has different units.)
There is a lovely relationship between the eigenvalues and eigenfunctions of \( H_1 \) and \( H_2 \), two seemingly different Hamiltonians. Let \( \Psi_n^{(1)}(x) \) and \( \Psi_m^{(2)}(x) \) be the \( n \)-th and \( m \)-th eigenfunctions of \( H_1 \) and \( H_2 \), respectively.

(h) Using the fact that \([H_W, Q_W] = [H_W, Q^*_W] = 0\), show that \( A^\dagger \Psi_m^{(2)}(x) \) is an eigenstate of \( H_1 \) and \( A \Psi_n^{(1)}(x) \) is an eigenstate of \( H_2 \). (Thus, if we know the eigenfunctions and eigenenergies of one of the Hamiltonians, we know them for the other.)

Let us work out a specific example. Consider \( W'(x) = (\pi \hbar /mL) \cot(\pi x/L) \).

(i) Show that \( H_1 \) is the particle-in-a-box Hamiltonian (Fig. N4.15) shifted by a constant to set its ground state energy to zero. Show that \( H_2 \) is a Hamiltonian with potential

\[
V(x) = \frac{\pi^2 \hbar^2}{2 m L^2} \left( 2 \csc^2 \left( \frac{\pi x}{L} \right) - 1 \right).
\]

(N4.67)

Using the first excited state \( \Psi_2^{(1)}(x) = \sqrt{2/L} \sin(2\pi x/L) \) of \( H_1 \) and the operator \( A \), generate the ground state of \( H_2 \) and show that it is proportional to \( \sin^2(\pi x/L) \). Explicitly show (taking the derivatives) that \( A \Psi_2^{(1)}(x) \) is an eigenfunction of \( H_2 \) and thus verify that its energy is the same as that of \( \Psi_2^{(1)} \).

Fig. N4.15 Supersymmetric eigenenergies and eigenstates. (Left) Eigenstates for \( H_1 \), the square well potential, displaced vertically by their eigenenergies. (Right) Eigenstates for \( H_2 \), the \( \csc^2 x \) potential, which is the supersymmetric pair for the square well.

While supersymmetry may not exist in nature, it has proved to be an excellent tool for gaining insight into the way theories with gauge symmetry behave. (For example, we have no proof that the strong interaction confines quarks, but Seiberg and Witten were able to demonstrate confinement in certain supersymmetric theories.) It also has allowed physicists to prove theorems in pure mathematics. Ed Witten, high-energy theorist at the Institute for Advanced Study, was awarded the Fields Medal (the Nobel equivalent in math) for his use of supersymmetry to figure out topological properties of a manifold (such as the Euler characteristic, related to the number of holes or handles a manifold has) by using the difference in the number of zero-energy fermion and boson wavefunctions on it.\(^{65}\)

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\(^{64}\)Note that the \( \Psi = 0 \) boundary conditions for the two Hamiltonians are the same for both \( H_1 \) and \( H_2 \).

Fourier series and group representations. (Math)

In class, we focused on finite-dimensional group representations for finite groups. In quantum mechanics, the most useful symmetries are often continuous, and Hilbert space is infinite dimensional. With some small modifications, all of our results can go through to the continuous case.

Here we apply group representation theory to the continuous rotations in the plane, $SO(2)$. Let $g_\phi \in SO(2)$ be the rotation by angle $\phi$.

(a) Show that every different $g_\phi$ is in its own conjugacy class. (This is true for any commutative group.)

Thus we may label the conjugacy classes by the angle $\phi$.

Consider the action of $g_\phi$ on a function $f(\theta)$:

$$R(g_\phi) : f(\theta) \to f(\theta - \phi).$$  \hfill (N4.68)

Here $\theta$ represents a point on a circle, the complex function $f(\theta)$ is a vector in the Hilbert space of complex functions on the circle, and $R(g_\phi)$ is a linear mapping of that function space into itself.

(b) Show that, for any nonnegative integer $m$, that the two-dimensional space spanned by the basis $\{\cos(m\theta), \sin(m\theta)\}$ is an invariant subspace under $SO(2)$. Give the explicit $2 \times 2$ matrix for $R(g_\phi)$ acting on this subspace in this basis. What is the character $\chi(\phi)$ of this representation? (Hint: Use the angle addition formulas. Check that the character of the identity is the dimension of the representation.)

In the space of complex functions on the circle, this two-dimensional representation is not irreducible. It can be decomposed into two invariant subspaces, with bases $\{e^{im\theta}\}$ and $\{e^{-im\theta}\}$.

(c) What is the character of the $m$-representation given by the one-dimensional invariant subspace of multiples of $\{e^{im\theta}\}$?

Thus the character table for $SO(2)$ would have an infinite number of rows (one for each integer $\pm m$) and a continuous infinity of columns (one for each angle $\phi$).

For finite groups, we decomposed representations into irreducible pieces using the little orthogonality theorem: for any two irreducible representations $\alpha$ and $\beta$, the sum over group elements $\sum_{g \in G} \chi^{(\alpha)}(g)\chi^{(\beta)}(g)^{} = o(G)\delta_{\alpha\beta}$, where $o(G)$ is the number of elements of the group. For continuous groups, the sum must be replaced by an integral over

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66 This exercise is mostly about understanding the definitions. If you find resources on the Web or elsewhere that are helpful, just properly acknowledge them. In particular, I found http://www.cmth.ph.ic.ac.uk/people/d.vvedensky/groups/Chapter8.pdf which discusses the application of group reps to $SO(2)$. No guarantees that my conventions agree with those in the literature, though.

67 Particularly, $L^2$ functions on the circle.

68 In the past, we viewed group representations as mappings of the group into spaces of matrices that preserve multiplication. But matrices are just linear transformations of vectors; here we are using infinite dimensional vectors instead. Thus $R(g)$ is a linear map taking a function to another function.
the group, and the number of elements of the group replaced by the “volume” of the group. For two-dimensional rotations, we find
\[
\int_0^{2\pi} d\phi \chi^{(\alpha)}(\phi)\chi^{(\beta)}(\phi)^* = 2\pi\delta_{\alpha\beta}.
\] (N4.69)

(d) Show that the characters of your irreducible representations from part (c) satisfy the orthogonality relation N4.69. Is the character of your reducible representation in part (b) orthogonal to all the irreducible representations? Use the little orthogonality relation explicitly to decompose this reducible representation into its irreducible components.

For finite-dimensional representations of finite groups, we knew that any representation could be decomposed into irreducible representations: that is, any general vector could be written as a sum of vectors in the different invariant subspaces. For example, in Alemi’s analysis of vibrations in a triangular molecule, he found the normal modes by using a projection operator
\[
P^{(\alpha)} = \left(\frac{1}{\mathcal{O}(G)}\right) \sum_{g \in G} \chi^{(\alpha)}(g) R(g).
\] (N4.70)

Here \( f^{(\alpha)} \) is the dimension of the representation \( \alpha \).

For example, any random deformation of the molecule, when averaged over the group, gave a uniform dialation of the triangle. This dialation is invariant under triangular symmetries—so it transforms under the representation \( A_1 \). Since \( \chi^{(A_1)}(g) \equiv 1 \), this is just what eqn N4.70 suggests. When Alex multiplied by the characters of the two-dimensional representation \( E \) (using \( P^{(E)} \) in eqn N4.70), though, he discovered a different normal mode that was doubly degenerate.

Let us return now to our infinite-dimensional space of complex functions on the circle, to connect our irreducible representation decomposition with the theory of Fourier series. For our continuous group \( SO(2) \), the corresponding projection operator is
\[
P^{(\alpha)} = \left(\frac{1}{2\pi}\right) \int_0^{2\pi} d\phi \chi^{(\alpha)}(\phi) R(g_{\phi}).
\] (N4.71)

Let \( f(\theta) \) be a particular complex function on the circle. Let \( R(g_{\phi}) \) be defined on the function as in eqn N4.68.

(e) Show that the projection operator in eqn N4.71, using the \( m \) representation of part (c), takes \( f(\theta) \) into a coefficient times the basis vector for that representation. How is the coefficient related to the Fourier series coefficient for \( f \)?

\[\text{footnote}\text{For } SO(2), \text{this is just an integral over } \phi. \text{More generally, and in particular for } SO(3), \text{you have an extra factor in the integral (the Haar measure).}
\[\text{footnote}\text{Alex didn’t bother with the factor } \left(\frac{f^{(\alpha)}}{\mathcal{O}(G)}\right), \text{since he just wanted a vector in the subspace. We want to make the sum over representations } \alpha \text{ equal to the original function. Alex also, I think, missed the complex conjugate (but all his characters were real).}
\[\text{footnote}\text{There are many different conventions for Fourier series. Clearly state which one you are using.}]}
If we sum the projections of a vector into all the invariant subspaces, we should get the vector back again. The inverse Fourier series takes the Fourier coefficients and resums them back into the original function.

(f) How is the inverse Fourier series related to the sum of the projections of \( f(\theta) \) over all the irreducible representations of \( SO(2) \)?

N4.45 Solving Schrödinger: WKB, instantons, and the double well. (Computation) 3

We study the problem of quantum tunneling in a symmetric double well potential. If the barrier between the two wells is large, the ground state and first excited state are well approximated as symmetric and antisymmetric superpositions of the ground states in the two separated wells. Indeed, the low-energy physics of the double-well system can be approximated by a two-level system (TLS), another example of a Q-bit. In the symmetric case, in a basis where \( \begin{pmatrix} 1 \\ 0 \end{pmatrix} \) is the state localized in the left well and and \( \begin{pmatrix} 0 \\ 1 \end{pmatrix} \) is in the right well, the Hamiltonian is

\[
H_{\text{TLS}} = \begin{pmatrix} 0 & -\Delta \\ -\Delta & 0 \end{pmatrix},
\]

where \( \Delta \) is called the tunneling matrix element.

(a) What is the energy splitting for \( H_{\text{TLS}} \)? Calculate the time evolution operator \( U(t) = \exp(-\frac{iH_{\text{TLS}}t}{\hbar}) \). If we start in the left well at \( t = 0 \), what is the probability being in the left well after time \( t \)?

It is traditional and useful to study the double well given by the quartic polynomial

\[
V(y) = V_0 \left( \frac{y}{Q_0} - 1 \right)^2 \left( \frac{y}{Q_0} + 1 \right)^2.
\]

In our notation, this potential has minima at \( \pm Q_0 \) separated by a barrier of height \( V_0 \); a particle of mass \( m \) has small oscillation frequency \( \omega \) near the minima in each of the two wells.

(b) Calculate the barrier height \( V_0 \) algebraically in terms of \( m \), \( Q_0 \), and \( \omega \), that sets the small oscillation frequency near \( y = Q_0 \) equal to \( \omega \).

(c) Calculate the classical instanton action (or, equivalently, the WKB exponent) \( S_0 = \int_{-Q_0}^{Q_0} \sqrt{2mV(y)}dy \) for our potential. Let \( Q_0 = na_0 \) with \( a_0 = \sqrt{\hbar/2m\omega} \) the width of the ground state in the harmonic approximation. What is \( S_0/\hbar \) in terms of \( n \)? What numerical value does it have for \( n = 5 \)? How big is \( \exp(-S_0/\hbar) \), the Euclidean action’s suppression to the contribution of the instanton path? (Hint: \( S_0/\hbar \) is dimensionless. As suggested by the latter part of the exercise, for our potential it should depend only on the dimensionless number \( n \), given that \( V_0 \) is given as in part (b).)

We shall assume our particle has the mass of a hydrogen atom, the small oscillation frequency in the left and right wells is 1THz, and \( Q_0 = 5a_0 \). Generate a grid of length
Let the initial wavefunction $\psi[0] = \psi_0(x + Q_0)$ be the harmonic ground state in the left well. As in the coherent-state exercise, generate the appropriate arrays $U_{pot}(dt/2)$ and $U_{kin}(dt)$ to evolve $\psi$ using your BCH formula, with $dt = P/20$ where $P$ is the period of the oscillator.

(d) Solve for the time evolution over 1,000 periods. Animate the probability density $|\psi(x,t)|^2$ versus time. Does it travel between the two wells periodically, as should be predicted by your answer to part (a)?

(e) Plot the probability $\int_{-\infty}^{0} |\psi(x,t)|^2$ that the particle is found in the left well. (The high-frequency wiggles are due to the small components of higher eigenstates of our initial state in the left well.) By eye, estimate the period of oscillation, and also $\Delta$, within 10%.

(f) Use a nonlinear least-squares method to fit your prediction from part (a) to the probability you calculated in part (e). Include your fit in the plot for part (e). What is $\Delta_{\text{Fit}}$? (If your answer from part (e) is off by much more than 30%, see if you can track down the problem.) What is the ratio of your estimated energy splitting (using the formula from part (a)), to the energy splitting between states in one of the two wells (in the harmonic approximation)? Quantum tunneling is one of the most important sources of low-energy, long-time behavior in physics.

The instanton formula for the tunnel splitting is closely related to the WKB formula. Both are of the form $\Delta = \hbar \omega_0 \exp(-S_0/\hbar)$. In the WKB formula, the prefactor $\hbar \omega_0$ is given by a matching calculation; in the instanton method it is given by a path integral incorporating small oscillations about the instanton path. Gildener and Patrascioiu (Phys. Rev. D 16, 423, 1977, referred to by Coleman) did the calculation explicitly for the quartic well using instanton methods, and got

$$\Delta \sim \hbar \omega \sqrt{\frac{6S_0}{\pi \hbar}} \exp(-S_0/\hbar)$$

so in our notation $\omega_0/\omega = \sqrt{6S_0/\pi \hbar}$.

(g) Calculate your numerical estimate of $\omega_0/\omega$, and compare with Gildener and Patrascioiu’s estimate.

There is an interesting story about communication between high-energy and condensed-matter physics. Sidney Coleman (high-energy Harvard theorist) and Jim Langer (condensed matter theorist on sabbatical at Harvard) were both working on barrier crossing at the same time. Coleman was stuck for months, because his instanton calculation was a factor of two higher than the WKB estimate (the 'double well done doubly well')

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72 Nonlinear least-squares routines typically get lost if you don’t start near the best fit.
73 The two turning points for the classical path in the harmonic well are not far enough apart for the traditional WKB formula to be accurate; a suitable generalization accounting for that proximity does agree with the instanton formula.
74 Coleman tells us the answer from the instanton calculation is given in terms of Wronskians.
appendix to his Les Houches lectures, calculating quantum fluctuations causing transitions through barriers). Unbeknownst to him, Jim Langer in a nearby office had solved the problem in the statistical mechanics context (critical droplet theory, calculating thermal fluctuations causing transitions over barriers). Eventually, Coleman realized as Langer had earlier that only half the fluctuations that reach the barrier actually cross it.

\[ \text{GHZ and } \sigma \text{ matrices.} \]  
(Weirdness)  

The GHZ state is the state of three spins held in spatial isolation from one another by Alice, Bob, and Carol: \[ |GHZ\rangle = \left( \frac{1}{\sqrt{2}} \right) (|↑z, ↑z, ↑z\rangle - |↓z, ↓z, ↓z\rangle), \] where the first label in the ket represents the state of the A-spin, the second the B-spin, etc. In class, we showed that \( XXX = X_A X_B X_C \) acting on \( |GHZ\rangle \) always gave \(-1\), but gave \(+1\) for \( XYY = X_A Y_B Y_C \) or \( YXY \) or \( YYX \), each time they are measured. Multiplying the results of measurements of the latter three, \((X_A Y_B Y_C)(Y_A X_B Y_C)(Y_A Y_B X_C)\) gave plus one, and not minus one (as one would presume if Bell was right and hidden variables stored “true” states of \( X \) and \( Y \)).

Instead of considering products of observations, consider the operator product

\[ \Omega = (X_A Y_B Y_C)(Y_A X_B Y_C)(Y_A Y_B X_C) \]  \( \text{(N4.75)} \)

In this exercise you shall show that \( \Omega \) is equal to \(-X_A X_B X_C\) as an operator (independent of the GHZ state). There are two steps in the proof.

First, since Alice, Bob, and Carol are isolated from one another, Alice’s measurements do not affect Bob’s, etc. This means that any operator \((X, Y, \text{or } Z)\) at \( A \) should commute with any operator at \( B \) or \( C \), and that operators at \( B \) and \( C \) should commute. Also, since all the operators have eigenvalues \( \pm 1 \), their squares are the identity (e.g., \( Y_B^2 = 1 \)).

(a) Using only this fact, rewrite \( \Omega \) of eqn N4.75 as the product of five operators.

The operator \( Z \) at a site has eigenvalue \(+1\) if the spin is quantized up along the \( z \) axis, and \(-1\) if the spin is down. Hence \( Z \) acting on a spin at that site\(^75\) gives \( Z = \left( \begin{array}{cc} 1 & 0 \\ 0 & -1 \end{array} \right) \). Similarly, in the \( z \) basis \( X = \left( \begin{array}{cc} 0 & 1 \\ 1 & 0 \end{array} \right) \) and \( Y = \left( \begin{array}{cc} 0 & -i \\ i & 0 \end{array} \right) \).

(b) Use this and your answer to part (a) to show that \( \Omega = -X_A X_B X_C \).

The GHZ state is unusual in that it is an eigenstate of \( XXX, XYY, YXY, \text{ and } YYX \), so that measurements of \( |GHZ\rangle \) by these operators leave it unchanged (up to a sign). This allowed us to view the conundrum that separate measurements of the three operators were incompatible with Bell. But we have shown that any state \( \psi \) will share the property of the GHZ state that \( \Omega |\psi\rangle = -XXX |\psi\rangle \).

\(^75\)Thus the spin operator \( \sigma_z = (\hbar/2)Z \). Also, \( X, Y, \text{ or } Z \) acting on a spin at another site is the identity.
N4.47 **No cloning theorem.** (Quantum Computing) 3

Can one make an exact copy of an arbitrary, unknown quantum state $|\psi\rangle$, without changing the original state? Sadly, this is impossible.76

\[
\begin{align*}
|\Psi_A\rangle & \quad |\Psi_A\rangle \\
|\Psi_B\rangle & \quad |\Psi_B\rangle \\
|0\rangle_B & \quad |0\rangle_B
\end{align*}
\]

**Fig. N4.16 Hypothetical cloning machine.** Our machine takes an unentangled state $|\psi\rangle_A$, and copies it using a blank state $|0\rangle_B$. In the process, the machine changes state in some fashion that can depend on $\psi$.

Pretend it was possible. In particular, let us start with three subsystems (Fig. N4.16): $A$ (to be cloned, with coordinates $x$), $B$ (to store the clone, coordinates $y$), and $M$ (the quantum cloning machine, with coordinates $Z$). Suppose subsystem $A$ is in state $|\psi\rangle_A = \psi(x)$; system $B$ starts in a blank system $|0\rangle_B = \beta(y)$, and our cloning machine starts in a resting state $|M_0\rangle_M = \mathcal{M}_0(Z)$. Before and after the machine operates, $A$, $B$, and $M$ are independent, uncoupled systems. The machine should act as a unitary operator $U_c$, taking $\Psi(x,y,Z) = \psi(x)\beta(y)\mathcal{M}_0(Z)$ into

\[
U_c|\Psi\rangle = \psi(x)\beta(y)\mathcal{M}_\psi(Z)
\]

where $\mathcal{M}_\psi(Z)$ is the final state of the machine after it processes the state $\Psi$. Our machine need only copy states $\psi(x)$ which are unentangled with the rest of the universe, and thus we expect the final state $\beta(y)$ should be also unentangled by our hypothetical machine.

(a) **Viewing the state in eqn N4.76 as a system $B$ in an environment made up of $A$ and $M$, is the quantum state entangled, or not? If not entangled, write it as a product; if entangled, give its Schmidt decomposition.** (Note: In writing the Schmidt decomposition $\sum_k \sigma_k u_k(y) v_k(x,Z)$, you must show $\sigma_k > 0$, $\sum_k \sigma_k^2 = 1$, the $|u_k\rangle_B$ are orthonormal, and that the $v_k(y,Z)$ are orthonormal. Hint: You can probably do this part by inspection.) If you do a Schmidt decomposition, explicitly check the orthogonality of the $AM$ states.

Suppose our initial cloning machine acts on two *orthonormal* input states $\psi(x)$ and $\phi(x)$, so in addition to eqn N4.76, we have

\[
U_c|\Psi\rangle = \phi(x)\phi(y)\mathcal{M}_\phi(Z)
\]

As with many impossible things, this would be very useful! Copying Qbits being used for secure quantum communication would allow one to listen in. Copying Mr. Spock while beaming up would have allowed exciting new episodes in the 1970’s . . .
(b) Let $U_c$ act on the superposition $\frac{1}{\sqrt{2}}(|\Psi\rangle + |\Phi\rangle)$; expand it using linearity and eqns N4.76 and N4.77. Is the final state entangled, or not? Again, write it as a product, or give its Schmidt decomposition. (Remember $\psi(x)$ and $\phi(x)$ are orthogonal. You may assume $|M_{\phi}\rangle$ and $|M_{\psi}\rangle$ are normalized after the unitary transformation, but they need not be orthogonal.) If you do a Schmidt decomposition, explicitly check the orthogonality of the AM states.

Thus our cloning machine must necessarily be unsuccessful in generating an unentangled clone of $\psi$, $\phi$, and their superposition. No general-purpose cloning operator is possible over a Hilbert space with more than one dimension.

N4.48 **Aharonov-Bohm wire: straightforward approach.** (Parallel Transport) ③

**Fig. N4.17 Aharonov-Bohm wire loop.** A circular wire of radius $R$ encloses a magnetic field of total flux $\Phi$.

An electron is confined to a one-dimensional wire loop of radius $R$ (Fig. N4.17). A uniform magnetic field of strength $B$ perpendicular to the loop has a net magnetic flux $\Phi = \pi R^2 B$. We pick a gauge in which the electromagnetic potential $A = A\theta = \Phi\theta/(2\pi R)$ at the radius of the wire, where $\theta$ is the unit vector tangent to the wire. Let $s$ measure distance around the wire. Schrödinger’s time-independent equation then becomes

$$E\psi(s) = -\frac{\hbar^2}{2m} \left( \frac{\partial}{\partial s} - \frac{i e A}{\hbar c} \right)^2 \psi(s)$$

$$E\psi(s) = -\frac{\hbar^2}{2m} \left( \frac{\partial}{\partial s} - \frac{i e \Phi}{2\pi R \hbar c} \right)^2 \psi(s)$$

where $e < 0$ is the charge on the electron. The energy eigenstates are of the form $\psi_n(s) \propto \exp(ik_n s)$. In this problem, we are not changing to a singular gauge.

(a) **What values of $k_n$ are allowed?** (Hints: The wavefunction must be periodic in arclength $s$ with period $2\pi R$. The answer does not depend on $\Phi$.)
(b) Show that $\psi_n$ is an eigenstate of $\left(\partial/\partial s - ieA/\hbar c\right)$. Using that, find the eigenenergies $E_n$. Are the eigenenergies of the system the same for $\Phi = 0$ and for $\Phi = \Phi_0 = hc/e = 2\pi hc/e$, as we showed in lecture? (Hint: The label $n$ of the energies may change, but is the spectrum of eigenenergies unchanged?)

The flux through the loop is set to exactly half a flux quantum, $\Phi = \Phi_0/2 = hc/2e = \pi hc/e$.

(c) What eigenstate or eigenstates have lowest energy? Are all the eigenstates doubly degenerate? (Show your work.)

N4.49 Number, Phase, and Josephson. (Quantum commutation) 3

In this exercise, we use the Josephson effect in superconductors to explore how the phase of the superconducting wavefunction is related to the indeterminacy in the number of particles.

A Josephson junction is a thin insulating layer or a weak link joining two regions of superconducting material or superfluid, see Fig. N4.18(a). Let the number of bosons (charge $2e$ Cooper pairs) in the two pieces be $N_1$ and $N_2$, and the superconducting phases be $\phi_1$ and $\phi_2$. There is a term in the energy $U(\phi_2 - \phi_1)$ that grows with the phase difference between the two sides. Naturally, this energy cost must be periodic in $\delta\phi = \phi_2 - \phi_1$. For weak links, this energy cost is very nearly some constant $U_0$ times $1 - \cos(\delta\phi)$,

$$U(\delta\phi) = U_0(1 - \cos(\delta\phi)),$$

as plotted in Fig. N4.18(b).

![Fig. N4.18 Josephson junction schematic.](image)

(a) A Josephson junction is made from a weak link (light small box) connecting two pieces of superconductor or superfluid. (b) The energy of the superconducting weak link as a function of the phase difference across it.

In class, we remarked that number and phase for superconductors are conjugate variables, $[\phi, N] = i$, similar to how position and momentum are conjugate $[x, p] = i\hbar$. We said this could be used to explain the behavior of Josephson junctions. Here we will work backward. We shall start with the experimental behavior of the junction (as predicted by Josephson from a more microscopic calculation), and deduce the commutation relation.

First, a Josephson junction experimentally has a supercurrent $I/(2e) = dN_2/dt = -dN_1/dt$ which depends on $\delta\phi$:

$$I = I_c \sin(\delta\phi).$$
This is the DC Josephson effect; electron pairs can tunnel across the insulating gap, but only up to a critical current $I_c$. Let $\delta N = (N_2 - N_1)/2$, the net number of 2e bosons that have crossed the junction. Note from eqn N4.78 that $d(\delta N)/dt = I (\delta \phi)/(2e) \propto -dU/d\delta \phi$, just as the rate of change of momentum $\dot{p}$ is given by the force (the negative derivative of the potential energy). We can use this to deduce the commutation of $\delta \phi$ and $\delta N$ up to an overall constant $x$.

(a) Use the general Heisenberg relation $dO/dt = (1/i\hbar)[O, H]$, and assume that $H$ is some (so far) unknown function of $\delta N$ (the “kinetic energy”) plus $U(\delta \phi)$ (the “potential energy”). Show that $\delta N = (x/i)\partial/\partial (\delta \phi)$ gives a DC Josephson relation for any value of $x$. Use this to write $[\delta \phi, \delta N]$ in terms of $x$. Write $x$ in terms of $I_c$ and $U_0$.

(b) Using your commutator $[\delta \phi, \delta N]$ from part (a), deduce that $\delta \phi = (y/i)\partial/\partial \delta N$ by showing that $[\delta \phi, \delta N^m] = (y/i)m\delta N^{m-1}$ for all positive integers $m$. (Hint: work by induction.) Write $y$ in terms of $x$.

Second, with a fixed voltage $V$ across a Josephson junction, we measure an AC current $I(t) = I_c \sin((2e/\hbar)Vt)$. This is the AC Josephson effect, and has been used in the past to define the volt. Using the DC Josephson relation (eqn N4.79), this implies $d(\delta \phi)/dt = 2eV/\hbar$. Note that $2eV$ is the energy to move one Cooper pair from the left to the right. For most junctions, a change of $\delta N = 1$ Cooper pair can be viewed as an infinitesimal change in $\delta N$. We can use this to deduce the constant $x$.

(c) Use $d(\delta \phi)/dt = (1/i\hbar)[\delta \phi, H]$ and the form of $\delta \phi$ from part (b), deduce $y$, $x$, and the relation between $U_0$ and $I_c$. In particular, verify that the commutator $[\delta \phi, \delta N] = i$, as suggested in class. (Hint: How is the voltage related to the energy, and thus the Hamiltonian?)

While the superconducting wavefunction is a macroscopic manifestation of quantum mechanics, it usually is not subject to large quantum fluctuations. Instead, it acts like a magnetization or a crystalline orientation—a macroscopic property of a material, that has a well-defined value. When will our Hamiltonian for the Josephson junction, which describes the classical evolution of the current and phase, have important quantum fluctuations?

Just as $[x,p] = i\hbar$ implies $\sigma_x \sigma_p \geq \hbar/2$, $[\delta \phi, \delta N] = i$ implies $\sigma_{\delta \phi} \sigma_{\delta N} \geq \hbar/2$. Under normal circumstances, the number of Cooper pairs $\delta N$ exchanged between the superconductors is enormous. In many ways, the superconducting condensate acts like a classical field. But for a tiny junction, the total number of Cooper pairs can be small—leading to a noticeable quantum fluctuation in the superconducting phase. A group at IBM tested ideas of Tony Leggett by building such a Josephson junction (Washburn et al., “Effects of dissipation and temperature on macroscopic quantum tunneling”, PRL 54, 25 (1985)).

Let the thin insulating layer in Fig. N4.18(a) be of thickness $t = 10$nm, and width and height $L$. Treat the Josephson junction as a capacitor, with a charge $Q = 2e\delta N$ transferred from one side to another. This gives us a model for the “kinetic energy” in $\mathcal{H}$. For many purposes, the complicated physics of a Josephson junction can be
summarized by the critical current $I_c$, the capacitance $C$, and a resistance (not discussed here).

(d) Including the capacitor energy as the kinetic energy, give the total Hamiltonian $\mathcal{H}$ for our junction, in terms of the junction capacitance $C$ and the critical current $I_c$. Take the harmonic approximation $U(\delta \phi) \approx \frac{1}{2} U_0 \delta \phi^2$, and derive the capacitance necessary to make $\langle \delta \phi^2 \rangle = 1$ in the ground state, in terms of the critical current $I_c$. Washburn et al. used a junction with a capacitance of $C = 0.02 \pm 0.005 \text{pF}$, and a critical current of $I_c = 55 \mu \text{A}$. How big is $\delta \phi$ for their junction?

Many experiments and theoretical calculations have been done on these quantum fluctuations. A superconducting loop with a tiny Josephson junction weak link, when put in a magnetic field, will allow flux to quantum tunnel out of the junction. This quantum tunneling can be viewed as quantum fluctuations allowing the phase $\delta \phi$ to tunnel from one minimum of $U(\delta \phi)$ to the next.

N4.50 Localization.⁷⁷ (Quantum, Condensed Matter) ³

In Section 7.4, we discussed how non-interacting electrons provide a useful model for metals, even though the electron-electron interactions are strong. The Fermi liquid of quasiparticles is a kind of adiabatic continuation of the noninteracting electron system, connected by perturbation theory. Here we shall study how a one-dimensional non-interacting metal responds to disorder. We shall discuss how metals with weak disorder are understood by perturbing around the clean state. We shall discover that strong disorder leads to an insulating system whose eigenstates are not extended, but localized. We will describe these localized states explicitly by perturbing about a state of isolated atomic states.

Consider a one-dimensional chain of atoms $n$, each with one noninteracting electron state $|n\rangle$ of energy $U_n$, that can be occupied by either zero or one spinless electrons. Electrons can hop between atoms with matrix element $t$, leading to the Hamiltonian

$$\mathcal{H} = \sum_{n=1}^{N} U_n |n\rangle \langle n|$$

(N4.80)

$$-t \sum_{n=1}^{N-1} (|n\rangle \langle n+1| + |n+1\rangle \langle n|)$$

$$= \begin{pmatrix}
U_1 & -t & 0 & \ldots & 0 & 0 \\
-t & U_2 & -t & \ldots & 0 & 0 \\
0 & -t & U_3 & \ldots & 0 & 0 \\
0 & 0 & -t & \ldots & 0 & 0 \\
\vdots & \vdots & \vdots & \ddots & \ddots & \ddots \\
0 & 0 & 0 & \ldots & -t & U_N
\end{pmatrix}$$

⁷⁷Hints for the computations can be found at the book web site [31].
We shall take the random energies $U_n$ as uniformly distributed between $-W$ and $W$. Without disorder ($W = 0$), this is a textbook model used to describe energy bands in crystals. Three dimensional analogs of this ‘tight-binding’ model are quite realistic models of Fermi surfaces and energy bands in real materials.\(^{78}\)

(a) Write a function that builds the Hamiltonian matrix of eqn N4.80 with size $N$, bandwidth $2W$, and hopping matrix element $t$. Studying zero disorder $W = 0$, find the eigenvectors for $t = 1$, and $N = 100$, sorted by their eigenvalues. Plot the eigenvectors for the four lowest energies. Check numerically that these four are sinusoidal with wavevectors $k_\alpha = \pi \alpha/(N + 1)$ appropriate for a box of size $N$ with hard-wall boundary conditions half a grid spacing to either side. Check that their four eigenvalues are the corresponding $E_{k_\alpha} = -2t \cos(k_\alpha)$

Imagine a 1D metal at zero temperatures with electrons filling the states up to a Fermi surface, here just two points at some $\pm k_{\text{Fermi}}$. Consider a packet of electrons made up of eigenstates near $k_{\text{Fermi}}$ traveling to the right. The wavepacket\(^{79}\) will travel, as usual, at the group velocity $dE_k/dk|_{k_{\text{Fermi}}}$, without dissipation.

Now let us explore what happens when we add a weak disorder.

(b) Build a Hamiltonian with weak disorder $W = W_{\text{weak}} = 0.04$, $t = 1$, and $N = 100$. Plot the lowest four eigenvectors. Are the eigenstates still extensive (reaching from one side of the box to the other)?

At this point, we could use perturbation theory to calculate the disordered eigenstates and energy levels. We could then create wavepackets and see how they evolve. In three dimensions, the scattering off of the disorder changes the electron transport qualitatively. Instead of wavepackets moving forever in one direction (ballistic transport, infinite conductivity), one gets diffusive motion of the electron probability through space (disorder providing an elastic scattering length, and a finite conductivity). In three dimensions, this is a good model for metals with impurities or dopants, illustrating how one can understand complex behavior by perturbing around solvable special cases.

Instead, let us examine what happens at large disorder $W$, or equivalently, small hopping $t$. (All of our eigenvectors depend only on $W/t$, and we will perturb in $t$ to study the localized states.)

(c) Set $t = t_{\text{weak}} = 0.1$, $W = 1$, and $N = 100$, plotting the ten eigenvectors with lowest energies. Also do a log-linear plot of the probability density (absolute square of

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\(^{78}\)We shall see that even a small disorder changes the metallic behavior of one-dimensional electrons in a qualitative and interesting way. Indeed, one dimensional electrons are unstable in many interesting ways. Adding interactions between electrons, they become \textit{Luttinger liquids}, with emergent scale invariance. Adding interactions with lattice vibrations, they can become topological insulators, with solitons and fractional charges.

\(^{79}\)Wavepackets are used to connect waves to particle-like motion. In a non-disordered system, one superimposes states with similar momenta to make a spatially localized wavefunction, which then moves with the group velocity of the wave. We discuss wavepackets to motivate the effects of disorder, but no knowledge about them is required to do this exercise.
the wavefunctions) for these eigenvectors. Do the eigenstates still look as if they will be extensive (stretching from one end of a macroscopic wire to the other)? What solvable special limit of $\mathcal{H}$ should we use to capture this new behavior?

Here we find the eigenstates appear localized – fixed in space near individual ‘atoms’. The probabilities in these states fall exponentially with distance from their centers. A wavepacket formed from localized states like these cannot transport current: for large disorder, our model describes an insulator.

Just as one can use perturbation theory to describe dirty metals in three dimensions from models like ours, we can use perturbation theory to calculate and understand these localized states. You should remember the use of second-order perturbation theory to describe the energies of a Hamiltonian $\mathcal{H} = \mathcal{H}_0 + V$ for small $V$. You may not remember that the first step was to use first-order perturbation theory to determine the eigenvectors. If $|\Psi_i^{(0)}\rangle$ has unperturbed eigenvalue $E_i^{(0)}$, then to first order

$$|\Psi_i^{(1)}\rangle = |\Psi_i^{(0)}\rangle + \sum_{i \neq j} \frac{\langle \Psi_j^{(0)} | V | \Psi_i^{(0)} \rangle}{E_i^{(0)} - E_j^{(0)}} |\Psi_j^{(0)}\rangle.$$  \hspace{1cm} (N4.81)

If the hopping is small compared to the disorder, let us perturb in $t$.

(d) **What are the eigenenergies for $\mathcal{H}$ in eqn N4.80 with $t = 0$?** Argue that, to first order in $t$, the new eigenstates will be confined to three adjacent sites.

(e) **Write a function, given $\mathcal{H}$, $i$, $t$, and $N$, that gives the perturbed eigenstate to lowest order in $t$ that is centered at site $i$.** Find the site of the ground state with largest probability. Plot the true ground state and your first-order approximation to it. (Hint: You may be unlucky, and happen to have a neighbor site with a near degenerate energy. Just create a new Hamiltonian and try again.)

What controls whether our model is a metal or an insulator? For a given $W/t$, are all the states either extended or all localized? Or could there be some mixture?

We can examine this by defining a rough measure of how spread out the wavefunction is, called the **participation ratio**:

$$P(\psi) = \frac{\left(\sum_n |\psi(n)|^2\right)^2}{\sum_n |\psi(n)|^4} = \frac{1}{\sum_n |\psi(n)|^4}. \hspace{1cm} (N4.82)$$

(f) **Show that a state whose probability is spread uniformly among $M$ sites has $P = M$.** At zero disorder, what is the participation ratio for the lowest energy state? For the long-wavelength next few states? What is the ratio for a localized state that decays exponentially, $\psi(j) \propto \exp(-|i - j|/\lambda)$, in an infinite chain, with $\lambda$ much larger than one?

If the participation ratio $P \ll N$ we can reasonably expect that the eigenstate is localized.
(g) Calculate the participation ratio for all the eigenstates for intermediate disorder $W_{\text{inter}} = 0.5$, $t = 1$, $N = 100$, and plot them against the energy. Is there a systematic variation? Plot the wavefunction for an energy in the middle of the energy band (eigenvalue $E$ near zero), and one at the top and bottom of the band. Which are less localized – the states near the edges of the band, or the states in the center?

In experiments, one finds a region of localized states at the edges of a band, and extended states in the middle of the band. Between these is a mobility edge, where a metal-insulator transition occurs as more electrons are added.

Finally, can we find a mobility edge for our model? One thing to check is if the wavefunctions might have decay lengths larger than our system (so they just look extended at $N = 100$).

(h) Find the eigenvectors and eigenvalues for the same parameters as in part (g), $W = W_{\text{inter}} = 0.5$ and $t = 1$, except for a much larger system ($N = 2000$ if it is feasible on your system). Plot the participation ratio verses energy, and plot eigenstate in the center of the band and at the two edges. Do the states in the middle of the band now appear localized? Are their participation ratios larger than 100 – the system size in part (g)? Does it make sense that they looked extended in the smaller system, but clearly in an infinite system are localized?

As it happens, disordered electrons in one dimension are always localized, even for tiny disorder. The spinless, noninteracting electrons we study here are also always localized in two dimensions. In two dimensions, they can become extended when interactions, spin orbit scattering, or strong magnetic fields are added. In particular, 2D electrons in a strong magnetic field exhibit the quantum Hall effect (with extended states around the edges). Even more interesting, interacting electrons in a strong magnetic field exhibit the fractional quantum Hall effect – our first example of an experimental system with fractional charges and fractional statistics.

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80 Each time we add one order to perturbation theory, we get a wavefunction extending outward by one atom. It appears from the first two terms that each order multiplies the terminal amplitude by a factor $|\psi(i + n)/\psi(i + (n - 1))| = |t/(U_i - U_{i+n})|$. Roughly speaking, if $U_i$ is at the edge of the band, this factor is twice as small as if $U_i$ is in the center, so there is less localization at the center. Notice, though, that this is useful only when $t > |U_i - U_{i+n}|$. Rare, nearly degenerate states can mix strongly, even at long distances, making the arguments subtle.
N4.51 **Eight-Fold Way.** (Particles, Group reps) ③

**Fig. N4.19 Baryon octet.** (a) Shown are a family of eight baryons, headed by the neutron and proton, organized by their electrical charge \( Q \), their *strangeness* \( S \), and their isospin \( I_3 \).\(^81\) Thus for example the neutron has isospin \(-\frac{1}{2}\), and \( \Sigma^- \) has isospin \(-1\). These baryons transform under an irreducible representation \( 8 \) of the group \( SU(3) \). The charge \( Q \) is a linear combination of \( I_3 \) and strangeness. (b) The same baryons, plotting mass versus isospin \( I_3 \).

Years before the realization that the strong interaction has an exact “color” gauge symmetry group \( SU(3) \), it was noticed that it had an approximate “flavor” \( SU(3) \) symmetry. This was noticed because the baryons and mesons naturally organized themselves into multiplets which mimicked representations of \( SU(3) \) (Fig. N4.19(a)). Figure N4.19(b) shows the masses of one family of baryons, indicating the original experimental evidence for the model.

In this exercise,\(^82\) we will explore the geometry shown in Fig. N4.19(a), exploring the rows, the angles, and their relation to the symmetry groups.

The story began with the neutron and the proton. They are amazingly similar, apart from their charge. (For example, the proton has a mass 0.9986 times that of the neutron—only about 2\( \frac{1}{2} \) electron masses different. A little lighter, and the neutron could not decay.) Werner Heisenberg made an analogy with spin-\( \frac{1}{2} \) particles—the proton has isospin \( I_3 = +\frac{1}{2} \) and the neutron isospin \( I_3 = -\frac{1}{2} \).

In a world with only the strong interaction, and where the up and down quark had the same mass, the strong interaction would have an \( SU(2) \) total isospin symmetry as well as an \( SU(2) \) rotational symmetry—thus an overall symmetry \( SU(2) \times SU(2) \). The nucleon ground state, with spin \( \frac{1}{2} \) and isospin \( \frac{1}{2} \), thus would have four degenerate states \((n^\uparrow, n^\downarrow, p^\uparrow, \text{and } p^\downarrow)\).\(^83\)

(a) Draw a figure corresponding to Fig. N4.19(a) for these four nucleon states, plotting \( I_3 \) horizontally but plotting spin as the vertical axis.

Consider now the deuteron—the bound state of a proton and a neutron—in a world with perfect isospin symmetry. In such a world, we can think of the bound states of

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\(^81\) Reading: I found K. Schulten, “Notes on Quantum Mechanics”, ch. 12 to be quite useful in developing this exercise ([http://www.ks.uiuc.edu/Services/Class/PHYS480/qm_PDF/chp12.pdf](http://www.ks.uiuc.edu/Services/Class/PHYS480/qm_PDF/chp12.pdf)).

\(^82\) It is traditional to ignore the rotational \( SU(2) \) symmetry when discussing gauge field theories, since it is an *external* symmetry inherited from the symmetry of space-time, rather than an *internal* symmetry of the particle fields. We break with tradition in part (a).
two identical nucleons, which can have two (ordinary) rotational spin states \( \pm \frac{1}{2} \) and two isospin states \( I_3 = \pm \frac{1}{2} \) (proton or neutron). We may assume that the ground state spatial wavefunction for the deuteron is symmetric under interchange of the two nucleons, with orbital angular momentum zero.\(^8\) Two spin \( \frac{1}{2} \) nucleons may thus form rotational spin either zero or one \( (\frac{1}{2} \otimes \frac{1}{2} = 1 \oplus 0) \). Their total isospins combine in the same way. That is, isospin has the same symmetry group structure, so we know that the eigenstates of our isospin-symmetric deuteron will also be eigenstates of isospin with total \( I = 1 \) or \( I = 0 \).

(b) First, write the three total isospin triplet states in terms of \(|p\rangle\) and \(|n\rangle\). Are there bound states with \( I_3 = \pm 1 \), to your knowledge? (See section 12.2 of Schulten if you get stuck.) Second, how does the total isospin singlet state transform under interchange of the two nucleons? Treating the two nucleons as identical fermions (differing by their isospin \( I_3 \)), use the antisymmetry of the wavefunction to argue that the rotational spin of the deuteron must be equal to one.

Isospin is indeed so close to a symmetry of the strong interaction\(^8\) that one can use isospin Clebsch-Gordon coefficients to predict branching ratios in particle collisions.

We now turn to a much more strongly broken symmetry. The strange quark is notably heavier than the up and down quarks. But if we extend the SU(2) total isospin symmetry to SU(3), incorporating the strange quark, we can rationalize the elementary particles into families organized by their isospin, strangeness, and charge. Figure N4.19(a) shows eight particles which transform under the irreducible representation of SU(3) called \( 8 \).

You remember that angular momentum has \( J_x, J_y, \) and \( J_z \), and that the ladder operators \( J_{\pm} = J_x \pm i J_y \) were useful in deducing facts about the different spin states. In particular, the fact that \([J_z, J_{\pm}] = \pm \hbar J_{\pm}\) told us that, given an eigenstate \(|j_z\rangle\) of \( J_z \), we can find other eigenstates \(|j_{\pm} \pm 1\rangle\) of \( J_\pm \) (raising and lowering the rung of the \( j_z \) ladder).\(^8\) Other calculations told us how the ladders ended, and gave us the rules that the states with total angular momentum \( J^2 = j(j + 1) \) came as multiplets of degeneracy \( 2j + 1 \), with \( j_z = (-j, \ldots, j) \). We will focus here just on generalizing these ladder operators to SU(3).

The three quarks

\[
|u\rangle = \begin{pmatrix} 1 \\ 0 \end{pmatrix}, \quad |d\rangle = \begin{pmatrix} 0 \\ 1 \end{pmatrix}, \quad |s\rangle = \begin{pmatrix} 0 \\ 0 \end{pmatrix}
\]

(N4.83)

form a three-dimensional representation of SU(3) by multiplying by the corresponding matrix. (Here the language gets confusing, since SU(3) denotes both the abstract

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\(^8\)At high energies, wavefunctions become less quantitatively useful. The internal state of a nucleus is actually a mixture of states with different numbers of particles. Also, for the deuteron, spin-orbit interactions lead to a small mixing of states with orbital angular momentum \( L = 2 \). We ignore such subtle effects.

\(^8\)It is of course not a symmetry of the electromagnetic interaction: the proton and neutron have different charges! It is also not a symmetry of the weak interaction: neutrons weakly decay into protons emitting isospin-free electrons and neutrinos.

\(^8\)As a quick review, this works because \( J_z |j_z \pm 1\rangle = J_z J_{\pm} |j_z\rangle = (J_{\pm} J_z + [J_z, J_{\pm}]) |j_z\rangle = J_{\pm} h j_z |j_z\rangle \pm h J_{\pm} |j_z\rangle = h(j_z \pm 1) J_\pm |j_z\rangle = h(j_z \pm 1) |j_z \pm 1\rangle \).
group and its traditional representation as a set of $3 \times 3$ complex unitary matrices that all have determinant one.) We call this representation $\mathbf{3}$, just as we call the octet representation $\mathbf{8}$.

The total isospin symmetry $SU(2)$ is the subgroup of $SU(3)$ which just mixes the first two components $u$ and $d$. In the $\mathbf{3}$ representation, it has three infinitesimal generators which can be turned into the isospin operator $I_3 = \begin{pmatrix} \frac{1}{2} & 0 & 0 \\ 0 & -\frac{1}{2} & 0 \\ 0 & 0 & 0 \end{pmatrix}$ and the two ladder operators $I_+ = \begin{pmatrix} 0 & 1 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{pmatrix}$ and $I_- = \begin{pmatrix} 0 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 0 \end{pmatrix}$. Just as for spin, these ladder operators exist in all representations. For example, in the octet representation $\mathbf{8}$ of Fig. N4.19(a), $I_\pm$ shifts one right and left, so $I_+|\Xi^-\rangle \propto |\Xi^0\rangle$ and $I_-|p^+\rangle \propto |n^0\rangle$. Similarly, we can use the generators of the $SU(2)$ subgroup that mixes the up and strange quarks to form ladder operators $V_+ = \begin{pmatrix} 0 & 0 & 1 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{pmatrix}$ and $V_- = \begin{pmatrix} 0 & 0 & 0 \\ 0 & 0 & 0 \\ 1 & 0 & 0 \end{pmatrix}$, shifting up and down along the axis in Fig. N4.19(a) up-and-to-the-right at roughly $60^\circ$. The subgroup mixing down and strange forms a third pair of ladders $U_\pm$ moving at roughly $-60^\circ$.

(c) Show that $[I_3, V_+] = \frac{1}{2} V_+$, using the representation $\mathbf{3}$ for the generators above.

The three-fold symmetry of the raising and lowering operators gives us the three-fold symmetry of the baryon octet of Fig. N4.19(a).

(d) If $V_+|\Sigma^-\rangle = \alpha|n^0\rangle$ for some constant $\alpha$, calculate $I_3 V_+|\Sigma^-\rangle$ and $V_+ I_3|\Sigma^-\rangle$. (Note that the isospin operator in $I_3 V_+|\Sigma^-\rangle$ evaluates the isospin of a neutron, while in $V_+ I_3|\Sigma^-\rangle$ it evaluates the isospin of the $\Sigma^-$. ) Check against your answer from part (c).

Let us finally consider the fact that this $SU(3)$ flavor symmetry is not exact. In particular, the strange quark is quite heavy: note that the masses in Fig. N4.19(b) are mostly proportional to strangeness. Adding a big mass to the strange quark breaks the $SU(3)$ symmetry, but retains the isospin symmetry $SU(2) \subset SU(3)$. By decomposing the various $SU(3)$ representations into irreducible representations of $SU(2)$, we can understand another feature of the octets.

(e) In the three-dimensional representation $\mathbf{3}$ used for the three quarks above, what are the two invariant subspaces$^{88}$ under isospin $SU(2)$, written in terms of the three quark directions in eqn N4.83? (Hint: Which quarks are mixed by the isospin symmetry group? This is not subtle.) Show that$^{89}$ the $SU(3)$ representation $\mathbf{3}$ decomposes into

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$^{87}$ To make the diagram in Fig. N4.19(a) reflect the permutation symmetry between the three quarks of $SU(3)$, we really should squash the vertical axis a bit, so that the horizontal distance between neutron and proton along the isospin direction $I_3$ equals the distance between $\Sigma^-$ and $n^0$ up-and-to-the-right. Because we plot strangeness $S$ along the vertical axis rather than the more symmetrical $\sqrt{3}S/2$, moving along $V_+$ actually is at $63.43^\circ$.

$^{88}$Reminder: a subspace of a 3D vector space is a line or plane through the origin—spanned by one or two vectors. Fundamental to group representation theory is the decomposition of representations into irreducible representations. Decomposing the representation involves finding subspaces which are invariant—mapped to themselves—under the group.

$^{89}$Do not be confused by the notation. It is traditional in SU(3) to label the representation by the number of dimensions of the corresponding vector space in **boldface**—hence $\mathbf{8}$ has eight dimensions. It is traditional
isospin SU(2) representations $0 \oplus \frac{1}{2}$. (Hint: you don’t need character tables—the different isospin representations all have different dimensions.)

![Fig. N4.20 Meson octet. The light mesons also share the same group representation 8 of SU(3). (Note that the strangeness here is zero for the pions. Note that the masses for these mesons is not linear in strangeness—it grows for both positive and negative strangeness away from zero. Note finally that the strangeness arrow is reversed both here and in Fig. N4.19(a) from the standard convention, which would have the neutron and proton have highest (least negative) strangeness of their family of baryons.)](image)

We now want to build particles out of quarks. Adding three quarks to make a baryon leads to several representations. Let us instead add a quark and an antiquark to make a meson (Fig. N4.20). The antiquarks transform under a related representation $\bar{3}$, which also decomposes into $0 \oplus \frac{1}{2}$ when SU(3) is broken to SU(2). A quark and an antiquark together form a nine-dimensional representation $3 \otimes \bar{3} = 8 \oplus 1$, where 1 is a one-dimensional representation corresponding in this case to the $\eta'$ meson.

We need not refer to character orthogonality relations to find the isospin decomposition of 8. Given that we know how the two three dimensional SU(3) reps 3 and $\bar{3}$ decompose when broken to SU(2), we can deduce the decomposition of 8. The direct sum of the decomposition of the nine-dimensional product rep $3 \otimes 3 = 8 \oplus 1$. Since the one-dimensional rep 1 must lead to a one-dimensional isospin-zero piece, the isospin subspaces of 8 then can be deduced by elimination.\(^{90}\)

\(^{90}\)Again, the notation is confusing: we use the same symbols for direct sums and products of vectors, vector spaces, and group representations. Let us give a review and summary, in case it is more helpful than it is distracting. The direct product of two vectors $a_\alpha$ and $c_\gamma$ of dimensions $d_A$ and $d_C$ is the $d_A d_C$-dimensional vector with components $(a \otimes c)_{\alpha \gamma} = a_\alpha c_\gamma$; thus a spin-up electron in a $p_x$ state $\chi(x)$ and a spin-up muon in a $d_{xy}$ state $\phi(y)$ make for a direct product wavefunction $\Psi(x, y) = \chi(x) \phi(y)$. The direct product of two vector spaces $A$ and $C$ is the space $A \times C$ spanned by all linear combinations of vectors in the two spaces. Thus a spin-up electron with $L = 1$ and a spin-up muon with $L = 2$ will have a Hilbert space of fifteen dimensions. The direct sum of $a$ with $b$ of dimension $d_B$ is a vector $a \oplus b$ of dimension $d_A + d_B$ with first $d_A$ components
To do this, we need to know that “direct product” is distributive over “direct sum”, for group representations. Let $A$, $B$, and $C$ be vector spaces, with elements $a$, $b$, and $c$. Let $R_A$, $R_B$, and $R_C$ be representations of the same group $G$, so for each $g \in G$ $R_A[g]$ is a matrix acting on vectors in $A$, etc. We want to show that $(R_A \oplus R_B) \otimes R_C$ is basically the same as (isomorphic to) $(R_A \otimes R_C) \oplus (R_B \otimes R_C)$.

The key is to label the vectors in the spaces $(A \oplus B) \otimes C$ and $(A \otimes C) \oplus (B \otimes C)$ in ways that make the correspondence clear. Let us write a vector in the spaces $(A \oplus B) \otimes C$ as $(a \oplus b) \otimes c$ if the $a \oplus b$ state will be in a fifty dimensional Hilbert space. In part (g), we are decomposing a state with $j = 2$ in $R_c[\mathbf{3} \otimes \mathbf{3}]$ into a direct sum of invariant spaces. In our electron-muon system this would correspond to $(3 \otimes 3) \oplus (4 \otimes 1) = (3 \oplus 3 \oplus 3 \oplus 3 \oplus 1 \oplus 5 \oplus 5 \oplus 3 \oplus 3 \oplus 1 \oplus 1)$; two reps of $L = 1$, 2, and 3, plus one rep each with $L = 4$ and 5. Do not forget—the SU(2) representation $S$ has dimension $2S + 1$, so $1 \otimes 2$ is of dimension $3 \times 5 = 15$.

### Footnote 91

If we are being fussy, we should show this is true also for a sum over triples of $a_n$, $b_n$, and $c_n$, since a general vector in the direct product of two spaces is a linear combination of direct products of vectors in the two spaces.
decompose $3 \otimes \bar{3}$ into irreducible representations of the isospin subgroup SU(2). (Hint: There should be five invariant subspaces of this nine-dimensional space.)

(g) How many of the isospin subspaces from part (f) should be part of the octet in Fig. N4.20? Give the total isospin and $I_3$ for each meson, grouping them into isospin multiplets (the mesons forming the different representations).

N4.52 Quantum dissipation from phonons. (Quantum) 2

Fig. N4.21 Atomic tunneling from a tip. Any internal transition among the atoms in an insulator can only exert a force impulse (if it emits momentum, say into an emitted photon), or a force dipole (if the atomic configuration rearranges); these lead to nonzero phonon overlap integrals only partially suppressing the transition. But a quantum transition that changes the net force between two macroscopic objects (here a surface and a STM tip) can lead to a change in the net force (a force monopole). We ignore here the surface, modeling the force as exerted directly into the center of an insulating elastic medium.92 See “Atomic Tunneling from a STM/AFM Tip: Dissipative Quantum Effects from Phonons” Ard A. Louis and James P. Sethna, Phys. Rev. Lett. 74, 1363 (1995), and “Dissipative tunneling and orthogonality catastrophe in molecular transistors”, S. Braig and K. Flensberg, Phys. Rev. B 70, 085317 (2004).

Electrons cause overlap catastrophes (X-ray edge effects, the Kondo problem, macroscopic quantum tunneling); a quantum transition of a subsystem coupled to an electron bath ordinarily must emit an infinite number of electron-hole excitations because the bath states before and after the transition have zero overlap. This is often called an infrared catastrophe (because it is low-energy electrons and holes that cause the zero overlap), or an orthogonality catastrophe (even though the two bath states aren’t just orthogonal, they are in different Hilbert spaces). Phonons typically do not produce overlap catastrophes (Debye–Waller, Frank–Condon, Mössbauer). This difference is usually attributed to the fact that there are many more low-energy electron-hole pairs (a constant density of states) than there are low-energy phonons ($\omega_k \sim ck$, where $c$ is the speed of sound and the wave-vector density goes as $(V/2\pi)^3 d^3 k$).

However, the coupling strength to the low energy phonons has to be considered as well. Consider a small system undergoing a quantum transition which exerts a net force at $x = 0$ onto an insulating crystal:

$$\mathcal{H} = \sum_k p_k^2/2m + 1/2 m\omega^2_k q_k^2 + F \cdot u_0.$$  \hspace{1cm} (N4.85)
Let us imagine a kind of scalar elasticity, to avoid dealing with the three phonon branches (two transverse and one longitudinal); we thus naively write the displacement of the atom at lattice site $x_n$ as $u_n = (1/\sqrt{N}) \sum_k q_k \exp(-ikx_n)$ (with $N$ the number of atoms), so $q_k = (1/\sqrt{N}) \sum_n u_n \exp(i k x_n)$.

Substituting for $u_0$ in the Hamiltonian and completing the square, find the displacement $\Delta_k$ of each harmonic oscillator. (Physically, the force $F$ adds a small linear term to the phonon mode with wavevector $k$, whose minimum becomes displaced by some amount $\Delta_k$.) Let $|F\rangle$ be the ground state of the harmonic oscillators under the force $F$. Write the formula for the likelihood $\langle F|0 \rangle$ that the phonons will all end in their ground states, as a product over $k$ of the phonon overlap integral $\exp(-\Delta_k^2/8\sigma_k^2)$ (with $\sigma_k = \sqrt{\hbar/2m\omega_k}$ the zero-point motion in that mode). Converting the product to the exponential of a sum, and the sum to an integral $\sum_k \sim (V/(2\pi)^3 \int d^3k$, do we observe an overlap catastrophe?

Note that you’ve calculated the probability of a zero-phonon transition—the likelihood that the quantum transition can happen without emitting any phonons is zero. But the same argument shows that there is zero probability of emitting one phonon, or any finite number of phonons. The only allowed transitions emit an infinite number of low-energy phonons. The initial and final ground states are in different Hilbert spaces—no finite number of excitations can connect them.
Bibliography


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