Inelastic Scattering
This book is dedicated to the new user – may you benefit from our mistakes.
Preface

The bright future has arrived for inelastic scattering studies of materials, molecules, and condensed matter. The international science community and national science agencies have invested heavily in new instruments and new sources of neutrons and x-rays that are overcoming many of the historical limitations of inelastic scattering research. The newest generation of transmission electron microscopes offers highly monochromatic electrons that enable measurements of thermal excitations that would otherwise be buried beneath the elastic line. A new family of inelastic laser scattering experiments are now possible, such as those that couple vibrations of the cavity itself to photons in from laser.

Higher intensities and instrument sensitivities are major and obvious improvements. Experimental inelastic scattering has been particularly constrained by low countrates, forcing experimental compromises in energy resolution and intensity. The ARCS inelastic neutron spectrometer with its location at the high-power target station of the Spallation Neutron Source and with its high detection efficiency, for example, provides unprecedented experimental productivity, overcoming many of the restrictions caused by the low countrates that have accompanied inelastic neutron scattering experiments to date.

As features in the data from inelastic scattering experiments become more visible, and averaging processes are less necessary, more information can be extracted from the experimental measurements. The goal of this book is to describe the underlying scattering physics and dynamic processes in materials, and show opportunities to elevate the level of science done with inelastic spectrometers. A large body of specialized knowledge is required to design modern experiments for inelastic instruments such as time-of-flight chopper spectrometers. This body of knowledge will only grow as new capabilities become available. Unfortunately, the underlying concepts are scattered over many disciplines. For example, the books on the theory of thermal neutron scattering by S. W. Lovesey and G. L. Squires are superb. Similarly, excellent solid-state physics texts by J. M. Ziman, C. Kittel, U. Rössler, and N. W. Ashcroft and N. D. Mermin are available for understanding the principles of excitations in condensed matter. Today the concepts from neutron scattering and condensed matter physics are not connected well by existing written texts. It is a challenge to organize a coherent presentation of this wide body of knowledge, connected by the needs of experimental inelastic scattering.

This document did not begin as a textbook. The original draft was a manual
of specifications for the ARCS data analysis software. Defining specifications is a major step in planning a software project and setting its scope. Writing a manual of specifications forces a high degree of detailed planning of classes and modules. As we struggled with these details on software structure, it became obvious that they should parallel as closely as possible the science and practice of experimental inelastic neutron scattering research. Besides the challenge of organizing the higher-level intellectual concepts, practical problems with notation became apparent almost immediately. (Should the scattering vector be \( \vec{Q}, \vec{\kappa} \) or \( \Delta \vec{k} \)? What about its sign?)

The present book is intended for a spectrum of readers spanning from graduate students beginning their doctoral research in inelastic neutron scattering, researchers who need to learn how to use chopper spectrometers and their data analysis, and ourselves, the authors, who need a reference manual. Our heartfelt concern, however, is for the graduate student who enters the field of inelastic neutron scattering with no experience with instruments and only a sketchy understanding of the scientific principles. This text was designed to help the reader understand the principles and methods of inelastic scattering, and do so efficiently.

Explanations are presented at the minimum level of detail required to understand physical concepts. There is some sacrifice of the care in development found in The Theory of Neutron Scattering from Condensed Matter by S. W. Lovesey and Introduction to the Theory of Inelastic Neutron Scattering by G. L. Squires, but our goal was to provide explanations that are “best buys,” producing the most physical insight for the amount of intellectual effort required to understand them. Another goal was to present the field of inelastic scattering as a codified intellectual discipline, showing inter-relationships between different topics. To do so, the notation from other books has been altered in places, for example \( \vec{Q} \), defined as \( \vec{k}_f - \vec{k}_i \), was selected for the scattering vector so that \( \kappa \), defined as \( \vec{\kappa} \equiv \vec{k}_f - \vec{k}_i \), could be consistent with its usage in The Theory of Lattice Dynamics in the Harmonic Approximation by A. A. Maradudin, et al. (It is an unfortunate, but well-established convention that the scattering vector in the diffraction literature is of opposite sign, \( \Delta \vec{k} \equiv \vec{k}_f - \vec{k}_i \).)

Graduate students learning the concepts presented in this book are assumed to have some understanding of scattering experiments – a good understanding of x-ray diffraction would be suitable preparation. The student should have some competence with the manipulation of Patterson functions in Fourier space to the level developed, for example, in Transmission Electron Microscopy and Difractometry of Materials by B. Fultz and J. W. Howe, and should have some understanding of solid-state physics at the level of Principles of the Theory of Solids by J. M. Ziman or Solid-State Physics by H. Ibach and H. Lüth.

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May, 2021
Acknowledgment

This work was supported by the DOE Office of Science, Basic Energy Sciences, under Award No. DE-FG02-03ER46055.
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Chapter 1

Scattering

This chapter on scattering assumes some background knowledge about elastic scattering as used for diffraction studies of atomic structure. (After all, x-ray diffraction is common currency for chemists, materials scientists and condensed matter physicists.) Nevertheless, some of the important concepts in diffraction physics are developed here, and recast it into a formulation based on Patterson functions.

We begin with a discussion of coherence and energy, in part because these concepts are sometimes taken for granted by persons having extensive experience with diffraction experiments. In Sect. 1.4, elastic scattering from static potentials is developed in the Born approximation, giving the basic Fourier transform relationship between the wave and the scattering factor distribution: \( \psi(\vec{Q}) = Ff(\vec{r}) \). This is followed by a brief explanation that shows how time-varying potentials cause inelastic scattering by modulating the frequency of the scattered wave.

More progress in understanding scattering experiments is possible by analyzing the correlation functions that are derived from the measured intensity, rather than the neutron wavefunction (which is not measured directly). Section
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4 returns to elastic scattering to develop the concept of the Patterson function, \( P(\vec{r}) \). The Patterson function is the spatial correlation function that includes all information about the diffracted intensity, as opposed to the diffracted wave. The corresponding correlation function for inelastic scattering is the Van Hove space-time correlation function. It is developed in Sect. 6.1 in Chapter 3 in a path that parallels the Patterson function, but is more general. The double Fourier transform of the Van Hove function, \( G(\vec{r}, t) \), provides all information about the scattered intensity (elastic and inelastic). The Patterson function is a special case of the Van Hove function for static scattering potentials. Graphical examples are used to demonstrate fundamental features of the scattered intensity for the elastic case with the Patterson function: \( I(\vec{Q}) = F_{\vec{r}} P(\vec{r}) \), and for the inelastic case with the Van Hove function: \( I(\vec{Q}, E) = F_{\vec{r}} F_{t} G(\vec{r}, t) \). The two Fourier relationships between the scattered intensity and the correlation functions are discussed in detail in this chapter and the next:

- \( I(\vec{Q}) = F_{\vec{r}} P(\vec{r}) \), which relates the diffracted intensity, \( I(\vec{Q}) \), to the Fourier transform (from space to momentum) of the Patterson function, \( P(\vec{r}) \).

- \( I(\vec{Q}, E) = F_{\vec{r}} F_{t} G(\vec{r}, t) \), which relates the scattered intensity, \( I(\vec{Q}, E) \), to two Fourier transforms (from space to momentum, and time to energy) of the Van Hove function, \( G(\vec{r}, t) \).

For purposes of orientation, we begin with an example.

1.1 A Sampler of Scattering Mechanisms

Neutrons are scattered from materials by several mechanisms. Each scattering mechanism can be taken as independent of the others, since in quantum mechanics the scattering either occurs with full effect, or it does not occur at all. You can usually assume that the scattering mechanisms do not mix, although the average over many scatterings gives an average behavior predicted by classical mechanics. Some categorizations of the different mechanisms of scattering are

- Nuclear or Magnetic. Is the neutron scattered by interaction with the nucleus, or with unpaired (magnetic) electron spins at the atom?

- Elastic or Inelastic. Is the neutron scattered without change of its kinetic energy, or is there an energy transfer between the neutron and the sample?

- Coherent or Incoherent. Is the phase of the neutron wavefunction predictable after scattering, or is there no relationship between the phase of the incident and scattered wavefunction?

- Multiphonon or single phonon. Does the scattering of the neutron cause only one phonon to be excited, or are two excited simultaneously?
1.1. A SAMPLER OF SCATTERING MECHANISMS

• Multiple or single scattering. Is the neutron scattered one time on its way through the sample, or is it scattered more than once?

Combinations across these categories are possible. There is a mechanism of coherent inelastic magnetic single scattering, for example. It is, however, an oxymoron to speak of incoherent coherent scattering. Obviously many mechanisms for neutron scattering are possible. The essentials of these processes will be covered in the next two chapters. For an overview, however, we illustrate several of these possibilities with Fig. 1.1.

Neutron scattering from vanadium is almost entirely incoherent, meaning that the scattering works with the intensities of scattering, but not the phases. This causes a loss of information along the Q direction in Fig. 1.1Va. For example, Bragg diffractions are not seen in the data of Fig. 1.1Va for vanadium. On the other hand, neutron scattering from aluminum is almost entirely coherent, and Fig. 1.1Ala shows rich, detailed structure in Q, and Bragg diffractions along the x-axis at E = 0. (The E = 0 corresponds to elastic scattering, which is essential to diffraction.)

Figures 1.1b-e show the different contributions to the experimental data. These cannot be measured, but they can be computed with reasonable reliability today. From Fig. 1.1Vb and 1.1Alc we can see that the dominant contribution to the scattering is incoherent scattering for V, and coherent scattering for Al. The coherent scattering from V and the incoherent scattering from Al are both small.

At higher temperatures, or with higher momentum transfers of the neutron, there is an increasing possibility of having two or more phonons created by one scattering event. In essence, the scattering is becoming more “classical,” where the scattering is considered to produce heat rather than individual phonons. These measurements of Fig. 1.1 were at low temperatures and with low neutron energies, however, so the multiphonon contribution makes only small changes in panels d. Nevertheless, it is easy to see that multiphonon scattering adds intensity at larger values of Q > 7 Å⁻¹ for both vanadium or aluminum. (in Fig. 1.1, panels d are similar to panels c for Q < 7 Å⁻¹). Finally, the samples were thin, so multiple scattering was unlikely as the neutron traversed the sample. Both multiphonon scattering and multiple scattering can be seen to add some elastic intensity over a broad range of Q.

In this book, all these mechanisms, and more, will be described quantitatively. Methods for optimizing one type of scattering or another will be described, along with experimental practices for measuring them. Different materials have different internal excitations, and some important types of magnetic and phonon excitations will be explained with enough care so that the reader is ready to take the next steps and learn how to calculate them in detail. There is a lot to know about inelastic neutron scattering, and a lot that it can do.

1With multiple scattering this can be possible, but one of the scatterings is fully incoherent, and the other is fully coherent.
Figure 1.1: See description in text. The vertical columns are scattering from bcc vanadium at left and fcc aluminum at right. For phonon scattering, V is nearly fully incoherent, and the Al is nearly fully coherent. Simulations were performed with MCViNE, as described in various parts of this book. (a) experimental data from polycrystalline V and Al. (b) incoherent phonon scattering (notice small scale for Al) (c) coherent phonon scattering (d) multiphonon scattering added to both incoherent and coherent scattering from b and c (e) multiple scattering added to the result of d. This should be comparable to the experimental data in a, since all effects are included.
1.2 Coherence and Incoherence

Diffraction requires “coherent scattering,” characterized by a precise relationship between the phases of the incident and scattered waves. The scattered wave is the sum of component waves, “wavelets” as we call them, emanating from the different atoms in the sample. In diffraction, phase differences between these outgoing wavelets cause constructive or destructive interferences at different angles around the sample, e.g., the appearance of Bragg diffraction peaks.

1.2.1 Wavefunctions

Phase

A wavefunction \( \psi(x, t) \) describes the structure of a wave (its crests and troughs) along position \( x \), at any time \( t \). The mathematical form \( \psi(kx - \omega t) \) accounts for how the wave amplitude shifts in position with increasing time. The argument of the wavefunction, \( kx - \omega t \), is called the “phase” of the wave. It includes two constants: \( k \) (the wavevector), and \( \omega \) (the angular frequency). The phase \( kx - \omega t \) is dimensionless, so it can be used as the argument of a sine function or a complex exponential, for example. Our mathematical form causes the entire structure of the wave \( \psi(kx - \omega t) \) to move to more positive \( x \) with increasing \( t \). This is clear if we recognize that a particular wavecrest in \( \psi \) exists at a particular value of phase, so for larger \( t \), the wave amplitude moves to larger \( x \) for the same value of \( kx - \omega t \).

One-Dimensional Wave

One-dimensional waves are simple because they have no vector character. Suppose the wave is confined a region of length \( L \). The wavefunction and its intensity are:

\[
\psi_{1D}(x, t) = \frac{1}{\sqrt{L}} e^{i(kx - \omega t)}, \tag{1.1}
\]

\[
I_{1D} = \psi_{1D}(x, t) \psi_{1D}^*(x, t) = \frac{1}{L} e^{i(kx - \omega t)} \frac{1}{L} e^{-i(kx - \omega t)} = \frac{1}{L}. \tag{1.2}
\]

If \( \psi_{1D}(x, t) \) were an electron wavefunction, the intensity, \( I_{1D} \), would be a probability density. The prefactor in (1.1) ensures proper normalization, giving a probability of 1 for finding the electron in the interval of length \( L \):

\[
P = \int_0^L I_{1D} \, dx = \int_0^L \frac{1}{L} \, dx = 1. \tag{1.4}
\]

\(^2\)We say \( \psi(kx - \omega t) \) travels to the right with a “phase velocity” of \( \omega/k \). The wave \( \psi(kx + \omega t) \) travels to the left.
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Plane Wave

In three dimensions, a plane wave is:

$$\psi_{3Dpl}(\vec{r}, t) = \frac{1}{\sqrt{V}} e^{i(\vec{k} \cdot \vec{r} - \omega t)},$$  \hspace{1cm} (1.5)

which has an intensity and a normalization analogous to those for the one-dimensional wavefunction. The spatial part of the phase, $\vec{k} \cdot \vec{r}$, is illustrated for a snapshot in time in Fig. 1.2 for two orientations of $\vec{r}$: with $\vec{k} \perp \vec{r}$, and with $\vec{k} \parallel \vec{r}$. Along the direction of $\vec{r}$ in Fig. 1.2a there is no change in the phase of the wave (here $\psi_{3Dpl}(\vec{r}, t) = 1/\sqrt{V} e^{i(0-\omega t)}$), whereas in Fig. 1.2b the phase changes most rapidly along $\vec{r}$ (here $\psi_{3DPlan}(\vec{r}, t) = 1/\sqrt{V} e^{i(kr-\omega t)}$). The dot product $\vec{k} \cdot \vec{r}$ for the phase in (1.5) gives the plane wave its anisotropy in space.

Spherical Wave

By placing the origin of a spherical coordinate system at the center of the spherical wave, the spherical wave has its simplest form:

$$\psi_{3Dsph}(\vec{r}, t) = \frac{1}{\sqrt{V}} e^{i(k|\vec{r}| - \omega t)}. \hspace{1cm} (1.6)$$

If the center of the spherical wave is the distance $\vec{r}'$ away from the origin of the coordinate system:

$$\psi_{3Dsph}(\vec{r}, t) = \frac{1}{\sqrt{V}} \frac{e^{i(k|\vec{r}' - \vec{r}| - \omega t)}}{|\vec{r}' - \vec{r}|}. \hspace{1cm} (1.7)$$

Figure 1.3 shows a vector construction for $\vec{r}' - \vec{r}$, which can be obtained by connecting the tail of $-\vec{r}'$ to the arrow of $\vec{r}$. At distances far from the scattering center, where the curvature of the spherical wave is not important, it is often useful to approximate the spherical wave as a plane wave with $\vec{r}' - \vec{r}$ pointing along the direction of $\vec{k}$.\(^3\)

\(^3\)This is often useful because real scatterers typically emit spherical waves, but Fourier transforms require plane waves.
1.2. COHERENCE AND INCOHERENCE

Figure 1.3: Spherical wave with \( \mathbf{k} \) oriented away from the center of wave emission. (a) with coordinate system for \( \mathbf{r} \) having its origin at center of wave emission. (b) with coordinate system for \( \mathbf{r} \) having an arbitrary origin.

### Phase Factor

A phase factor, \( e^{-i\mathbf{\Delta k} \cdot \mathbf{R}_j} \) or \( e^{-i(\mathbf{\Delta k} \cdot \mathbf{R}_j + \omega t)} \), has the mathematical form of a plane wave (1.5), and is associated with a particular wavelet, but beware. A phase factor is not a wave. A phase factor proves handy when two or more wavelets are scattered from different points in space at \( \{\mathbf{R}_j\} \), typically separated by some atomic distances. What is important after the long path to the detector is how the wavelets interfere with each other – constructively or destructively – and this is accounted for by sums of phase factors like this:

\[
\psi_{\text{phf}}(\mathbf{\Delta k}) = \sum_{\mathbf{R}_j} e^{-i\mathbf{\Delta k} \cdot \mathbf{R}_j}.
\]  

(1.8)

The definition \( \mathbf{\Delta k} \equiv \mathbf{k} - \mathbf{k}_0 \) (illustrated in the chapter title image) is repeated a number of times in this book. This \( \mathbf{\Delta k} \) is a difference in the wavevectors of two actual waves. Dot products like \( \mathbf{\Delta k} \cdot \mathbf{R}_j \) give phase differences between wavelets, but \( \mathbf{\Delta k} \) is not an actual wavevector. Chapter 5 in Fultz and Howe develops these concepts, but the reader is hereby warned that exponentials containing \( \mathbf{\Delta k} \) are not waves, but phase factors.

### 1.2.2 Coherent and Incoherent Scattering

Coherent scattering preserves the relative phases of the wavelets, \( \{\psi_{\mathbf{r}_j}\} \), scattered from different locations, \( \{\mathbf{r}_j\} \), in a material. For coherent scattering, the total scattered wave, \( \Psi_{\text{coh}} \), is constructed by adding the amplitudes of the scattered wavelets:

\[
\Psi_{\text{coh}} = \sum_{\mathbf{r}_j} \psi_{\mathbf{r}_j}.
\]  

(1.9)

The total coherent wave therefore depends on the constructive and destructive interferences of the wavelet amplitudes. Diffraction experiments measure the total coherent intensity, \( I_{\text{coh}} \):

\[
I_{\text{coh}} = \Psi_{\text{coh}}^* \Psi_{\text{coh}} = \left| \sum_{\mathbf{r}_j} \psi_{\mathbf{r}_j} \right|^2.
\]  

(1.10)
On the other hand, “incoherent scattering” does not preserve a phase relationship between the incident wave and the scattered wavelets. For incoherent scattering it is incorrect to add the amplitudes of the scattered wavelets, $\{\psi_{r_i}\}$. Incoherently-scattered wavelets do not maintain phase relationships, so they cannot interfere constructively or destructively. The total intensity of incoherent scattering, $I_{\text{inc}}$, is the sum of individual scattered intensities:

$$I_{\text{inc}} = \sum_{r_i} I_{r_i} = \sum_{r_i} |\psi_{r_i}|^2 .$$

Because measurable intensities are added in incoherent scattering, the angular distribution of incoherent scattering from a group of $N$ identical atoms is the same as for a single atom, irrespective of how these $N$ atoms are positioned in space. The total intensity is simply $N$ times larger. Some types of incoherent scattering occur with a transfer of energy from the wave to the material, and these processes can be useful for spectroscopic analysis of the atom species in a material.

It is important to emphasize the difference between the right-hand sides of (1.10) and (1.11). Because the intensity of coherent scattering in (1.10) first involves the addition of wave amplitudes, coherent scattering depends on the relative phases of the scattered wavelets and the relative positions of the $N$ atoms in the group. Coherent scattering is useful for diffraction experiments. Incoherent scattering is not. This chapter describes in sequence the four types of scattering having coherent components that allow for diffraction experiments on materials:

- x-rays, which are scattered when they cause the atomic electrons to oscillate and re-radiate,
- electrons, which are scattered by Coulomb interactions when they penetrate the positively-charged atomic core,
- neutrons, which are scattered by nuclei (or unpaired electron spins), and
- $\gamma$-rays, which are scattered when they resonantly excite a nucleus, which later re-radiates.

1.2.3 Elastic and Inelastic Scattering

Besides being “coherent” or “incoherent,” scattering processes are “elastic” or “inelastic” when there is, or is not, a change in energy of the wave after scattering. We can therefore construct four word pairs:

(coherent elastic) (coherent inelastic)
(incoherent elastic) (incoherent inelastic)

Diffraction experiments need coherent elastic scattering, whereas spectroscopies that measure intensity versus energy often use incoherent inelastic scattering. The case of incoherent elastic scattering is also common, and occurs, for example, when phase relationships between scattered wavelets are disrupted by
disorder in the material. Incoherent elastic intensity does not show the sharp diffractions associated with crystalline periodicities, but has a broad angular dependence. Finally, coherent inelastic scattering is used in neutron scattering studies of excitations in materials, such as phonons (vibrational waves) or magnons (spin waves), that have precise energy-wavevector relationships. In some phonon studies, a neutron loses energy when creating a phonon (so it is inelastic), but the scattering amplitude depends on the phases of the atom movements in the phonon with respect to the neutron wavevectors (so it is coherent).

A deeper and more rigorous distinction between coherent and incoherent scattering involves our knowledge about the internal coordinates of the scatterer:

- Consider a simple oscillator (a bound electron, for example) that is driven by an incident wave and then re-radiates. There is a transfer of energy from the incident wave to the oscillator, and then to the outgoing wave. Suppose we know in full detail how the coordinates of the oscillator respond to the incident wave. Since the scattering process is determined fully, the phases of all outgoing wavelets have a precise and known relationship to the phase of the incident wave. The scattering is coherent.

- On the other hand, suppose the coordinates of this oscillator were coupled to another system within the material (a different electron, for example), and furthermore suppose there is freedom in how the oscillator can interact with this other system. (Often differing amounts of energy can be transferred from the oscillator to the other system because the transfer occurs by a quantum mechanical process that is not deterministic.) If this energy transfer is different for different scatterings, we cannot predict reliably the phase of the scattered wavelet. The scattering is incoherent.

It is therefore not surprising that incoherence is often associated with inelastic scattering, since inelastic scattering involves the transfer of energy from the scatterer to another component of the material. Incoherence does not imply inelastic scattering, however, and inelastic scattering is not necessarily incoherent.

1.2.4 Wave Amplitudes and Cross-Sections

Cross-Sections

X-rays, electrons, neutrons, and \( \gamma \)-rays are detected one-at-a-time in scattering experiments. For example, the energy of an x-ray is not sensed over many positions, as are ripples that spread to all edges of a pond of water. Either the entire x-ray is detected or not within the small volume of a detector. For x-ray scattering by an individual atomic electron as described in the next section, the scattering may or may not occur, depending on a probability for the x-ray–electron interaction.
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Figure 1.4: These 7 scatterers occupy the fraction 0.2 of the sample area, $A$, and therefore remove the fraction 0.2 of the rays from the incident beam. From (1.12):

$$\sigma = (0.2/7)A.$$ In the thin sample limit, the number of scatterers and the amount of scattering increase in proportion to thickness, $x$, but $\sigma$ remains constant.

An important quantity for scattering problems is the “cross-section,” $\sigma$, which is the effective “target area” presented by each scatterer. With cross-sections it is handy to think of a number, $N$, of scatterers in a sample of area $A$ as in Fig. 1.4. The probability of scattering is equal to the fraction of sample area “blocked” by all $N$ scatterers. For thin samples when the scatterers do not overlap, the $N$ scatterers block an area equal to $N\sigma$. The fraction of rays removed from the incident beam is the blocked area divided by the total area:

$$N \frac{\sigma}{A} = N \frac{\alpha x}{Ax} = \rho \sigma x.$$ (1.12)

Here the density of scatterers, $\rho \equiv N/(Ax)$ has units [scatterers cm$^{-3}$].

To illustrate a salient feature of coherent scattering, consider the elastic scattering of neutrons through the interaction of their spin with the spin polarization of electrons in an antiferromagnet. The total cross-section depends on the total number of unpaired electrons in the material. As mentioned after (1.11), for incoherent scattering the picture would then be complete – the spatial distribution of the scattered intensity is obtained by adding the intensities from independent scattering events from different atoms.

Coherent scattering requires further consideration of the wave amplitudes before calculating the cross-section. A hierarchy of wave interference processes can occur between spin structures on different length scales:

- the unpaired electrons in the same atom (atomic form factor),
- the atoms in the unit cell of the crystal (structure factor),
- the unit cells in the crystal (shape factor), and
- density variations across a material (small angle scattering).

The spatial redistribution of scattered intensity can be spectacularly large in the case of Bragg diffractions, but the total coherent cross-section remains constant. By rearranging the atom positions in a material, the constructive and destructive interferences of coherent scattering are altered and the angles of scattering are redistributed, but for the same incident flux the scattered energy
is conserved (for x-rays or γ-rays), or the total number of scattered particles remains the same (electrons and neutrons).

The flux of scattered x-rays, electrons, neutrons, or γ-rays at the distance \( r \) from the scatterer decreases as \( 1/r^2 \) along \( \hat{r} \). A scattered photon carries energy, so the radiated energy flux also decreases as \( 1/r^2 \) from the scatterer. The energy of a photon is proportional to \( E^*E \), so the electric field, \( E \), has an amplitude that must decrease as \( 1/r \) from the center of scattering. For scattered x-rays, we relate the electric field along \( \hat{r} \) to the incident electric field at the scatterer, \( E_0 \):

\[
E(\vec{r}) \propto E_0 \frac{r}{r},
\]

where the constant of proportionality would include any angular dependence. The electric fields \( E(\vec{r}) \) and \( E_0 \) in (1.13) have the same units, of course, so the constant of proportionality has units of length. The square of this “scattering length” is the cross section per steradian, as we next show for electron scattering (but the argument pertains to all waves).

The problem is similar and actually simpler for neutron scattering, for which this “coherent scattering length” is a constant, \( b \). This \( b \) can be converted to a cross-section by first squaring it (actually taking \( b^*b \) because it is often convenient for \( b \) to be a complex number). The second step is to integrate over all solid angle, which is \( 4\pi \). For all intents and purposes the nucleus that scatters a typical 1 Å neutron is a point and has isotropic scattering, so the relationship between the scattering length and the total coherent cross-section, \( \sigma_{coh} \), is:

\[
\sigma_{coh} = 4\pi|b|^2.
\]

### Cross-Section for Wave Scattering

Here we find the cross-section for wave scattering. Imagine a large sphere of radius \( R \) around the scatterer, and consider the total flux, \( J_{sc}(R) \), scattered through a unit area of surface of this sphere. The incident beam has a flux \( J_{in} \) over an area \( A \). The ratio of all scattered electrons to incident electrons, \( N_{sc}/N_{in} \), is:

\[
\frac{N_{sc}}{N_{in}} = \frac{J_{sc}(R)}{J_{in}} \frac{4\pi R^2}{A \psi_{sc}(R)^2} \frac{4\pi R^2}{\psi_{in}^2 A}.
\]

We consider elastic scattering for which the incident and scattered electrons have the same velocity, \( v \), but for inelastic scattering these factors do not cancel. We use the spherical wave (1.6) for \( \psi_{sc}(R) \) and the plane wave (1.5) for \( \psi_{in} \). For both waves, the exponential phase factors, multiplied by their complex conjugates, give the factor 1. The normalization factors also cancel, so (1.15) becomes:

\[
\frac{N_{sc}}{N_{in}} = \frac{|f_{el}|^2 4\pi R^2}{R^2 A},
\]

(1.16)
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Figure 1.5: The differential scattering cross-section, $d\sigma/d\Omega$, for three paths past a scatterer. The third path, $d\sigma_3/d\Omega_3$, misses the scatterer and contributes only to the forward beam. The paths with areas $d\sigma_1$ and $d\sigma_2$ make contributions to the total cross-section for scattering, $\sigma$, and these contributions are included when the intensity is integrated over the differential solid angles $d\Omega_1$ and $d\Omega_2$.

where $f_{el}/R$ is the fraction of the incident electron amplitude that is scattered into a unit area of the sphere at radius $R$. Figure 1.4 helps demonstrate the fact that the ratio of the cross-section $\sigma$ to the area $A$ of the incident beam equals the ratio of scattered to incident electrons, $N_{sc}/N_{in}$:

$$\frac{\sigma}{A} = \frac{N_{sc}}{N_{in}} = \frac{4\pi|f_{el}|^2}{A},$$  \hspace{0.5cm} (1.17)

$$\sigma = 4\pi|f_{el}|^2.$$  \hspace{0.5cm} (1.18)

The scattering of an x-ray by a single atomic electron can be treated in the same way, but we need to account for the electric dipolar pattern of x-ray radiation with a factor of $2/3$ in the cross-section,

$$\sigma_{xle} = \frac{8\pi}{3}|f_{xle}|^2,$$  \hspace{0.5cm} (1.19)

where $f_{xle}$ is the scattering length. This $f_{xle}$ is the actual constant of proportionality to convert (1.13) into an equality.

Anisotropic scattering is the rule rather than the exception, however, so simple cross-sections like those of (1.18) are usually inadequate, even if altered by factors like the $2/3$ used in (1.19). The “differential scattering cross-section,” written as $d\sigma/d\Omega$, contains the angular detail missing from the total cross-section, $\sigma$.

The differential scattering cross-section, $d\sigma/d\Omega$, is the piece of area offered by the scatterer, $d\sigma$, for scattering an incident x-ray (or electron or neutron) into a particular increment in solid angle, $d\Omega$.

The concept of $d\sigma/d\Omega$ is depicted Fig. 1.5. Note that $d\sigma/d\Omega$ relates an increment in area (on the left) to an increment in solid angle (on the right).

For the simple case of isotropic scattering,

$$\frac{d\sigma}{d\Omega} = |f|^2,$$  \hspace{0.5cm} (1.20)
1.2. COHERENCE AND INCOHERENCE

which is a constant. For anisotropic scattering, (1.20) is generalized with a scattering length, \( f(k_0, \vec{k}) \), that depends on the directions of the incident and outgoing wavevectors, \( \vec{k}_0 \) and \( \vec{k} \), respectively:

\[
\frac{d\sigma}{d\Omega} = |f(k_0, \vec{k})|^2.
\]  
(1.21)

We recover the total cross-section, \( \sigma \), by integrating \( d\sigma/d\Omega \) over all solid angle,

\[
\sigma = \int_{\text{sphere}} d\sigma d\Omega.
\]  
(1.22)

As a check, substituting the constant (1.20) into (1.22) and integrating gives (1.18), as expected.

Special Characteristics of Coherent Scattering

Compare the differential scattering cross-sections for coherent x-ray scattering by a single electron at \( \vec{r}_j, \frac{d\sigma_{x1e,\vec{r}}}{d\Omega} \), and an atom having \( Z \) electrons, \( \frac{d\sigma_{\text{atom}}}{d\Omega} \):

\[
\frac{d\sigma_{x1e,\vec{r}}}{d\Omega}(k_0, \vec{k}) = |f_{x1e,\vec{r}}(k_0, \vec{k})|^2, \\
\frac{d\sigma_{\text{atom}}}{d\Omega} = |f_{\text{atom}}|^2.
\]  
(1.23), (1.24)

In coherent scattering we sum wave amplitudes (cf., (1.9)), so for coherent scattering we sum the scattering lengths of all \( Z \) electrons to obtain the scattering length of an atom:

\[
f_{\text{atom}}(k_0, \vec{k}) = \sum_{\vec{r}} f_{x1e,\vec{r}}(k_0, \vec{k}).
\]  
(1.25)

Note that (1.25) is a sum of the \( f_{x1e,\vec{r}} \), but (1.24) is the square of this sum. Equation (1.24) can predict that the coherent x-ray scattering from an atom with \( Z \) electrons is \( Z^2 \) times stronger than for a single electron, and this proves to be true in the forward direction. However, the total cross-section for coherent scattering must increase linearly with the number of scatterers (here the number of electrons, \( Z \)). Consequently the coherent scattering is suppressed in other directions if a scaling with \( Z^2 \) is allowed in special directions. The angular distribution of coherent scattering must be different for the atom and for the single electron. That is, \( f_{x1e}(k_0, \vec{k}) \) and \( f_{\text{atom}}(k_0, \vec{k}) \) must have different shapes (they must depend differently on \( \vec{k}_0 \) and \( \vec{k} \)). The following is an inequality for coherent scattering (although its analog for incoherent scattering is an equality):

\[
\frac{d\sigma_{\text{atom,coh}}}{d\Omega}(k_0, \vec{k}) \neq \sum_{\vec{r}} \frac{d\sigma_{x1e,\vec{r},\text{coh}}}{d\Omega}(k_0, \vec{k}).
\]  
(1.26)
Integrating (1.26) gives an equality for coherent (and incoherent) scattering:

\[ \int \frac{4\pi}{\text{sphere}} d\sigma_{\text{atom}, \text{coh}} (\vec{k}_i, \vec{k}_f) d\Omega = \int \frac{4\pi}{\text{sphere}} \sum Z \frac{d\sigma_{\text{xl}, \text{coh}}}{d\Omega} (\vec{k}_i, \vec{k}_f) d\Omega , \quad (1.27) \]

because with (1.22) we see that (1.27) equates the individual electron cross-sections to the total cross-section of the atom:

\[ \sigma_{\text{atom}, \text{coh}} = Z \sigma_{\text{xl}, \text{coh}} . \quad (1.28) \]

The process of actually performing the sum in (1.25) evidently requires delicacy in accounting for the phase relationships between the x-ray wavelets scattered into different angles, and knowledge about the electron density of the atom. This is the subject of atomic form factor calculations (see Chapter 3 in Fultz and Howe, for example, or (1.76)).

1.3 Momentum Transfer and Phase Shifts

1.3.1 Momentum and Energy

All processes of scattering redirect the incident wave, causing a change in wavevector, \( \vec{Q} \). Momentum conservation then requires a transfer of momentum, \( \Delta \vec{p} \), from the incident wavefunction to the sample

\[ \vec{Q} = \vec{k}_i - \vec{k}_f, \quad (1.29) \]
\[ \Delta \vec{p} = \hbar \vec{Q}, \quad (1.30) \]

where “i” and “f” denote initial and final. Equation 1.29, shown graphically in Fig. 1.6 is familiar from the elastic scattering in diffraction experiments, where \( |\vec{k}_i| = |\vec{k}_f| \) (and where the quantity \( \Delta \vec{k} = -\vec{Q} \) is often used). The transfer of momentum to the sample (Eq. 1.30) occurs in diffraction experiments, but is rarely considered.

There can be a transfer of energy accompanying a transfer of momentum. Consider the scattering of a particle such as an electron or neutron. The transfer
of energy $E$ is

$$E = \frac{p_i^2}{2M} - \frac{p_f^2}{2M},$$  \hspace{1cm} (1.31)$$

$$E = \frac{\hbar^2}{2M} \left( k_i^2 - k_i^2 \right),$$  \hspace{1cm} (1.32)$$

$$E = -\frac{\hbar^2}{2M} \left( Q^2 - 2k_i \cdot \vec{Q} \right),$$  \hspace{1cm} (1.33)$$

where the last line was obtained after substituting $\vec{k}_f = \vec{k}_i - \vec{Q}$. These relationships between $E$ and $\vec{Q}$ are called the “kinematics” of scattering. Three cases of Eq. 1.33 are noteworthy:

1. When $\vec{k}_i = \vec{k}_f$ and $\vec{Q} = 0$, there is no momentum transfer, and there is no scattering, either. The particle continues with no deflection along the path of the forward beam. Here $E = 0$, of course.

2. When $\vec{Q} = \vec{k}_i$, and $\vec{k}_f = 0$, the incident particle is stopped by the scatterer. This gives maximum energy transfer $E = +\frac{\hbar^2 Q^2}{2M}$.\footnote{Three examples of $\{|Q = \vec{k}_i, \vec{k}_f = 0\}$ are noteworthy:}

3. When $|\vec{k}_i| = |\vec{k}_f| = k$ and $Q = 2k \sin \theta$, the energy transfer $E = 0$ because the scattered particle has its original wavevector $\vec{k}$, and all of its original kinetic energy. For this to occur, the mass of the scatterer must be high, so the scatterer undergoes no recoil.

So one reason why Eq. 1.33 is not used to interpret diffraction experiments is that many atoms participate in the scattering. The mass that absorbs the momentum is therefore large, perhaps the weight of a crystal. This causes $E$ to be small owing to the large $M$ in the denominator of Eq. 1.33. There is little energy in the recoil of a heavy object\footnote{The sample then transfers its momentum to the even more massive Earth, which undergoes a negligible change in its orbit.} when it is hit by a light particle, even if there is a big change in momentum of the lightweight particle. We know that many atoms participate in the diffraction process because the diffraction pattern arises from wave interferences from coherent scattering from many different atoms.
What is more interesting is when the scattering of the particle causes an excitation within the sample. The classical analog is when the scattering generates heat, and is called “inelastic scattering” because kinetic energy is not conserved. The proper accounting for the internal heat allows total energy to be conserved. Our interest is when the internal heat is quantized, however, and the creation of individual magnons or phonons is used in the energy balance. It is appropriate to consider this as “inelastic scattering,” because the incident particle or wave transfers energy to or from the sample, much as a classical particle creates heat.

1.3.2 Phase Shifts

A transfer of momentum always occur in scattering. Here we show why all scattering processes include a factor \( \exp(i\vec{Q} \cdot \vec{R}) \) in the scattering amplitudes (where \( \vec{Q} = \Delta p/\hbar \)).

Both elastic scattering and inelastic scattering involve momentum transfer to the sample, \( \Delta \vec{p} \), as the incident wave (or wavefunction) changes its wavevector from \( \vec{k}_i \) to \( \vec{k}_f \)

\[
\Delta \vec{p} = \hbar \vec{Q} \quad \text{where} \quad \vec{Q} \equiv \vec{k}_i - \vec{k}_f \quad \text{and} \quad \vec{k}_i = \vec{k}_i - \vec{Q},
\]

(1.34)

where i and f designate “initial” and “final.” To accommodate this change in momentum, it is more convenient to work with the momentum representation of a neutron state \( \phi_{\alpha}(\vec{p}) \), rather than the usual spatial representation \( \psi_{\alpha}(\vec{x}) \).

These are related by Fourier transformation

\[
\phi_{\alpha}(\vec{p}) = \frac{1}{\sqrt{2\pi \hbar}} \int_{-\infty}^{\infty} e^{-i\vec{p} \cdot \vec{x}/\hbar} \psi_{\alpha}(\vec{x}) \, d\vec{x},
\]

(1.35)

\[
\psi_{\alpha}(\vec{x}) = \frac{1}{\sqrt{2\pi \hbar}} \int_{-\infty}^{\infty} e^{+i\vec{p} \cdot \vec{x}/\hbar} \phi_{\alpha}(\vec{p}) \, d\vec{p}.
\]

(1.36)

The \( \alpha \) denotes the state of the neutron, and changes after scattering.

Phase Shifts in Elastic Scattering

Consider first an elastic scattering, where the incident and final waves are plane waves, and \( |k_i| = |k| \). After scattering, the final wave \( \phi_f(\vec{p}) \) is shifted in
momentum by $\Delta p = -\vec{Q}/\hbar$ with respect to the initial $\phi_\alpha(\vec{p})$, so Eq. 1.36 becomes

$$\psi_f(\vec{r}) = \frac{1}{\sqrt{2\pi\hbar}} \int_{-\infty}^{\infty} e^{i(\vec{p}_f/\hbar) \cdot \vec{r}} \phi_i(\vec{p} + \hbar \vec{Q}) \, d^3 \vec{p}, \quad (1.37)$$

$$\psi_i(\vec{r}) = \frac{1}{\sqrt{2\pi\hbar}} \int_{-\infty}^{\infty} e^{i(\vec{p}_i/\hbar) \cdot \vec{r}} \phi_i(\vec{p}) \, d^3 \vec{p}, \quad (1.38)$$

$$\psi_f(\vec{r}) = e^{-i\vec{Q} \cdot \vec{r}} \psi_i(\vec{r}), \quad (1.39)$$

$$\psi_i(\vec{r}) = e^{-i\vec{Q} \cdot \vec{r}} \psi_f(\vec{r}), \quad (1.40)$$

The change of variable from $\vec{p}$ to $\vec{p}' = \vec{p} + \hbar \vec{Q}$ from Eq. 1.37 to Eq. 1.38 has no effect on the infinite limits of integration, the phase factor $e^{-i\vec{Q} \cdot \vec{r}}$ is constant in $\vec{p}'$ for Eq. 1.39, and Eq. 1.36 was recognized in going to Eq. 1.40.

Equation 1.40 gives a general effect of momentum transfer in elastic scattering, sometimes described as the effect of “Gallilean invariance” of the scattering process. The amplitude for elastic scattering at a selected $\vec{Q}$ is defined as $f(\vec{Q})$, where

$$f(\vec{Q}) = K \int_{-\infty}^{\infty} \psi_f^*(\vec{r}) \psi_i(\vec{r}) \, d^3 \vec{r}, \quad (1.41)$$

$$f(\vec{Q}) = K \int_{-\infty}^{\infty} e^{i\vec{Q} \cdot \vec{r}} \psi_f^*(\vec{r}) \psi_i(\vec{r}) \, d^3 \vec{r}, \quad (1.42)$$

The constant $K$ depends on the physical process of scattering. For neutrons, $K$ might be the coherent scattering length $b$ of a nucleus. For electron scattering by an atom, $K$ might be $2Z/(a_0 Q^2)$, where $a_0$ is the Bohr radius, $Z$ is atomic number, and the $Q^{-2}$ originates with the Fourier transform of the Coulomb potential. This $f(\vec{Q})$ is often called the atomic form factor for elastic scattering.

**Phase Shifts in Inelastic Scattering**

After scattering, the final state of the neutron is $\beta$, which always differs from the initial state $\alpha$ owing to a change in energy and wavelength after inelastic scattering. We seek the inelastic form factor, $f(\vec{Q}, E)$, which is the amplitude of $\psi_\beta$ for the incident amplitude $\psi_\alpha$. For coherent neutron scattering, this scales with the scattering length $b$, and with the integral $\langle \beta | \alpha \rangle$. In what follows, we integrate over momentum coordinates, but the initial and final coordinates are not the same. The final wavefunction has a shift in its momentum coordinates (complementary to that of the sample in Eq. 1.34)

$$-\vec{Q} = \vec{p}_f/\hbar - \vec{p}_i/\hbar. \quad (1.43)$$

We can use the same momentum coordinate for the final state of the neutron if we make a shift of all momentum coordinates by $-\hbar \vec{Q}$. Equation 1.36 becomes

$$\psi_\beta(\vec{x}) = \frac{1}{\sqrt{2\pi\hbar}} \int_{-\infty}^{\infty} e^{i(\vec{p}/\hbar) \cdot \vec{x}} \phi_\beta(\vec{p}) \, d\vec{p}. \quad (1.44)$$
The $\vec{Q}$ is fixed by our selection of detector angle and energy transfer, so its exponential can be removed from the integration over $\vec{p}$

$$\psi_{\beta f}(\vec{x}) = \frac{1}{\sqrt{2\pi\hbar}} \int_{-\infty}^{\infty} e^{i\vec{Q} \cdot \vec{r}/\hbar} \psi_{\beta j}(\vec{p}) \, d\vec{p}. \quad (1.45)$$

Comparing Eq. 1.45 to Eq. 1.36 we find that the final state of the sample after momentum transfer is

$$\psi_{\beta f}(\vec{x}) = e^{i\vec{Q} \cdot \vec{x}} \psi_{\beta j}(\vec{x}). \quad (1.46)$$

The amplitude of the scattering, or form factor, depends on the overlap of the initial and final states as

$$\langle \beta | \alpha \rangle = \frac{1}{\sqrt{2\pi\hbar}} \int_{-\infty}^{\infty} e^{i\vec{Q} \cdot \vec{x}} \psi^*_{\beta j}(\vec{x}) \psi_{\alpha i}(\vec{x}) \, d\vec{x}. \quad (1.47)$$

For scattering in general

$$f(\vec{Q}) = K \langle \vec{f} | e^{i\vec{Q} \cdot \vec{r}} | \vec{i} \rangle, \quad (1.48)$$

an expression that is often stated with little proof or context. The phase factor $e^{i\vec{Q} \cdot \vec{r}}$, a complex number of modulus unity, is central to all types of scattering. It can be extended to assemblages of many atoms, such as a crystal in Eq. 6.48 below. As a word of warning, explained in Chapter 6, for quantum mechanical scattering by exciting of phonons or magnons, the phase factor becomes an operator expression, where $\vec{Q}$ and $\vec{r}$ are replaced with operators $\vec{Q}$ and $\vec{r}$. Evaluation of the exponential requires its expansion in a power series. Unfortunately, the operators $\vec{Q}$ and $\vec{r}$ do not commute, so subtlety is required when doing the quantum mechanics (Sect. 7.2).

### 1.4 Born Approximation

Almost without a second thought, we treat neutron scattering as a wave phenomenon with the neutron wavefunction satisfying the Schrödinger wave equation. A neutron diffraction pattern, with its sharp peaks, is certainly evidence of wave behavior. The interpretation of the neutron wavefunction is different from that of a simple wave, however. Suppose we were to turn on a monochromatic neutron beam and watch the formation of a diffraction pattern, using an area detector capable of displaying the impacts of individual neutrons. When the neutron beam is turned on, individual counts are recorded at different pixels in the detector array. With time, an obvious bias appears, where the points of detection are most frequently at the positions of the rings and spots of the diffraction pattern. This behavior motivates the interpretation of the neutron wavefunction in terms of probabilities – specifically, the neutron probability is
1.4. BORN APPROXIMATION

the neutron wavefunction times its complex conjugate (which gives a real number). Usually this probabilistic interpretation can be ignored when we consider a diffraction pattern from many neutrons, and we can consider neutron diffraction as the diffraction of any other type of wave. When individual neutron events are considered, however, we may have to recall the probabilistic interpretation of the neutron wavefunction because individual neutron detections look like particles rather than waves.

Another point to remember is that the wave behavior is a characteristic of an individual neutron. When considering a diffraction pattern involving multiple neutrons, we do not add the amplitudes of multiple wavefunctions. Neutrons are fermions, and do not form coherent states as in Bose condensation, for example. At the viewing screen, we add the intensities of individual neutrons. The interactions between different neutrons are not coherent.

Our picture of scattering begins with one neutron as a wave incident on an atom. This wave looks like a plane wave because it comes from a distant source. The wave interacts with the nucleus or magnetic electron cloud of the atom, and an outgoing wave is generated. This outgoing wave is something like a spherical wave originating at the atom, although its intensity is not isotropic. Figure 1.7 shows the geometry, wavevectors and position vectors for our neutron scattering problem. Here both $\vec{r}$ and $\vec{p}$ are large compared to the size of the scatterer. Our plane wave incident from the left, $\Psi_{\text{inc}}$, is of the standard form:

$$\Psi_{\text{inc}} = e^{i(k_i \cdot \vec{r} - \omega t)}.$$  

In what follows we neglect the time dependence to emphasize the manipulations of the spatial coordinates. We later recover the time-dependence by multiplying our results by $e^{-i\omega t}$. A spherical wave, $\Psi_{\text{sc}}$, travels outwards from the center of scattering. The scattered wave has the form:

$$\Psi_{\text{sc}} = f(\vec{k}_i, \vec{k}_f) \frac{e^{i\vec{k}_f \cdot \vec{r}'}}{r} e^{i k_i \cdot \vec{r}'} ,$$  

where the scattering length $f(\vec{k}_i, \vec{k}_f)$ of Sect. 1.2.4 varies with the orientation of $\vec{k}_i$ and $\vec{k}_f$, $\vec{r}'$ is now used to locate the center of the scatterer, and the difference, $\vec{r}' - \vec{r}$, is the distance from the scatterer to the detector. The intensity of $\Psi_{\text{sc}}$ falls off with distance as $1/r^2$, as we expect:

$$I_{\text{sc}} = |\Psi_{\text{sc}}|^2 \Psi_{\text{sc}}^*,$$  

$$I_{\text{sc}} = |f(\vec{k}_i, \vec{k}_f)|^2 \frac{1}{|\vec{r}' - \vec{p}|^2} .$$
To obtain the scattering length \( f(k_i, k_f) \), we must solve the Schrödinger equation for the incident neutron inside the scattering atom (the mass of the neutron is \( m \), and its coordinates in the atom are \( \vec{r}' \)):

\[
-\frac{\hbar^2}{2m} \nabla^2 \Psi(\vec{r}') + V(\vec{r}') \Psi(\vec{r}') = E \Psi(\vec{r}'),
\]

(1.53)

\[
\frac{\hbar^2}{2m} \nabla^2 \Psi(\vec{r}') + E \Psi(\vec{r}') = V(\vec{r}') \Psi(\vec{r}'),
\]

(1.54)

which we write as:

\[
\left( \nabla^2 + k_i^2 \right) \Psi(\vec{r}') = U(\vec{r}') \Psi(\vec{r}'),
\]

(1.55)

after having made the two definitions:

\[
k_i^2 \equiv \frac{2mE}{\hbar^2},
\]

(1.56)

\[
U(\vec{r}') \equiv \frac{2mV(\vec{r}')}{\hbar^2}.
\]

(1.57)

The formal approach to finding the solution of the Schrödinger equation in this problem makes use of Green’s functions. A Green’s function, \( G(\vec{r}, \vec{r}') \), provides the response at \( \vec{r} \) for a point scatterer at \( \vec{r}' \):

\[
\left( \nabla^2 + k_i^2 \right) G(\vec{r}, \vec{r}') = \delta(\vec{r}').
\]

(1.58)

We find the Green’s function in a quick way by starting with an identity:

\[
\frac{\nabla^2}{r} e^{ikr} = \frac{e^{ikr}}{r} \frac{1}{r} - k^2 \frac{e^{ikr}}{r},
\]

(1.59)

\[
\frac{(\nabla^2 + k^2)}{r} e^{ikr} = e^{ikr} \frac{1}{r},
\]

(1.60)

Recall that:

\[
\frac{\nabla^2}{r} \frac{1}{r} = -4\pi \delta(r), \quad \text{so}
\]

\[
\left( \nabla^2 + k_i^2 \right) \frac{1}{r} = -e^{ikr} 4\pi \delta(r).
\]

(1.61)
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The \( \delta \)-function simplifies the right-hand side, forcing it to zero everywhere except at \( r = 0 \). At \( r = 0 \), however, \( e^{ikr} = 1 \). From our identity (1.59) we therefore obtain:

\[
\left( \nabla^2 + k^2 \right) \frac{e^{ikr}}{r} = -4\pi \delta(r) .
\] (1.63)

We make a shift of the origin: \( \vec{r} \rightarrow \vec{r} - \vec{r}' \) (so we can see more easily how the outgoing wave originates at the scatterer – see Fig. 1.7). After doing so, we identify our Green’s function by comparing (1.58) and (1.63):

\[
G(\vec{r}, \vec{r}') = -\frac{1}{4\pi} \frac{e^{ik|\vec{r}-\vec{r}'|}}{|\vec{r} - \vec{r}'|} .
\] (1.64)

With our Green’s function in hand, we construct \( \Psi_{\text{sc}}(\vec{r}) \) by integrating. The idea is that to obtain the total wave amplitude at \( \vec{r} \), we need to add up the spherical wavelet amplitudes emanating from all \( \vec{r}' \) (each of form (1.64)), weighted by their strengths. This weight is the right-hand side of (1.55). Formally, the limits of integration cover all of space, but in fact it is only important to extend them over the \( \vec{r}' \) where \( U(\vec{r}') \) is non-zero (approximately the volume of the atom).

\[
\Psi_{\text{sc}}(\vec{r}) = \int U(\vec{r}') \Psi(\vec{r}') G(\vec{r}, \vec{r}') \, d^3\vec{r}' .
\] (1.65)

The total wave at \( \vec{r} \), \( \Psi(\vec{r}) \), has both incident and scattered components:

\[
\Psi(\vec{r}) = \Psi_{\text{inc}} + \Psi_{\text{sc}} , \quad \Psi(\vec{r}) = e^{ik\vec{r}' \cdot \vec{r}} + \frac{2m}{\hbar^2} \int V(\vec{r}') \Psi(\vec{r}') G(\vec{r}, \vec{r}') \, d^3\vec{r}' .
\] (1.66, 1.67)

1.4.2 First Born Approximation

Up to here our solution is exact. It is, in fact the Schrödinger equation itself, merely transformed from a differential equation to a integral equation appropriate for scattering problems. The problem with this integral equation (1.65) is that \( \Psi \) appears both inside and outside the integration, so an approximation is generally required to proceed further. The approximation that we use is the “first Born approximation.” It amounts to using a plane wave, the incident plane wave, for \( \Psi \) in the integral:

\[
\Psi(\vec{r}') \approx e^{ik\vec{r}' \cdot \vec{r}} .
\] (1.68)

The first Born approximation assumes that the wave is undiminished and scattered only once by the material. This assumption is valid when the scattering is weak.

We simplify (1.64) by making the approximation that the detector is far from the scatterer. This allows us to work with plane waves at the detector, rather
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than outgoing spherical waves. To do so we align the outgoing wavevector $\vec{k}_f$ along $(\vec{r} - \vec{r}')$ as shown in Fig. 1.7. The product of scalars, $k_f |\vec{r} - \vec{r}'|$, in the exponential of a spherical wave emitted from $\vec{r}'$, is then equal to $\vec{k}_f \cdot (\vec{r} - \vec{r}')$ of a plane wave,

$$G(\vec{r}, \vec{r}') \approx -\frac{1}{4\pi} \frac{e^{i\vec{k}_f \cdot (\vec{r} - \vec{r}')}}{|\vec{r}|}.$$  (1.69)

In (1.69) we also assumed that the origin is near the scatterer, so $|\vec{r}| \gg |\vec{r}'|$, simplifying the denominator of our Green’s function.

Returning to our exact integral equation (1.67), we obtain the approximate scattered wave (the first Born approximation for the scattered wave) by using (1.68) and (1.69) in (1.67):

$$\Psi(\vec{r}) \approx e^{i\vec{k}_i \cdot \vec{r}} - \frac{m}{2\pi \hbar^2} \int V(\vec{r}') e^{i\vec{k}_i \cdot \vec{r}'} e^{i\vec{k}_f \cdot (\vec{r} - \vec{r}')} d^3\vec{r}'$$  (1.70)

$$\Psi(\vec{r}) = e^{i\vec{k}_i \cdot \vec{r}} - \frac{m}{2\pi \hbar^2} \frac{e^{i\vec{k}_f \cdot \vec{r}'} \int V(\vec{r}') e^{i\vec{k}_i \cdot \vec{r}'} d^3\vec{r}'}{|\vec{r}|}.$$  (1.71)

If we define:

$$\vec{Q} \equiv \vec{k}_i - \vec{k}_f,$$

$$\Psi(\vec{r}) = e^{i\vec{k}_i \cdot \vec{r}} - \frac{m}{2\pi \hbar^2} \frac{e^{i\vec{k}_f \cdot \vec{r}'} \int V(\vec{r}') e^{i\vec{Q} \cdot \vec{r}'} d^3\vec{r}'}{|\vec{r}|}.$$  (1.72)

The scattered part of the wave is:

$$\Psi_{sc}(\vec{Q}, \vec{r}) = \frac{e^{i\vec{Q} \cdot \vec{r}}}{|\vec{r}|} f(\vec{Q}),$$  (1.74)

where:

$$f(\vec{Q}) \equiv -\frac{m}{2\pi \hbar^2} \int V(\vec{r}') e^{i\vec{Q} \cdot \vec{r}'} d^3\vec{r}'.$$  (1.75)

The factor $f(\vec{Q})$ is the scattering factor of (1.50), which we have found to depend on the incident and outgoing wavevectors only through their difference, $\vec{Q} \equiv \vec{k}_i - \vec{k}_f$. We recognize the integral of (1.75) as the Fourier transform of the potential seen by the incident neutron as it goes through the scatterer. In the first Born approximation:

The scattered wave is proportional to the Fourier transform of the scattering potential.

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6If we neglect a constant prefactor, this assumption of $|\vec{r} - \vec{r}'| = |\vec{r}|$ is equivalent to assuming that the scatterer is small compared to the distance to the detector.

7Sadly, the diffraction vector for elastic scattering is defined as $-\vec{Q} = \Delta \vec{k} \equiv \vec{k}_f - \vec{k}_i$
The factor $f(\vec{Q})$ of (1.75) is given various names, depending on the potential $V(\vec{r})$ (we changed notation: $\vec{r} \to \vec{r}$). When $V(\vec{r})$ is the potential of a single atom, $V_{\text{at}}(\vec{r})$, we define $f_{\text{at}}(\vec{Q})$ as the "atomic form factor":

$$f_{\text{at}}(\vec{Q}) \equiv -\frac{m}{2\pi\hbar^2} \int V_{\text{at}}(\vec{r}) e^{i\vec{Q} \cdot \vec{r}} d^3\vec{r}.$$  \hspace{1cm} (1.76)

Alternatively, we can use the potential for the entire crystal for $V(\vec{r})$ in (1.75) (and develop the kinematical theory of diffraction). When $V(\vec{r})$ refers to the entire crystal, however, the first Born approximation of 1.73 is sometimes unreliable because multiple scattering invalidates the assumption of (1.68). This assumption is, nevertheless, the basis for the "kinematical theory of diffraction," which we develop for its clarity and its qualitative successes. It is possible to transcend formally the single scattering approximation, and develop a "dynamical theory" of neutron diffraction by considering higher-order Born approximations, but this has not proved a particularly fruitful direction. Modern dynamical theories take a completely different path.

### 1.4.3 Higher-Order Born Approximations

Nevertheless, it is not difficult in principle to extend the Born approximation to higher orders. Instead of using an undiminished plane wave for $\Psi(\vec{r})$, we could use a $\Psi(\vec{r})$ that has been scattered once already. Equation (1.67) gives the second Born approximation if we use do not use the plane wave of (1.68) for $\Psi(\vec{r})$, but rather:

$$\Psi(\vec{r}) = e^{i\vec{k} \cdot \vec{r}} + \frac{2m}{\hbar^2} \int V(\vec{r}''') \Psi(\vec{r}''') G(\vec{r}', \vec{r}'''') d^3\vec{r}'''',$$  \hspace{1cm} (1.77)

where we now use a plane wave for $\Psi(\vec{r}''')$:

$$\Psi(\vec{r}''') \approx e^{i\vec{k} \cdot \vec{r}'''}.$$  \hspace{1cm} (1.78)

The second Born approximation involves two centers of scattering. The first is at $\vec{r}''$ and the second is at $\vec{r}$ (as shown in Fig. 1.8). The second and higher Born approximations are not used very frequently. If the scatterer is strong enough to violate the condition of weak scattering used in the first Born approximation, the scattering will also violate the assumptions of the second Born approximation.
Figure 1.8: Coordinates for the second Born approximation. The neutron path is shown as the dark arrows, which are labeled by the relevant Green’s functions.
Chapter 2

Inelastic Electron Scattering and Spectroscopy

2.1 Inelastic Electron Scattering Principles

This Chapter 2 first describes how high-energy electrons are scattered inelastically by materials, and then explains how electron energy-loss spectrometry (EELS) is used in materials research. Inelastic scattering occurs by the processes listed below in order of increasing energy loss, $E$. Although the scattering is called “inelastic,” energy is conserved – the spectrum of energy gains by the sample is mirrored in the spectrum of energy losses of the high-energy electrons.

Electrons undergoing energy losses to crystal vibrations, quantized as phonons with $E \sim 10^{-2}$ eV, are indistinguishable from elastically scattered electrons, given the present state-of-the-art for EELS in a TEM.

With modern instrumentation, it is possible to measure interband transitions of electrons from occupied valence bands to unoccupied conduction bands of semiconductors and insulators. With $E \sim 2$ eV, these spectral features are quite close in energy to the intense zero-loss peak from elastic scattering, so resolving them has been a challenge.

In many solids, especially metals, the bonding electrons can be understood as a gas of free electrons. When a high-energy electron suddenly passes through this electron gas, plasmons may be created. Plasmons are brief oscillations of the free electrons, giving broadened peaks in EELS spectra. Plasmon energies ($E \sim 10$ eV) increase with electron density, so plasmon spectra can be used to estimate free electron density. Plasmon spectra are also useful for measuring the thickness of a TEM specimen because more plasmons are excited as the electron traverses a thicker specimen.

Electrons that ionize atoms by causing core excitations are used for micro-chemical analysis. Chemical spectroscopy with EELS measures the intensities
of “absorption edges,” which are jumps in spectral intensity at the threshold energies for ejecting core electrons from atoms in the material ($10^2 < E < 10^4$ eV). After a core electron has been excited from the atom, the remaining “core hole” decays quickly, often by the emission of a characteristic x-ray. Characteristic x-rays with energies from $>10^2$ to $>10^4$ eV are used in energy dispersive x-ray spectrometry (EDS) for chemical analysis.

Methods

“Analytical transmission electron microscopy” uses EDS or EELS to identify the elements in a specimen, and to measure elemental concentrations or spatial distributions. To quantify chemical concentrations, a background is subtracted to isolate the heights of absorption edges (EELS) or the intensities of peaks in an x-ray energy spectrum (EDS). These isolated intensities are then compared for the different elements in the specimen, and often converted into absolute concentrations with appropriate constants of proportionality. The accuracy of quantification depends on the reliability of these constants, so significant effort has been devoted to understanding them.

In this chapter, after brief descriptions of an EELS spectrometer and features of a typical EELS spectrum, plasmon energies are discussed with a simple model of a free electron gas. The section on “core excitations” provides a higher-level treatment of how a high-energy electron can cause a core electron to be ejected from the atom. It turns out that the probability of a core electron excitation is proportional to the square of the Fourier transform of the product of the initial and final wavefunctions of the excited electron. The cross-section for inelastic scattering also has an angular dependence that must be considered when making quantitative measurements with EELS. Some experimental aspects of EELS measurements are presented, including energy-filtered TEM imaging.

This chapter then presents the principles of EDS in the TEM, which involves more physical processes than EELS. Interestingly, the cross-section for core ionization decreases with atomic number, but the cross-section for x-ray emission increases with atomic number in an approximately compensating way. This gives EDS spectrometry a balanced sensitivity for most elements except the very lightest ones.

2.2 Electron Energy-Loss Spectrometry (EELS)

2.2.1 Instrumentation

Spectrometer

After electrons have traversed a TEM specimen, a significant minority of them have lost energy to plasmons or core excitations, and exit the specimen with energies less than $E_0$, the energy of the incident electrons. ($E_0$ may be $200,000 \pm 0.5$ eV, for example.) To measure the energy spectrum of these losses, an EELS spectrometer can be mounted after the projector lenses of a TEM. The heart of a
transmission EELS spectrometer is a magnetic sector, which serves as a prism to disperse electrons by energy. In the homogeneous magnetic field of the sector, Lorentz forces bend electrons of equal energies into arcs of equal curvature. Some electron trajectories are shown in Fig. 2.1.

The spectrometer must allow an angular range for electrons entering the magnetic sector, both for reasons of intensity and for measuring how the choice of scattering angle, $\phi$, affects the spectrum (cf., (2.44)). A well-designed magnetic sector provides good focusing action. Focusing in the plane of the paper (the equatorial plane) is provided by the magnetic sector of Fig. 2.1 because the path lengths of the outer trajectories are longer than the path lengths of the inner trajectories. It is less obvious, but also true, that the fringing fields at the entrance and exit boundaries of the sector provide an axial focusing action. With good electron optical design, the magnetic sector is “double-focusing” so that the equatorial and axial focus are at the same point on the right of Fig. 2.1. Since the energy losses are small in comparison to the incident energy of the electrons, the energy dispersion at the focal plane of typical magnetic sectors is only a few microns per eV.

Electrons that lose energy to the sample move more slowly through the magnetic sector, and are bent further upwards in Fig. 2.1. In a “serial spectrometer,” a slit is placed at the focal plane of the magnetic sector, and a scintillation counter (see Sect. ??) is mounted after the slit. Intensity is recorded only from those electrons bent through the correct angle to pass through the slit. A range of energy losses is scanned by varying the magnetic field in the spectrometer. A “parallel spectrometer,” shown in the chapter title image, covers the focal plane of the magnetic sector with a scintillator and a position-sensitive photon detector such as a photodiode array. The post-field lenses Q1–Q4 magnify the energy dispersion before the electrons reach the scintillator. A parallel spectrometer has an enormous advantage over a serial spectrometer in its rate of data acquisition, but it requires calibrations for variations in pixel sensitivity.

The optical coupling of a magnetic sector spectrometer to the microscope usually puts the object plane of the spectrometer at the back focal plane of

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1Their longer time in the magnetic field overcomes the weaker Lorentz forces.
the final projector lens (Fig. ??). This back focal plane contains the diffraction pattern of the sample when the microscope is in image mode. When the microscope is operated in image mode, the spectrometer is therefore said to be “diffraction-coupled” to the microscope. With diffraction coupling, the collection angle, \( \beta \), of the spectrometer is controlled by the objective aperture of the microscope. Alternatively, when the microscope is operated in diffraction mode, the back focal plane of the projector lens contains an image, and the spectrometer is said to be “image-coupled” to the microscope. With image-coupling, the collection angle, \( \beta \), is controlled by an aperture at the entrance to the spectrometer (at top of Fig. 2.1).

**Monochromator**

The typical energy resolution for EELS spectrometers was about 1 eV or so for many years, but recent developments have allowed energy resolutions better than 0.1 eV on commercial microscopes. This is accomplished by starting with a field emission gun, often a Schottky effect gun (Sect. ??), followed by an electron monochromator, often a Wien filter as described here. Electrons traveling through a Wien filter encounter a region with crossed electric and magnetic fields that induce competing forces on the electron. In a Wien filter these electric and magnetic forces are tuned to cancel for electrons of one velocity, \( v_0 \), which avoid deflection and pass through the exit aperture of the filter. Specifically, for an electron with velocity \( v_z \) down the optic axis along \( \hat{z} \), a magnetic field oriented along \( \hat{y} \), produces a force along \( \hat{x} \), \( F_x^{\text{mag}} = ev_zB_y \). A Wien filter has an electric field along \( \hat{x} \) in the same region, generating a force on the electron of \( F_x^{\text{el}} = -eE_x \). The special condition of cancelling forces, \( F_x^{\text{mag}} = -F_x^{\text{el}} \), can be true for electrons of only one velocity, \( v_0 \):

\[
\begin{align*}
e v_0 B_y &= eE_x, \\
v_0 &= \frac{E_x}{B_y}.
\end{align*}
\]

Electrons with velocities differing from \( v_0 \) are deflected, and do not pass through the exit aperture of the Wien filter. In practice, it is typical to operate the Wien filter at a voltage close to that of the electron gun itself, so the electron velocity through the filter will be slow enough that sub-eV resolution is possible with reasonable values of electric field, magnetic field, and aperture size. Biasing the Wien filter assembly near –100 or –200 keV can be challenging, however.

The Wien filter first disperses electrons of different energies into different angles, and then allows electrons of only a selected energy to pass through its exit aperture. Monochromatization therefore discards a substantial fraction of electrons – perhaps 80% of the electrons are lost when monochromating to 0.1 eV. When operating in STEM mode, the electron current is also reduced substantially when forming the smallest electron probes. It is typical to make compromises between the brightness of the image, the electron monochromatization, and the size of the probe – an increase in one usually requires a decrease
in another. Manufacturers are constantly trying to find ways to improve these aspects of microscope performance.

### 2.2.2 General Features of EELS Spectra

A typical EELS spectrum is presented in Fig. 2.2. The enormous “zero-loss peak” is from electrons of 200,000 eV that passed through the specimen without any energy loss. The sharpness of this peak indicates that the energy resolution is about 1.5 eV. The next feature is at the energy loss $E = 25$ eV, from electrons having energies of 199,975 eV. It is the “first plasmon peak,” caused by the excitation of one plasmon in the sample. With thicker specimens there may also be peaks at multiples of 25 eV from electrons that excited two or more plasmons in the specimen. The small bump in the data at 68 eV is not a plasmon peak, but rather a core loss. Specifically it is a Ni $M_{2,3}$ absorption edge caused by the excitation of 3$p$ electrons out of Ni atoms. An enormous feature is seen at an energy loss of about 375 eV, but it is an artifact of the serial data acquisition method, and not a feature of the material. (At 375 eV the detector operation was changed from measuring an analog current to the counting of individual electron events.)

The background in the EELS spectrum falls rapidly with energy (the denominator of $\Delta k^2$ in (2.28) is partially responsible for this), and the next feature in the Ni spectrum of Fig. 2.2 is a core loss edge at 855 eV. This feature is caused by the excitation of 2$p^{3/2}$ electrons out of the Ni atom, and is called the “$L_3$ edge.” The $L_2$ edge at 872 eV is caused by the excitation of 2$p^{1/2}$ electrons out of the atom. Right at the $L_2$ and $L_3$ edges are sharp, intense peaks known as “white lines” that originate from the excitation of 2$p$ electrons into unoccupied 3$d$ states at a Ni atom. Such features are typical of transition metals and their alloys as described in Section 2.2.3. More generally, unoccupied states such as antibonding orbitals are often responsible for sharp peaks at core edges.

Compared to plasmon excitations, the cross-sections for inner-shell ioniza-
tions are relatively small, and become smaller at larger energy losses. To obtain good intensities, for many elements it is preferable to use absorption edges at lower energy losses (e.g., $L$ and $M$). Some of the nomenclature of electronic transitions was given previously in Sect. ?? Figure 2.3 shows an orbital representation and associated nomenclature for EELS edges.

### 2.2.3 * Fine Structure*

**Near-Edge Fine Structure**

The region in an EELS spectrum around a core-loss edge often shows clear and reproducible structure that can be used to identify the local chemical environment. This “electron energy-loss near-edge structure” (ELNES) depends on the number and energy of unoccupied states at the excited atom. Chemists call these low-lying unoccupied states “lowest unoccupied molecular orbitals,” and they include antibonding orbitals. Physicists call them “states above the Fermi energy,” and they include the conduction band. A core electron can be excited into these unoccupied states, and the energy gained by the core electron during this transition is mirrored in the energy-loss spectrum of the high-energy electron. Simple metals with nearly-free electrons show core edges in EELS spectra that are smooth and without sharp features. On the other hand, materials with high densities of states just above the Fermi level, such as transition metals and rare-earth metals, have sharp features at their absorption edges associated with transitions into unoccupied $d$- and $f$-states, respectively. These features do not appear at all absorption edges, owing to the dipole selection rule where the angular momentum must change by $\pm 1$. This selection rule allows transition metals with unoccupied $d$-states to have intense white lines at their $L_{2,3}$ edges, which involve excitations from core $p$-electrons, but not at their $L_1$ edges, which involve excitations from $s$-electrons (see Fig. 2.2).

The intensity of the white lines at the $L_{2,3}$ edge of Ni in Fig. 2.2 can be understood with the inelastic cross section for core shell ionizations (2.37), where $\psi_\beta$ is an unoccupied $3d$ state and $\psi_\alpha$ is an occupied $2p$ core state, both centered at the Ni atom. The intensities of the white lines are larger when there are more unoccupied $3d$ states (the factor $\rho(E)$ in (2.37)). If the integral in (2.37) is evaluated, integrated intensities of the white lines can be used
to quantify $\rho(E)$, the number of unoccupied $3d$ states at Ni atoms, and how this number changes with alloying or chemical bonding. Likewise, rare earth metals with unoccupied $f$-states have sharp features at their $M_{4,5}$ edges, which involve core $d$-electrons (but not at their $M_2$ or $M_3$ edges, which involve $p$-electrons). Semiconductors and insulators usually show distinct structure at their absorption edges, owing to the excitation of core electrons into unoccupied states above the band gap.

Because the number of unoccupied states is sensitive to the chemical and structural environment around the excited atom, ELNES can be used as a “fingerprint” of its local environment, even when the experimental systematics are not simple, or when electronic structure calculations are not possible. Figure 2.4 shows that the oxygen $K$-edge ELNES is sensitive to the local environment around the O atom in a variety of manganese oxides. The structure around 527-532 eV is dominated by the effects of chemical bonding on the density of electron states at the O atom, but the peak from 537-545 eV is more sensitive to the local positions of Mn atoms near the O atom – it is part of the “extended fine structure,” described below.

Changes to the chemical environment around an atom alter the energy of the lowest unoccupied state, and therefore shift the onset energy of the core edge. Chemical shifts of absorption edges therefore reflect changes in the energies of the unoccupied states. More subtly, however, they also reflect changes in the energies of the core states. Any change to the outer atomic electrons, as caused by changes in chemical bonding for example, alters the intra-atomic electron-electron interactions. The energies of core electrons are therefore altered by changes in the outer electrons. For example, if an outer electron of a Li atom is transferred to a neighboring F atom, one may expect a lower-energy unoccupied state about the Li, and a shift of the Li $K$ edge to lower energy. In fact, however, the loss of this electron in Li$^+$ reduces the screening of the core 1s electrons, causing them to be more tightly bound to the nucleus. This causes the absorption edge to shift to higher energies. Lithium has only three
electrons, so this effect is anomalous, but chemical shifts of absorption edges for all elements depend in part on the shifts in energies of the core electrons caused by intra-atomic screening.

Finally, we note that the core hole itself alters the energies of the atomic electrons. It is sometimes assumed that the removal of a core electron serves to increase the effective nuclear charge from $Z$ to $Z + 1$, but the effects of a core hole on the energy levels of an unstable atom are not easy to understand.

**Extended Fine Structure**

Extended electron energy-loss fine structure (EXELFS) starts at energies where the outgoing electron state can be considered free of the atom, perhaps about 30 eV beyond the absorption edge. The state of the outgoing electron from the “central atom” is affected by the surrounding atoms, and self-interference occurs as the outgoing electron is backscattered from the nearest-neighbor shells of atoms. This process is illustrated schematically in Fig. 2.5. With changes in the wavelength of the outgoing electron, constructive and destructive interference occurs, causing the EXELFS signal, $\chi$, to be oscillatory:

$$\chi(k) = \sum_j \frac{N_j f_j(k)}{r_j^2} N_j f_j(k) e^{-2r_j/\lambda} e^{-2\sigma_j^2} \sin(2kr_j + \delta_0 + \delta_j).$$  \hspace{1cm} (2.2)

Equation (2.2) includes a number of different effects, and its factors are best justified one-by-one. The sine function is the oscillatory interference of the outgoing electron wavefunction with itself as it travels the distance $2r_j$ from the central (excited) atom to a neighboring atom at $r_j$ and back again. The phase of this electron wave is shifted by the amount $\delta_j$ upon scattering by the neighboring atom at the distance $r_j$, and by $\delta_0$ from the central atom. These phase shifts generally depend on the electron wavevector, and this $k$-dependence must be known for quantitative work. The other factors in (2.2) are the number and backscattering strength of the neighboring atoms, $N_j$ and $f_j(k)$, a qualitative decay factor to account for the finite lifetime of the outgoing electron state, $e^{-2r_j/\lambda}$ (where $\lambda$ is the electron mean-free-path), and a Debye-Waller factor, $e^{-2\sigma_j^2}$, that attenuates $\chi(k)$. Here $\sigma_j^2$ is a mean-squared displacement of the central atom relative to its neighboring atoms, typically originating with temperature or disorder in the local structure. The sum in (2.2) is over the neighboring atoms, and typically includes the first- and second-nearest-neighbor (1nn and 2nn) shells around the central atom.

Figure 2.6 shows some steps in a typical EXELFS analysis, in this case for the $L_{2,3}$ edge from a slightly-oxidized sample of bcc Fe metal. Figure 2.6a shows the absorption edge after correction for the pre-edge background. The region of interest begins above the $L_3$ and $L_2$ edges. Unfortunately, the $L_1$ edge ($2s$ excitation) appears as a feature in the region of interest, so it is best to work with data at energies beyond the $L_1$ edge. The useful data range did include the oscillations with broad peaks at about 920 and 1000 eV (barely visible in Fig.
2.6. Extracting these small oscillations from the monotonic decay characteristic of an isolated atom is usually done by fitting a cubic spline function through the EXELFS oscillations. Subtracting this spline fit reveals the oscillations in energy, which are converted to $k$-space as in Fig. 2.6b, using the wavevector dependence on energy above the absorption edge, $E_a$ (where $k$ is in Å$^{-1}$):

$$E - E_a = \frac{\hbar^2 k^2}{2m_e} = 3.81 \, k^2 \, [\text{eV}],\; (2.3)$$

Real space periodicities are obtained from the data of Fig. 2.6b by taking their Fourier transform. The periodicities in real space are not affected significantly if $\chi(k)$ is multiplied by a power of $k$, and doing so helps to sharpen the peaks in the real space data. The real-space function of Fig. 2.6c, called a “pseudo-” or “raw-” radial distribution function, was obtained by taking the Fourier transform of $k\chi(k)$. The peak at 2.25 Å corresponds approximately to the position of the 1nn shell of Fe atoms in bcc Fe (2.02 Å), but a discrepancy is expected because the phase shifts $\delta_j$ and $\delta_0$ of (2.2) were not included in the data analysis. For comparative work with similar specimens, however, this simple Fourier transform method may be adequate.

Better-known than EXELFS is EXAFS (extended x-ray absorption fine structure) spectroscopy, performed with tuneable synchrotron radiation. EXAFS is identical to EXELFS, except that the excitation of the central atom is caused by a photon. The energy of the incident photon is tuned from below an absorption edge to well above it. The self-interference of the backscattered photoelectron is seen in the data as decreased or increased photon transmission through the sample (or electron yield in another variant of the EXAFS technique). The

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2. It is an approximation to ignore the slight phase difference between the $L_1$ and $L_2$ EXELFS oscillations, and to neglect the $L_1$ EXELFS, but the approximation is not too bad.
Figure 2.6: (a) Fe $L$-edge from pure Fe metal at 97 K. Pre-edge background was subtracted, but no corrections were performed for plasmon excitations, which do not affect the gradual EXELFS structure. (b) Fe $L_{2,3}$ edge EXELFS extracted from data in a. (c) Magnitude of Fourier transform of data in b. After [5.3].
2.3. PLASMON EXCITATIONS

analysis of the $\chi(k)$ data is identical to that of EXELFS, and (2.2) was originally proposed for EXAFS.

There is a stronger $E$-dependence for EXELFS spectra than for EXAFS spectra, causing EXELFS to be more practical than EXAFS for energies below about 2 keV. Nevertheless, EXAFS is more practical at higher energies, and higher energies have two advantages. Atomic levels at higher energies are better separated in energy, making it easier to obtain wide ranges of energy where the extended fine structure can be measured without interruption from other absorption edges. The second advantage of EXAFS is its ability to work with $K$-shell excitations of many elements, whose simpler structure allows their $\chi(k)$ to be interpreted more reliably. On the other hand, EXELFS can be performed readily on local regions of material identified in TEM images. Synchrotron beamline optics including x-ray mirrors and Fresnel zone plates now allow EXAFS measurements on areas smaller than $\sim 1 \mu m$, however.

2.3 Plasmon Excitations

2.3.1 Plasmon Principles

A fast electron jolts the free electrons when it passes through a material. The displaced charge creates an electric field to restore the equilibrium distribution of electrons, but the charge distribution oscillates about equilibrium for a number of cycles. These charge oscillations, called “plasmons,” are quantized in energy. Larger energy losses correspond to the excitation of more plasmons, not to an increase in the energy of a plasmon. In most EELS spectra, the majority of inelastic scattering events are plasmon excitations.

To find the characteristic oscillation frequency of a plasmon, consider the rigid translation of a wide slab of electron density by a small amount, $x$, as in Fig. 2.7. At the bottom surface of the slab all the electrons are removed, but at the top the electron density is doubled. This charge disturbance therefore sets up the electric field, $E$, of a parallel-plate capacitor:

$$E = 4\pi \sigma_s ,$$

where $\sigma_s$ is the surface charge density equal to the electron charge, $e$, times $\rho$, the number of electrons per unit volume, times the displacement, $x$:

$$\sigma_s = e \rho x .$$

The field, $E$, provides the restoring force to move the electron slab back to its original position. The electrostatic restoring force per unit volume of slab is:

$$F = -e\rho E .$$

Substituting (2.5) into (2.4), and then into (2.6) gives a restoring force linear in the displacement:

$$F = -e^2 \rho^2 4\pi x .$$
Figure 2.7: Displacement of a slab of electric charge, leading to doubling of the charge density at the top of the slab over thickness $x$, and depletion of charge at the bottom. A wide, flat slab idealizes the problem as one dimensional.

The Newtonian equation of motion per unit volume of the electron slab is:

$$F = \rho \frac{d^2 x}{dt^2}.$$  

(2.8)

Substituting (2.7) into (2.8) gives:

$$\frac{d^2 x}{dt^2} = - \left( \frac{4\pi e^2 \rho}{m_e} \right) x.$$  

(2.9)

Equation (2.9) is the equation of motion for an undamped harmonic oscillator with the characteristic frequency:

$$\omega_p = \sqrt{\frac{4\pi e^2 \rho}{m_e}} = 5.64 \times 10^4 \sqrt{\rho},$$  

(2.10)

where the units of $\rho$ are [electrons cm$^{-3}$], and $\omega_p$ is [Hz]. With analogy to a mechanical oscillator, the electron density provides the stiffness. The higher the electron density, the higher the plasmon frequency. For metals, assuming an approximate free electron density of $\rho = 10^{23}$ electrons cm$^{-3}$, $\omega_p \approx 2 \times 10^{16}$ Hz. The characteristic energy of such an oscillation is the plasmon energy, given by:

$$E_p = \hbar \omega_p,$$  

(2.11)

and for our example $E_p = (6.6 \times 10^{-16} \text{ eV s}) (2 \times 10^{16} \text{ s}^{-1}) \approx 13 \text{ eV}$.

In EELS, intense plasmon peaks are prominent at energy losses of 10–20 eV. Plasmons are not long-lived, however, often because they promote excitations of electrons near the Fermi energy. Plasmon peaks therefore tend to be broadened in energy. Free electron metals such as aluminum have sharper plasmon peaks than transition metals, which have a high density of states at the Fermi energy. Compared to core electron excitations, however, plasmon excitations do not provide much detailed information about the individual atom species in the material.

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3 The present one-dimensional approach is not necessarily reliable for relating the free electron density to the plasmon energy. A more general approach uses the imaginary component of the dielectric constant of the material, and the theory uses the mathematics discussed in Problem 5.6.

4 “Lifetime broadening” is understandable from the uncertainty principle: $\Delta E \Delta t \approx \hbar$. A short lifetime $\Delta t$ comes with a large uncertainty in energy, $\Delta E$. 


2.3. PLASMON EXCITATIONS

2.3.2 * Plasmons and Specimen Thickness

The characteristic length or “mean free path,” $\lambda$, over which a 100 keV electron excites one plasmon is about 100 nm in metals and semiconductors. This is an average length, so in a TEM specimen of even 50 nm, some electrons excite one, two, or more plasmons. The probability $P_n$ for the excitation of $n$ plasmons in a sample of thickness, $t$, is determined by the statistics of Poisson processes:

$$P_n = \frac{1}{n!} \left( \frac{t}{\lambda} \right)^n e^{-t/\lambda} = \frac{I_n}{I_t},$$

where $I_n$ is the number of counts in the $n$th plasmon peak, and $I_t$ is the number of counts in all plasmon peaks with $n \geq 0$ ($I_t$ includes the zero-loss peak for which $n = 0$). The EELS spectrum in Fig. 2.8 shows distinct plasmon peaks. After subtracting a background from other inelastic processes (originating with the Al L-edge and a contribution from oxides and the substrate), $P_n$ is obtained as the fractional area of the $n$th plasmon peak.

Good samples for TEM imaging are several times thinner than the sample used for Fig. 2.8, but plasmon peak areas still offer a practical way to determine thicknesses of thin samples. Setting $n = 0$ in (2.12), the thickness, $t$, is:

$$t = \frac{\lambda}{\ln \left( \frac{I_t}{I_0} \right)}.$$

Measurement of $I_t$ and $I_0$ (the zero-loss or the $n = 0$ plasmon peak) involves the choice of the energies $\epsilon$, $\delta$ and $\Delta$, which define the limits of integration, as illustrated in Fig. 2.9. The lower limit ($-\epsilon$) of the zero-loss region can be taken anywhere to the left of the zero-loss peak where the intensity has decreased to zero, the separation point $\delta$ between the zero-loss and inelastic

---

5If a collection aperture limits the angles recorded by the spectrometer to a maximum angle $\beta$, $\lambda$ in (2.13) must be interpreted as an effective mean free path, $\lambda(\beta)$. 
regions may be taken as the first minimum in intensity, and $\Delta \approx 100$ eV is usually sufficient to include most of the inelastic scattering in relatively thin, low $Z$ materials (for high $Z$ and/or thick specimens, several hundred eV should be used since the scattered intensity is shifted to higher energy loss). Equation (2.13) has been shown to give 10% accuracy for relative thickness measurements on samples as thick as $t = 5\bar{\lambda}$. Some deviations from the intensities of (2.12) are of course expected when the electron beam passes through regions of non-uniform thicknesses or composition, and when other contributions are present in the low-loss spectrum.

Absolute determinations of specimen thickness require values for the total inelastic mean free path. For materials of known composition, it is possible to calculate a value for the mean free path according to the semi-empirical equation [5.5]:

$$\bar{\lambda} \approx \frac{106 F E_0}{\ln(2\beta E_0/E_m) E_m},$$

(2.14)

where units for $\bar{\lambda}$ are [nm], $\beta$ is the collection semiangle [mrad], $E_0$ is the incident energy [keV], $E_m$ is a mean energy loss [eV] that depends on the composition of the sample, and $F$ is a relativistic factor:

$$F = \frac{1 + E_0/1022}{(1 + E_0/511)^2},$$

(2.15)

and $F = 0.768$ for $E_0 = 100$ keV and 0.618 for $E_0 = 200$ keV. For a specimen of average atomic number $Z$, $E_m$ can be obtained from the semi-empirical formula:

$$E_m \approx 7.6Z^{0.36}.$$  

(2.16)

For large collection apertures, i.e., $\beta > 20$ mrad for $E_0 = 100$ keV or $\beta > 10$ mrad at 200 keV, (2.14) becomes inapplicable and the mean free path saturates at a value independent of $\beta$.

Specimen thickness measurement by this EELS plasmon technique has some advantages over other techniques for measuring specimen thickness (such as CBED) because it can be applied for a wide range of specimen thicknesses, including very thin specimens, and for specimens that are highly disordered or amorphous. Table 2.1 lists some values for calculated (with (2.10) and (2.11)) and measured plasmon energies $E_p$, widths of the plasmon peaks $\Delta E_p$, characteristic scattering angles for plasmons $\phi_E$, and calculated mean free paths $\bar{\lambda}$, for 100 keV incident electrons.
### 2.4 Core Excitations

#### 2.4.1 Scattering Angles and Energies – Qualitative

When a high-energy electron undergoes inelastic scattering, its energy loss, $E$, is actually a transfer of energy to the sample. When this energy is transferred to an atomic electron, the atomic electron may find an unoccupied electron state about the same atom, or it may leave the atom entirely (i.e., the atom is ionized). The total energy and the total momentum are conserved, but the scattering redistributes the energy and momentum between the high-energy electron and the atomic electron. The two electrons have coupled behavior. In particular, the probabilities and energies for the allowed excitations of the atomic electron are mirrored in the spectrum of energy losses of the high-energy electron. Relevant energies and their notation are listed in Table 2.2.

When a high-energy electron transfers energy to a core electron, the wavevector of the high-energy electron is changed in both magnitude and direction. The change in energy is obtained from the change in magnitude of the wavevector. The change in momentum is obtained from both the change in direction.
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Figure 2.10: Kinematics of inelastic electron scattering. (a) Definitions, with sphere of constant $E$. (b) Enlargement valid for small $\phi$, or equivalently for small $\Delta k$.

...and the change in magnitude of the wavevector. Total momentum is conserved, and before scattering the total momentum is with the incident electron, $p_0 = m_e v_0 = \hbar k_0$. After scattering, the momentum transfer to the atomic electron must be $\hbar \Delta k = \hbar (k - k_0)$. This same $\Delta \mathbf{k} \equiv \mathbf{k} - k_0$ is used for elastic scattering (Fig. 1.5), but inelastic scattering has an extra degree of freedom because $k \neq k_0$. Figure 2.10a shows that increasing $\phi$, the scattering angle, gives larger values of $\Delta k$ for the same $E$. Momentum conservation requires that the head of the wavevector $\Delta \mathbf{k}$ lies along the circle of radius $k$. The scattering vector, $\Delta \mathbf{k}$, can be zero only when both $\phi = 0$ and $E = 0$. When $E = 0$ but $\phi \neq 0$, $\Delta k$ cannot be zero – this is the case for elastic scattering in diffraction experiments.

We first consider general features of how inelastic scattering depends on $E$ and $\Delta k$. For $E$ only slightly larger than an absorption edge energy, $E_a$, the inelastic scattering is forward-peaked with a maximum intensity at the smallest $\Delta k$. Figure 2.10a shows that when $\phi = 0$ and $E \neq 0$, there is a nonzero minimum value of $\Delta k$, corresponding to inelastic scattering in the forward direction: $\Delta k_{\text{min}} \equiv (|k| - |k_0|)k_0$. In a particle model, these low-angle scatterings correspond to soft collisions with large impact parameters (meaning that the high-energy electron does not pass close to the center of the atom). The energy transfer is still large ($E \approx E_a$), unlike most soft classical collisions, but the outgoing core electron carries insignificant kinetic energy and $\Delta k$ is small. For small $\Delta k$, the scattering is sensitive to the large $r$ (long-range) features of the scattering potential.

On the other hand, at larger energy losses ($E \gg E_a$), the scattering is at higher angles, corresponding to hard collisions with small impact parameters. The outgoing core electron carries significant kinetic energy (equal to $E - E_a$), and...
2.4. CORE EXCITATIONS

the momentum transfer deflects the high-energy electron. For sufficiently large $E$, we expect the momentum transfer and energy transfer to be understandable by collisional kinematics, with little influence from the characteristics of the atom such as $E_a$. In fact, for larger energy transfers the inelastic intensity becomes concentrated around a specific value of $\Delta k$ such that:

$$\Delta k_B = \frac{\sqrt{2mE}}{\hbar}. \quad (2.17)$$

This peak in $\Delta k$ corresponds to the momentum transfer in classical “elastic” scattering of a moving ball (electron) by another ball initially at rest. This peak is called the “Bethe peak,” and in a two-dimensional plot of inelastic scattering intensity versus $\Delta k$ and $E$, these peaks become a “Bethe ridge” (cf., Fig. 2.11). Substituting into (2.17) a handy expression involving the Bohr radius, $a_0$, and the Rydberg energy, $E_R = \hbar^2(2ma_0^2)^{-1}$, we obtain for $\Delta k_B$:

$$(\Delta k_B a_0)^2 \approx \frac{E}{E_R}, \quad (2.18)$$

for which the equivalent scattering angle for the Bethe ridge, $\phi_r$, is:

$$\phi_r \approx \sqrt{\frac{E}{E_0}}. \quad (2.19)$$

The results of (2.17)–(2.19) are valid for small $\phi$ and non-relativistic electrons.

Experimentally, we count electrons. The energy spectrum of these electrons, $\rho(E) \, dE$, varies with solid angle, $\Omega$. With reference to Fig. 1.5, the three different $d\Omega_j$ will have different energy spectra. The most detailed experimental measurements would provide an energy spectrum at each differential solid angle, $d\Omega$. The number of electrons detected in a range $d\Omega$ around $\Omega$ and a range $dE$ around $E$ is proportional to the “double-differential cross-section,” $d^2\sigma / d\phi \, dE$.

In practice, there is often cylindrical symmetry around the forward beam, so we may need only the $\phi$-dependence (where $\phi = 2\theta$ in scattering angle). Experimental EELS spectra are measurements of intensity versus energy loss, $I(E)$, over a finite range of scattering angles, $\phi$.

Theoretically, we calculate the probability that a transfer of energy, $E$, and momentum, $\hbar \Delta k$, occurs between a high-energy electron and an atomic electron. To relate the theory to measured EELS spectra, we then need:

- The variation of the inelastic scattering over the parameter space of $(\phi, E)$. This is given by a double-differential cross-section, $d^2\sigma_{in} / d\phi \, dE$, described in Sect. 2.4.3 (d$\phi$ refers to rings of solid angle). This $d^2\sigma_{in} / d\phi \, dE$ includes the “generalized oscillator strength” of the specific atom.

- EELS spectra, $I(E)$, are measured over a range of $\phi$, so we need to integrate $d^2\sigma_{in} / d\phi \, dE$ over angle to obtain the differential cross-section, $d\sigma_{in} / dE$, described in Sect. 2.4.5.
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- Compositional analysis by EELS uses total intensities, given by the total cross-section, \( \sigma \) (or more typically by partial cross-sections, corresponding to a finite range in energy), as described in Sect. 2.4.6. This total probability for ionizing an atom is also needed for understanding EDS spectra, which measure x-ray emissions after the atom is ionized.

2.4.2 Inelastic Form Factor

Here we calculate the probability of an inelastic scattering process involving the excitation of a core electron. In this process, a high-energy electron excites a second electron from a bound atomic state into a state of higher energy. Since two electrons are involved, for conciseness we employ the Dirac bra and ket notation.\(^8\) The high-energy electron, “electron 1,” is initially in a plane wave state \( |k_0\rangle \), and after scattering it is in the state \( |k\rangle \). The atomic electron, “electron 2,” is initially in the bound state \( |\alpha\rangle \). After scattering, electron 2 is in the state \( |\beta\rangle \), which may be either a bound state that is initially unoccupied, or a spherical (or plane) wave state if electron 2 is ejected from the atom. For inelastic scattering, \( |k\rangle \neq |k_0\rangle \) and \( \alpha \neq \beta \). The Schrödinger equation with the initial state is written as:

\[
H_0 |\kappa_0, \alpha\rangle = (E_0 + E_\alpha) |\kappa_0, \alpha\rangle .
\] (2.20)

So long as the two electrons are far apart and therefore non-interacting, the two-electron system obeys the unperturbed Hamiltonian:

\[
H_0 = -\frac{\hbar^2}{2m_e} \nabla_1^2 - \frac{\hbar^2}{2m_e} \nabla_2^2 + V(r_2) .
\] (2.21)

The coordinates of the high-energy electron 1 are \( r_1 \), and the coordinates of the atomic electron 2 are \( r_2 \). With different coordinates, each Laplacian in (2.21) acts on only one of the two electrons, and the potential energy term involves only electron 2. In such problems we can express the initial state as a product of one-electron wavefunctions: \( |\kappa_0, \alpha\rangle = |k_0\rangle |\alpha\rangle \), and the final state as: \( |k, \beta\rangle = |k\rangle |\beta\rangle \).

When using a product wavefunction in (2.21), the factor for electron 2, \( |\beta\rangle \), is a constant under the operations of \( \nabla_2^2 \), and \( |k\rangle \) is a constant under the operations of \( \nabla_1^2 \) and \( V(r_2) \). A “constant factor” does not affect the solution of the Schrödinger equation for the other wavefunction of the product. The Hamiltonian of (2.21) is therefore equivalent to two independent Hamiltonians for two independent electrons. This is as expected when the two electrons have no interaction.

\(^8\)Recall that Dirac notation is free of spatial coordinates and explicit functional forms of wavefunctions, but these are obtained with the position operator for coordinate set 1, \( \hat{r}_1 \), as: \( \hat{r}_1 |k\rangle = \psi(\hat{r}_1) \).

Actual evaluations of integrals require expressions such as: \( \langle \alpha | H | \beta \rangle = \int \psi_\alpha^* \hat{H} \psi_\beta \, d^3r \). When \( |\alpha\rangle \) is an eigenstate of \( H_1 \), \( \langle \alpha | H | \alpha \rangle = E_\alpha |\alpha\rangle = E_\alpha \), since the state functions are normalized. State functions are orthonormal, so \( \langle \alpha | \beta \rangle = 0 \) and \( \langle \alpha | \alpha \rangle = 1 \).
2.4. CORE EXCITATIONS

As the high-energy electron approaches the atom, we must consider two perturbations of our two-electron system. One perturbation is the Coulombic interaction of electron 2 with the electron 1, which is 
\[ \frac{e^2}{\left| r_1 - r_2 \right|} \]. The second perturbation is the interaction of the high-energy electron 1 with the potential from the rest of the atom, \( V(r_1) \). The perturbation Hamiltonian, \( H' \), is:

\[ H' = \frac{e^2}{\left| r_1 - r_2 \right|} + V(r_1). \] (2.22)

This perturbation \( H' \) couples the initial and final states of the system. The stronger the coupling, the more probable is the transition from the initial state \( |\alpha\rangle \) to the final state \( |\beta\rangle \). It is a result from time-dependent perturbation theory that the wavefunction of the scattered electron 1 is an outgoing spherical wave times the form factor, \( f(\vec{k}, \vec{k}_0) \) (cf., (1.50)), where:

\[ f(\vec{k}, \vec{k}_0) = -\frac{me}{2\pi\hbar^2} \langle \beta | \left\{ \frac{1}{\left| r_1 - r_2 \right|} \right. \langle \vec{k} | H' | \vec{k}_0 \rangle | \alpha \rangle . \] (2.23)

Substitution of (2.22) into (2.23) gives:

\[ f(\vec{k}, \vec{k}_0) = -\frac{me}{2\pi\hbar^2} \left[ e^2 \langle \beta | \left( \frac{1}{\left| r_1 - r_2 \right|} \langle \vec{k} | H' | \vec{k}_0 \rangle | \alpha \rangle \right. \right. \\
+ \left. \left. \langle \beta | V(r_1) | \vec{k}_0 \rangle | \alpha \rangle \right] . \] (2.24)

When evaluating the second term of (2.24), the coordinates of electron 2 appear only in the atomic wavefunctions \( |\alpha\rangle \) and \( |\beta\rangle \), so these wavefunctions are moved out of the integral involving the coordinates of electron 1:

\[ f(\vec{k}, \vec{k}_0) = -\frac{me}{2\pi\hbar^2} \left[ e^2 \langle \beta | \left( \frac{1}{\left| r_1 - r_2 \right|} \langle \vec{k} | H' | \vec{k}_0 \rangle | \alpha \rangle \right. \right. \\
+ \left. \left. \langle \beta | V(r_1) | \vec{k}_0 \rangle | \alpha \rangle \right]. \] (2.25)

For inelastic scattering we have \( \alpha \neq \beta \), so the second term\(^\text{10}\) is zero by the orthogonality of the atomic wavefunctions. To be explicit in notation, we denote the inelastic contribution to \( f(\vec{k}, \vec{k}_0) \) as \( f_{\text{in}}(\vec{k}, \vec{k}_0) \), and call it the “inelastic form factor.” To calculate \( f_{\text{in}}(\vec{k}, \vec{k}_0) \), we use spatial coordinate representations.

\(^9\)For the potential from the rest of the atom, we could use the potential of an atom without a core electron, since we consider electron 2 separately.

\(^{10}\)For elastic scattering there is no transfer of energy from the high-energy electron (electron 1) to the atomic electron (electron 2), so \( \alpha = \beta \). By the orthonormality of the atomic wavefunctions we know that \( \langle \alpha | \alpha \rangle = 1 \), so this second term is nearly equal to the right hand side of (1.76). The difference is that the scattering potential from electron 2 is considered separately as the first term in (2.23), but together the two terms in (2.25) account for the scattering from the entire atom.
for our wavefunctions. The non-zero first term of (2.25) is:

\[
\psi^*_{\beta}(\vec{r}_2) \psi_{\alpha}(\vec{r}_2) d^3 r_2 \ .
\]

We change variables: \( \vec{r}' \equiv \vec{r}_1 - \vec{r}_2 \) (so \( \vec{r}'_1 = \vec{r} + \vec{r}_2 \)), and \( \vec{\Delta k} \equiv \vec{k} - \vec{k}_0 \), and separate the integrations:

\[
\psi^*_{\beta}(\vec{r}_2) \psi_{\alpha}(\vec{r}_2) d^3 r_2 .
\]

Equation (2.27) shows that the only dependence of \( f_\in \) on \( \vec{k} \) and \( \vec{k}_0 \) is through their difference, \( \vec{\Delta k} \). The integral over \( \vec{r}' \) is \( 4\pi/\Delta k^2 \) (A.27):

\[
\psi^*_{\beta}(\vec{r}_2) \psi_{\alpha}(\vec{r}_2) d^3 r_2 .
\]

This inelastic form factor, \( f_\in(\vec{\Delta k}) \), is the amplitude of the outgoing high-energy electron wavefunction along the direction \( \vec{k} = \vec{k}_0 + \vec{\Delta k} \) when the high-energy electron excites the atomic transition \( \psi_\alpha \rightarrow \psi_\beta \). The inelastic form factor has many similarities to the elastic form factor, \( f_\el(\vec{\Delta k}) \), of (1.76). Specifically, the second term of (??) for \( f_\el(\vec{\Delta k}) \), which describes elastic scattering from the atomic electron density, \( \rho(\vec{r}) \), has the same form as (2.28). It is convenient to think of both the inelastic and elastic form factors in a common way. Along the direction \( \vec{k} = \vec{k}_0 + \vec{\Delta k} \), wavelets are emitted from all sub-volumes, \( d^3 r_2 \), of the atom. Each wavelet has a relative phase \( e^{-i\vec{k} \cdot \vec{r}_2} \), and its amplitude for elastic scattering is proportional to an electron density. The full wave is the coherent sum (integration) of wavelets from all volumes of the atom, weighted by an electron density. For elastic scattering the electron density is the usual electron density, \( \rho(\vec{r}) = \psi^*_\alpha(\vec{r}) \psi_\alpha(\vec{r}) \). For inelastic scattering, however, this “density” is the overlap of the initial and final wavefunctions, \( \rho'(\vec{r}) = \psi^*_\beta(\vec{r}) \psi_\alpha(\vec{r}) \).

Note the common prefactors of \( f_\el(\vec{\Delta k}) \) of (??) and \( f_\in(\vec{\Delta k}) \) of (2.28). Recall that the factor of \( \Delta k^{-2} \) originates with the Fourier transform of the Coulomb potential (??). Using the definition of the Bohr radius, \( a_0 = \hbar^2/(me^2) \), this prefactor is \( 2/(a_0 \Delta k^2) \), which has dimensions of length. We now obtain the
2.4. CORE EXCITATIONS

differential cross-section for inelastic scattering, $d\sigma_{\text{in}}/d\Omega$, as $f_1^* f_2$ (1.21)\(^{11}\):

$$
\frac{d\sigma_{\text{in}}(\Delta k)}{d\Omega} = \frac{4}{a_0^2 \Delta k^4} \left| \int_{-\infty}^{+\infty} e^{-i\Delta \vec{k} \cdot \vec{r}} \psi_\beta^*(\vec{r}) \psi_\alpha(\vec{r}_2) d^3\vec{r}_2 \right|^2. \quad (2.29)
$$

Although energy is transferred from the high-energy electron 1 to the atomic electron 2, the total energy is conserved. In the transition $|\tilde{k}_0\rangle |\alpha\rangle \rightarrow |\tilde{k}\rangle |\beta\rangle$, the total energy before scattering equals the total energy after scattering:

$$
E_0 + E_\alpha = (E_0 - E) + E_\beta, \quad (2.30)
$$

$$
E = E_\beta - E_\alpha = E_{\alpha\beta}. \quad (2.31)
$$

A spectrum of electron energy losses shows enhanced intensity when $E = E_{\alpha\beta}$. Owing to the Pauli principle, however, the state $\psi_\beta$ must be initially empty for it to be allowed as a final state for electron 2. The EELS spectrum usually shows a jump in intensity, or “edge jump,” when $E_{\alpha\beta} = E_\alpha$, where $E_\alpha$ corresponds to the lowest energy of an unoccupied state $\psi_\beta$. Enhanced intensity extends for $E > E_\alpha$, because other unoccupied states of higher energy are available to the atomic electron 2.

With actual wavefunctions for $\psi_\alpha$ and $\psi_\beta$, we could use (2.29) to calculate the strength of this inelastic scattering,\(^{12}\) and the measured intensity of the electron energy-loss spectrum at the various energies $E_{\alpha\beta} > E_\alpha$. To do this, however, we must first relate the experimental conditions to the cross-section of (2.29). Specifically, we need to know how experimental detector angles, $\phi$, select $\Delta k$ at various $E$. This is the topic of the next subsection.

2.4.3 * Double-Differential Cross-Section, $d^2\sigma_{\text{in}}/d\phi dE$

In EELS, we measure the spectrum of energy losses from electrons in some range of $\Delta k$, set by the angle, $\beta$, of a collection aperture (see Fig. 2.1). To understand the intensity of core-loss spectra, we need to know how the inelastic scattering depends on both scattering angle, $\phi$, and energy loss, $E$. This dependence of the intensity on $\phi$ and $E$ is provided by a double-differential cross-section, $d^2\sigma_{\text{in}}/d\phi dE$. We start with the $\phi$-dependence for fixed $E$. For small $\Delta k$ we can approximate, as shown in Fig. 2.10b:

$$
\Delta k^2 = k^2 \phi^2 + \Delta k_{\text{min}}^2. \quad (2.32)
$$

\(^{11}\)A correction factor at high energy losses accounts for how the outgoing flux of scattered electrons is reduced when the electron is slowed (cf. (1.15)), but we safely ignore this effect for energy losses of a few keV.

\(^{12}\)There is a subtle deficiency of (2.28) and (2.29). The excitation of a core electron changes the electronic structure of the atom. It is not necessarily true that atomic wavefunctions are appropriate for $\psi_\alpha$ or $\psi_\beta$ when a core hole is present. The atomic electrons change their positions somewhat in response to the core hole, so the second term in (2.25) may not be strictly zero by orthogonality.
The increment in solid angle covered by an increment in $\phi$ (making a ring centered about $k_0$) is:

$$d\Omega = 2\pi \sin \phi \, d\phi .$$  \hspace{1cm} (2.33)

By differentiating (2.32) (for fixed $E$, $\Delta k_{\text{min}}$ is a constant):

$$\phi \, d\phi = \frac{\Delta k}{k^2} \, d\Delta k ,$$  \hspace{1cm} (2.34)

so for the small $\phi$ of interest:

$$d\Omega = 2\pi \frac{\Delta k}{k^2} \, d\Delta k .$$  \hspace{1cm} (2.35)

Substituting (2.35) into (2.29), and re-defining $\vec{r}_2 \rightarrow \vec{r}$, provides:

$$d\sigma_{\text{in}}(\Delta k) \, d\Delta k = \frac{d\sigma_{\text{in}} \, d\Omega \, d\Delta k}{d\Omega \, d\Delta k} = \frac{8\pi}{a_0^2 k^2 \Delta k^3} \left[ \int_{-\infty}^{+\infty} e^{-i\Delta k \cdot \vec{r}} \psi^*_\beta(\vec{r}) \psi_\alpha(\vec{r}) \, d^3 \vec{r} \right]^2 ,$$ \hspace{1cm} (2.36)

where the right-hand side is averaged for all $\Delta k$ of the detected electrons.

When $\psi_\beta$ is a bound state of the atom, (2.36) can be used directly to obtain an EELS intensity at the energy corresponding to the transition $\alpha \rightarrow \beta$. In the more typical case, $\psi_\beta$ lies in a continuum of states, such as free electron states when the atomic electron leaves the atom with considerable energy, or a band of unoccupied antibonding states for energies $E$ that are close to the absorption edge energy, $E_\alpha$. We then need to scale the result of (2.36) by the number of states in the energy interval of the continuum, which is $\rho(E) \, dE$. Here $\rho(E)$ is the “density of unoccupied states” available to the atomic electron when it is excited. Accounting for the density of states of $\psi_\beta$ gives the double-differential cross-section:

$$\frac{d^2\sigma_{\text{in}}(\Delta k, E)}{d\Delta k \, dE} = \frac{8\pi}{a_0^2 k^2 \Delta k^3} \rho(E) \left[ \int_{-\infty}^{+\infty} e^{-i\Delta k \cdot \vec{r}} \psi^*_\beta(\vec{r}) \psi_\alpha(\vec{r}) \, d^3 \vec{r} \right]^2 .$$ \hspace{1cm} (2.37)

The convention is to rewrite (2.37) to isolate the scattering properties of the atom. This is done by defining the “generalized oscillator strength,” GOS, or $G_{\alpha\beta}(\Delta k, E)$:

$$G_{\alpha\beta}(\Delta k, E) \equiv E_{\alpha\beta} \frac{2m_e}{\hbar^2 \Delta k^2} \left[ \int_{-\infty}^{+\infty} e^{-i\Delta k \cdot \vec{r}} \psi^*_\beta(\vec{r}) \psi_\alpha(\vec{r}) \, d^3 \vec{r} \right]^2 .$$ \hspace{1cm} (2.38)
2.4. CORE EXCITATIONS

Here $E_{\alpha\beta}$ is the difference between the energies of the states $\psi_\alpha$ and $\psi_\beta$. Using (2.38) in (2.37):

$$
\frac{d^2\sigma_{in}(\Delta k, E)}{d\Delta k dE} = \frac{2\pi\hbar^4}{\alpha_0^2 \mu_e^2 E_{\alpha\beta} T} \frac{1}{\Delta k} \rho(E) G_{\alpha\beta}(\Delta k, E) .
$$

(2.41)

To make connection to experimental EELS spectra, we convert the $\Delta k$-dependence of (2.41) into a dependence on the scattering angle $\phi$ of Fig. 2.10. We do so by arranging (2.34) as a relationship between $d\Delta k$ and $d\phi$, and substituting into (2.41):

$$
\frac{d^2\sigma_{in}(\Delta k, E)}{d\phi dE} = \frac{2\pi\hbar^4}{\alpha_0^2 \mu_e^2 E_{\alpha\beta} T} \frac{k_0^2 \phi}{\Delta k^2} \rho(E) G_{\alpha\beta}(\Delta k, E) .
$$

(2.42)

Figure 2.10b shows the definition of $\phi_E \equiv \Delta k_{\text{min}}/k_0$ and the approximation:

$$
\Delta k^2 \approx k_0^2 \left( \phi^2 + \phi_E^2 \right) ,
$$

(2.43)

from which we obtain a useful expression:

$$
\frac{d^2\sigma_{in}(\phi, E)}{d\phi dE} = \frac{2\pi\hbar^4}{\alpha_0^2 \mu_e^2 E_{\alpha\beta} T} \frac{\phi}{\phi^2 + \phi_E^2} \rho(E) G_{\alpha\beta}(\Delta k, E) .
$$

(2.44)

From Newtonian mechanics we would expect $\phi_E$, which is a ratio of $k$-vectors, to depend on the energies in the collision problem as $\sqrt{E/T}$. There is, however, a change in mass energy equivalent of the high-energy electron after scattering. This energy loss from the change in mass is significant, so the wavelength change is considerably smaller than the non-relativistic prediction. The result from relativistic kinematics is:

$$
\phi_E = \frac{E}{2\gamma^2 T} \approx \frac{E}{2E_0} .
$$

(2.45)

As an example, for the C K-edge at 284 eV in a 200 kV microscope $\phi_E = 0.7$ mrad.

2.4.4 * Scattering Angles and Energies – Quantitative

We revisit the angular dependence of the inelastic scattering. At lower energy losses ($E$ slightly larger than $E_a$), and at smaller scattering angles, the main angular dependence in (2.44) is from the Lorentzian factor, $(\phi^2 + \phi_E^2)^{-1}$, peaked

\[13\] For accuracy, we have written (2.41) with the incident kinetic energy, $T$, that differs from the incident energy, $E_0$, as:

$$
T \equiv \frac{1}{2} m_e v^2 = E_0 \frac{1 + \gamma}{2} \gamma^2 ,
$$

(2.39)
owing to the relativistic correction:

$$
\gamma \equiv \frac{1}{\sqrt{1 - (v/c)^2}} = 1 + \frac{E_0}{mc^2} ,
$$

(2.40)

($\gamma \approx 1.4$ for 200 keV electrons).
at $\phi = 0$, with $\phi_E$ (2.45) as the half-width of the angular distribution. (The factor $\phi$ in the numerator of (2.44) merely accounts for the larger radius of a ring at larger $\phi$.)

We first compare this characteristic angle for inelastic scattering, $\phi_E$, to the characteristic angle for elastic scattering, $\phi_0$. The elastic angle $\phi_0$ is associated with the atomic form factor, which is a measure of the size of the atom. For convenience we select $r_0$, the Bohr radius of the Thomas–Fermi atom as this size (??), and obtain $\phi_0$ as:

$$\phi_0 = \frac{1}{k_0 r_0}, \quad (2.46)$$

Putting typical values into (2.45) and (2.46), we find that $\phi_E$ is generally a few tenths of a milliradian while $\phi_0$ is a few tens of milliradians, i.e., $\phi_0 \approx 100\phi_E$. The inelastic scattering is concentrated into a much smaller range of angles about the forward beam than the elastic scattering, especially when $E \approx E_a$.

Section 2.4.1 discussed the other extreme case where $E \gg E_a$, and the collision kinematics are insensitive to the shape of the atom – recall that the intensity became bunched into angles characteristic of classical “billiard-ball” collisions.

The generalized oscillator strength, $G_{\alpha\beta}(\Delta k, E)$ of (2.38), helps complete the picture of how the inelastic intensity varies between these two extremes of $E \approx E_a$ and $E \gg E_a$. The generalized oscillator strength, $G_{\alpha\beta}(\Delta k, E)$, is the probability of the transition $\alpha \rightarrow \beta$, normalized by a factor related to the energy and momentum transfer. Figure 2.11 shows the GOS on the two-dimensional space of $\{\ln(\phi), E\}$ in a plot known as a “Bethe surface.” The individual curves in Fig. 2.11 show the angular dependence of the inelastic scattering for each energy loss above the carbon $K$-edge. Likewise, the energy dependence of the GOS may be obtained by taking sections through the Bethe surface at constant scattering angle. The Bethe ridge is marked on Fig. 2.11. Although distinct at large $E$, the Bethe peak is less well-defined at energy transfers closer to $E_a$ (the C $K$-edge threshold at 0 eV in Fig. 2.11).

In EELS measurements, an entrance aperture having an acceptance semi-angle $\beta$ is placed around the forward beam (Fig. 2.1). This aperture cuts off the scattering beyond a certain $\phi$. The measured spectrum of intensity versus energy is therefore an integration of the scattering intensity over combinations of $E$ and $\Delta k$ that fall below this cutoff. At energies significantly above the edge, Fig. 2.11 shows that a substantial portion of the intensity is concentrated in the Bethe ridge at larger scattering angles. A relatively large objective aperture (> 10 mrad or so) is needed to include this intensity in the EELS spectrum. On the other hand, at energies just above the edge, a small aperture will collect most of the intensity. This small aperture may be useful for removing background intensity at large $\Delta k$ that originates from tails of other elements with lower $E_a$.

2.4.5‡ * Differential Cross-Section, $d\sigma_{\text{in}}/dE$

Ignoring any truncation of the scattered inelastic intensity caused by the spectrometer entrance aperture, $\beta$, we integrate (2.44) over all possible scattering
angles, $\phi$, from 0 to $\pi$. This provides the total inelastic differential cross-section, $d\sigma_{in,\alpha\beta}(E)/dE$ for exciting an atomic electron from state $|\alpha\rangle$ to state $|\beta\rangle$:

$$\frac{d\sigma_{in,\alpha\beta}(E)}{dE} = \frac{2\pi\hbar^4}{\alpha_0^2 m_e^2 E_{\alpha\beta} T} \rho(E) G_{\alpha\beta}(\Delta k, E) \int_0^{\pi} \frac{\phi}{\phi^2 + \phi_E^2} d\phi .$$  \hspace{1cm} (2.47)

Here we have ignored the $\phi$-dependence of the GOS, $G_{\alpha\beta}(\Delta k, E)$. With the reasonable approximation that $\phi_E \ll \pi$, the integration of (2.47) gives:

$$\frac{d\sigma_{in,\alpha\beta}(E)}{dE} = \frac{\pi\hbar^4}{\alpha_0^2 m_e^2 E_{\alpha\beta} T} \rho(E) G_{\alpha\beta}(\Delta k, E) \ln \left( \frac{\pi^2}{\phi_E^2} \right) .$$ \hspace{1cm} (2.48)

With (2.48) and (2.45) we obtain the inelastic differential cross-section:

$$\frac{d\sigma_{in,\alpha\beta}(E)}{dE} = \frac{2\pi\hbar^4}{\alpha_0^2 m_e^2 E_{\alpha\beta} T} \rho(E) G_{\alpha\beta}(\Delta k, E) \ln \left( \frac{2\pi^2 T}{E} \right) .$$ \hspace{1cm} (2.49)

Figure 2.12 shows a plot of the energy-differential cross-section for K-shell ionization of C ($E_a = 284$ eV), calculated for different collection semiangles $\beta$ using hydrogenic wavefunctions.\textsuperscript{14} Logarithmic axes are used to illustrate the approximate behavior:

$$\frac{d\sigma_{in,\alpha\beta}(E)}{dE} \propto E^{-r} ,$$ \hspace{1cm} (2.50)

where $r$ is the downward slope in Fig. 2.12 and is constant over various ranges in energy loss. The value of $r$ depends on the size of the collection aperture.

\textsuperscript{14}A hydrogenic atom uses the wavefunctions of a hydrogen atom, but with radial coordinates rescaled by a larger nuclear charge. There are no electron-electron interactions for a hydrogenic atom, but analytical expressions for the wavefunctions are available.
For large $\beta$, when most of the inner-shell scattering contributes to the energy loss spectrum, $r$ is typically about 3 at the ionization edge, decreasing towards 2 with increasing energy loss. The asymptotic $E^{-2}$ behavior occurs because for $E \gg E_a$, practically all of the scattering lies within the Bethe ridge. It approximates Rutherford scattering from a free electron (??), for which $d\sigma_{\text{in},\alpha\beta}(E)/dE \propto \Delta k^{-4} \propto E^{-2}$.

For small $\beta$, $r$ increases with increasing energy loss, the largest value (just over 6) corresponding to large $E$ and small $\beta$. The breaks in slope in Fig. 2.12 correspond to the condition where $E$ is large enough so that the Bethe ridge moves to angles outside the collection aperture. It is usually important to avoid this transition in experimental practice because it complicates the $E$-dependence of the measured intensity. It may be a good idea to calculate $\phi_r$ with (2.19), and use a collection angle, $\beta$, a few times larger than this, as mentioned in the context of (2.53).

2.4.6 ‡ Partial and Total Cross-Sections, $\sigma_{\text{in}}$

In quantitative elemental analysis, the inelastic intensity measured with an aperture angle $\beta$ is integrated over an energy range of width $\delta$ beyond an absorption edge. Assuming a thin specimen with negligible multiple scattering, the integrated intensity above $E_a$, is:

$$I_a(E_a, \delta, \beta) = N I_0 \sigma_{\text{in},\text{a}}(E_a, \delta, \beta),$$

(2.51)

where $N$ is the number of atoms per unit specimen area, and $I_0$ is the integrated zero-loss intensity. In (2.51), the “partial cross-section” $\sigma_{\text{in},\text{a}}(E_a, \delta, \beta)$ is
the integral of (2.44) over the range of collection angle and energy:

$$
\sigma_{in,a}(E_a, \beta, \delta) = \int_0^{\beta} \int_{E_a}^{E_a + \delta} d^2 \sigma_{in}(\phi, E) \frac{dE}{d\phi} dE d\phi.
$$

(2.52)

For numerical integration of $d^2 \sigma_{in}(\phi, E)/dEd\phi$, it is sometimes convenient to use the power-law behavior of (2.50).

Figure 2.13 shows the calculated angular dependence of $K$-shell partial cross-sections for the first-row (second-period) elements. The figure illustrates the dependence of the cross-sections on collection angle $\beta$, incident electron energy $E_0$, and ionization energy $E_K$, for constant integration width $\delta$. The cross-sections saturate at large values of $\beta$, i.e., above the Bethe ridge angle, $\phi_r$, owing to the fall-off in $G_{\alpha\beta}(\Delta k, E)$ at large $\Delta k$. The median scattering angle (for energy losses in the range $E_a$ to $E_a + \delta$) corresponds to a partial cross-section equal to one-half of the saturation value, and is typically $5 \phi_E$, where:

$$
\langle \phi_E \rangle = \frac{E_K + \delta/2}{2\gamma T},
$$

(2.53)

with $\gamma \equiv (1 - v^2/c^2)^{-1/2}$. Figure 2.13 shows that the saturation cross-sections decrease with increasing incident electron energy, although the low-angle values increase. This is because a small collection aperture accepts a greater fraction of the scattering when the incident energy is high and the scattering is more strongly forward-peaked.

For a very large range of energy integration $\delta$, the partial cross-section becomes the “integral cross-section” $\sigma_{in,k}(E_K, \beta)$ for inner shell scattering up to $\beta$ and all permitted values of energy loss. By setting $\beta = \pi$, the integral cross-section becomes the “total cross-section” $\sigma_{in,k}(E_K)$ for inelastic scattering from the $K$-shell. An approximate expression for $\sigma_{in,k}(E_K)$ is the “Bethe asymptotic cross-section”:

$$
\sigma_{in,k}(E_K) = 4\pi a_0^2 N_K b_K \frac{E_K^2}{T_E K} \ln \left( \frac{c_K T}{E_K} \right),
$$

(2.54)

where $N_K$ is the number of electrons in the $K$-shell (2, but for the $L$ and $M$ shells this would be 8 and 18, respectively), $E_K \equiv h^2(2m_e a_0^2)^{-1}$, $b_K \approx f_K/N_K$, $c_K \approx 4E_K/\langle E \rangle$, where $f_K \approx 2.1 - Z/27$ is the dipole oscillator strength for $K$-shell ionization and typically $\langle E \rangle \approx 1.5E_K$. A similar expression, useful for calculating the integral cross-section as a function of collection angle $\beta$, is given in Problem 5.10.

Computer programs are available to calculate differential cross-sections for $K$, $L$ and $M$ shell ionizations using various atomic models [5.5]. Figures 2.14a and 2.14b compare experimental $N$-$K$ and $Cr$-$L$ edges to those calculated with the widely-used SIGMAK and SIGMAL programs of Egerton [5.5]. These programs calculate inelastic cross-sections for individual, isolated atoms with
hydrogenic wavefunctions. As shown in Fig. 2.14, their integrated intensities are generally reliable, but they cannot provide information about local chemical effects in the near-edge region of the spectrum as discussed in Sect. 2.2.3. Figure 2.14b shows that the L-shell calculation with the SIGMAL program is good on the average, but it cannot model the white line peaks at the edge onsets. It does estimate their average intensity, however, based on the number of unoccupied d-states of the element.

* Dipole Approximation and X-Ray Absorption Edges

For energy losses near the absorption edge (small $E - E_a$) where most of the intensity occurs with small $\phi$ and small $\Delta k$, it is sometimes convenient to use the “dipole approximation” for the integral in (2.38). The dipole approximation is obtained by approximating $e^{-i\Delta k \cdot \vec{r}} \approx 1 - i\Delta k \cdot \vec{r}$, and recognizing that the integration of the first term of 1, i.e., $\langle \beta | 1 | \alpha \rangle$, is zero owing to the orthogonality of $\psi_\alpha$ and $\psi_\beta$. The dipole approximation therefore amounts to replacing the factor $e^{-i\Delta k \cdot \vec{r}}$ in the integrand of (2.38) with the simpler factor $-i\Delta k \cdot \vec{r}$. Electric dipole radiation is the dominant transition process in EELS, but non-dipole transitions are observed at large $\Delta k$ when higher order terms must be considered in the expansion $e^{-i\Delta k \cdot \vec{r}} \approx 1 - i\Delta k \cdot \vec{r} - (\Delta k \cdot \vec{r})^2/2 + ...$

For atomic transitions induced by x-rays, the GOS for inelastic x-ray scatter-
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Figure 2.14: (a) Comparison of an experimental N K-edge and a hydrogenic fit to the edge using the SIGMAK program, and (b) comparison of an experimental Cr L_{2,3}-edge and a modified hydrogenic approximation to the edge using the SIGMAL program. After [5.6].

ing differs from (2.38) in that the exponential, \( e^{-\Delta \vec{k} \cdot \vec{r}} \), is replaced by the dipole operator, \( \vec{e} \vec{r} \). For small values of \( \Delta k \), the integral in (2.38) is identical for both electron and photon inelastic scattering, and x-ray and electron absorption edges look very similar. Although the dipole approximation provides the same selection rules for the allowed atomic transitions for both EELS and for inelastic x-ray scattering, the \( E \)-dependence of EELS spectra is significantly different from that for inelastic scattering spectra of photons. This difference originates from the nature of electron scattering by a Coulomb potential, whose Fourier transform causes (2.28) to decrease strongly with \( \Delta k \). Since large energy losses, \( E \), are associated with the larger \( \Delta k \), it becomes difficult to acquire EELS spectra at \( E > 4 \) keV. In practice, inelastic x-ray scattering, using a synchrotron radiation source for example, is performed for energies from 5–50 keV or so, whereas EELS experiments measure energy losses less than 5 keV.

2.5 Energy Dispersive X-Ray Spectrometry (EDS)

2.5.1 Electron Trajectories Through Materials

This section explains how high-energy electrons traverse thin TEM specimens and generate x-ray emissions from atoms. Some issues of instrumentation and artifacts are discussed. The following Sect. 2.6 describes procedures for quantitative analysis of x-ray spectra to obtain chemical concentrations in the sample. We first consider the trajectories of high-energy electrons through the sample because these paths determine where the x-rays come from. For thin samples, most electrons go straight through. Some electrons undergo high-angle deflections from elastic Rutherford scattering, so the electron beam broadens as it traverses the sample. Along an electron trajectory we need to know the probabilities, or at least the relative probabilities, that the electron
will ionize atoms of different types – this was the topic of Sect. 2.4. After an atom is ionized, it is important to know the probability that it will emit an x-ray, and the probability that the x-ray will leave the sample and be counted by the detector.

Large-angle scatterings of electrons are primarily elastic in origin,\(^\text{15}\) and occur when the high-energy electron passes close to an atomic nucleus. In these scatterings the shielding effects of the atomic electrons can be ignored, and the result is the Rutherford scattering cross-section, \(d\sigma_R/d\Omega\), of (??), written with \(\theta\equiv\phi\) as:

\[
\frac{d\sigma_R}{d\Omega} = \frac{Z^2e^4}{16\pi^2 \sin^4(\phi/2)}.\tag{2.55}
\]

Equation (2.55) is also useful for understanding the occurrence of electron backscattering from the sample. “Backscattered electrons” are defined as electrons scattered by angles so large that they reverse direction and go back out through the same surface they entered. Because of the \(T^{-2}\) dependence in (2.55), electron backscattering is relatively rare for electrons of several hundred keV passing through thin specimens.\(^\text{16}\)

Electron trajectories are typically calculated individually with a Monte Carlo algorithm. The computer code allows for random occurrences of scattering events, consistent with a user-specified density of nuclei of charge \(Ze\), electron energy, and Rutherford cross-section of (2.55). The electrons move along straight paths between these elastic collisions, which occur with randomness in the path length and scattering angle.

Along the straight paths between the Rutherford scattering events, the electron is assumed to lose energy at random to inelastic processes, both core excitations and plasmons. The core electron excitations are the ionization events that enable the subsequent emission of x-rays. Sections 2.4.1–2.4.6 described how the probability for ionizing an atom depends on the scattering angle, \(\phi\), and energy loss, \(E\), of the incident electron. To calculate x-ray emission, we need to integrate over all \(\phi\) and \(E\) to obtain a total cross-section for inelastic scattering by core electron ionization, \(\sigma_{\text{in}}\). This was obtained as (2.49) and (2.54), which depend on the energy of the incident electron as: \(1/\tau \ln(\tau)\), where \(\tau \sim 2\pi T/E_{\text{in}} \gg 1\). We therefore expect that as the high-energy electron loses energy in a thick specimen, the inelastic scattering events become more frequent, at least until its kinetic energy, \(T\), becomes too small. Monte Carlo codes have been developed to model electron trajectories in solids with all the physical phenomena mentioned in this section, and typical results from a Monte Carlo code in Sect. 2.4.4 showed that the electron energy-loss spectrum tends to be forward-peaked, especially at small energy losses, owing to the \(\phi\)-dependence of (2.44).

\(^{15}\)The discussion of ionization cross-sections in Sect. 2.4.4 showed that the electron energy-loss spectrum tends to be forward-peaked, especially at small energy losses, owing to the \(\phi\)-dependence of (2.44).

\(^{16}\)Backscattered electrons are much more common in scanning electron microscopy, which uses incident electrons of a few keV. Although these electrons tend to be multiply-scattered, backscattered electrons provide some chemical analysis capability to the SEM image; the factor of \(Z^2\) in (2.55) causes the backscattered electron image (BEI) to be brighter in regions containing heavier elements.
2.5. ENERGY DISPERSIVE X-RAY SPECTROMETRY (EDS)

Carlo simulations of electron trajectories (top), and assumed locations of x-ray emission (bottom). (In reality, most individual paths generate zero x-rays.) After [5.9].

Figure 2.15: Monte Carlo simulations of electron trajectories (top), and assumed locations of x-ray emission (bottom).

A schematic map of the electron trajectories in a thick bulk specimen is shown in Fig. 2.16a. The deep penetration and lateral broadening of high-energy electrons in bulk material causes the region of x-ray emission to be approximately 1 $\mu$m in diameter. This is a typical spatial resolution of an electron microprobe, for example. Specimens used in TEM may be only tens of nm in thickness, however. A thin specimen, as depicted in Fig. 2.16b, lacks the bulk of the material where most broadening of the electron beam occurs. Spatial resolution in an analytical TEM is therefore much better than in an electron microprobe.\(^{17}\) As a rule of thumb, the spatial resolution is significantly smaller than the width of the probe beam plus the thickness of the sample. Monte Carlo simulations that implement the model of elastic–inelastic scattering described in this section provide an approximation for the beam broadening, $b$, in [cm]:

$$b = 6.25 \times 10^5 \frac{Z}{E_0} \sqrt{\frac{\rho t^3}{A}}, \hspace{1cm} (2.56)$$

where $A$ is the atomic weight of the element [g/mole], $\rho$ is density [g cm$^{-3}$], $t$ is thickness [cm], and $E_0$ is incident energy [eV].

“Secondary electron” emission is especially important in scanning electron microscopy (SEM). A secondary electron is an electron that is weakly bound to the sample and is ejected with a few (at most tens of) electron volts of energy. Since these electrons have little energy, they can traverse only short distances through a material (less than about 100 Å), and therefore originate from the near-surface region. The detected secondary electrons are highly sensitive to surface topography, being more likely to emerge from the peaks than the valleys of the surface drawn in Fig. 2.17. Secondary electron imaging (SEI) is the main technique of SEM, and can be performed in much the same way in the TEM. The instrument is operated in scanning mode with a secondary electron detector attached to the microscope column as illustrated in Fig. 2.18. The number of secondary electrons emitted per incident electron is defined as the “secondary

\(^{17}\)On the other hand, x-ray emission from the large volume on the left of Fig. 2.16 provides much greater intensity. This high intensity, and the higher current of the incident electron beam, allow electron microprobes to use wavelength dispersive x-ray spectrometers, which have low collection efficiency but excellent energy resolution.
Figure 2.16: Differences in beam broadening in a bulk specimen (a), and a thin film (b). Part a shows regions of electron penetration, electron escape, and x-ray emission. For high-energy electrons, dimensions of regions of x-ray emission are typically a few microns, microns for backscattered electrons, tens of Å for secondary electrons. The larger dimensions do not exist for the thin specimen in b.
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2.5.4 Fluorescence Yield

After a core electron has been emitted from an atom, the ionized atom quickly decays from its excited state. This may occur by “radiative” or “non-radiative” processes, in which the atom emits either an x-ray or an Auger electron, respectively. Both processes (described in Sect. ??) compete for the atomic decay. For a K-shell ionization, for example, the “fluorescence yield,” \( \omega_K \), is defined as the fraction of decays that occur by the emission of a K-shell x-ray. A calculation of \( \omega_K \) requires knowledge of the relative rates of decay of the atom by Auger and by x-ray processes. The rate of x-ray emission is calculated for an electric dipole transition between the two atomic states of the atom, \( |\alpha\rangle \) and \( |\beta\rangle \). The x-ray emission rate is proportional to factors like \( |\langle \alpha | e | \beta \rangle|^2 \). The rate of Auger electron emission involves two electrons, and is calculated for a Coulomb interaction between them. The Auger electron emission rate is proportional to factors like \( |\langle k | (\beta | e^2 / (r_1 - r_2) | \alpha \rangle | \gamma \rangle|^2 \), where \( |\alpha\rangle \), \( |\beta\rangle \), and \( |\gamma\rangle \) are atomic states. The
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Figure 2.19: K-shell fluorescence yield of the elements. The difference, $1 - \bar{\omega}_K$, is the yield of Auger electrons. After [5.10].

state $|k\rangle$ is that of a free electron with the Auger energy (the difference in energy between the states $|\alpha\rangle$ and $|\beta\rangle$, minus the binding energy of state $|\gamma\rangle$). The fluorescence yield is the ratio of the x-ray rate to the total rate, where the total rate is the sum of x-ray plus Auger rates. Empirically, for a K-shell emission, $\bar{\omega}_K$ depends approximately on atomic number, $Z$, as:

$$\bar{\omega}_K = \frac{Z^4}{10^6 + Z^4}.$$  \hspace{1cm} (2.57)

Heavier elements tend to emit x-rays, and lighter elements tend to emit Auger electrons. The K-fluorescence yield of the elements is presented in Fig. 2.19. The fluorescence yield increases rapidly with $Z$. On the other hand, the K-shell ionization cross-section decreases strongly with $Z$. This decrease in total ionization cross-section, denoted $Q_k$ but equal to $\sigma_{in}$ of Sect. 2.4.6, can be obtained from (2.54), or can be calculated with actual wavefunctions as in (2.38) (substituted into (2.44) and (2.52)). It also can be approximated as $Z^{-4}$. This $Z$-dependence of $Q_k$ is opposite to that of $\bar{\omega}_K$ in (2.57). The probability of generating an x-ray depends on the product of $\bar{\omega}_K$ and $Q_k$, and this product turns out to be relatively constant in the energy range from 1–20 keV. The EDS method therefore has a well-balanced sensitivity to the elements from Na to Rh.

The detection of x-ray fluorescence radiation is the most widely-used technique for microchemical analysis in a TEM. A solid state detector, whose characteristics were described in Sect. ?? (Fig. ??), is positioned near the specimen.

\^Approximately, the Auger emission probability is independent of $Z$, whereas the x-ray emission probability increases strongly with $Z$. Unfortunately, it is generally impractical to use a TEM for chemical analysis by measuring the energies of Auger electrons. Auger electrons lose a significant fraction of their energy through nanometer distances in a material. Auger energies characteristic of atomic transitions are obtained only for those few atoms at the very surface of a sample. Unfortunately, the vacuum in a TEM is not particularly good, and the sample is heated under the electron beam. The surfaces of a TEM specimen quickly become contaminated, even if they are not oxidized already.
The energy spectrum of the x-rays emitted from the specimen is acquired in a multichannel analyzer (Sect. ??, Fig. ??). A typical EDS spectrum, in this case from SiC, is presented in Fig. 2.20. The widths of the peaks are set by the energy resolution of the detector, and not by the atoms in the specimen. The detector characteristics also affect the intensities of the peaks. Notice that in spite of the equiatomic stoichiometry of SiC, the intensity of the C peak in the spectrum is much less than that of the Si peak.

Factors for converting x-ray intensities to elemental concentrations are a necessary part of quantitative EDS measurements. Fortunately, the thinness of a TEM sample simplifies the conversion process – for a particular sample geometry these conversion factors can often be regarded as a set of constants (Sect. 2.6.1). Simple constants of conversion are not appropriate when there is significant x-ray absorption and secondary x-ray fluorescence events in the sample, as illustrated in Fig. 2.16a, and this is typically the case for measurements on bulk samples in an electron beam microprobe or a scanning electron microscope. The thinness of the TEM sample minimizes problems with x-ray absorption and fluorescence (as illustrated in Fig. 2.16b), and quantitation is often straightforward.

2.6 Quantitative EDS

2.6.1 Thin-Film Approximation

Cliff–Lorimer Factors

Microchemical analysis by EDS begins by removing the background from the measured x-ray spectrum. The background originates primarily from bremsstrahlung radiation, which we found in Sect. ?? to depend weakly on energy, especially for thin specimens where multiple scatterings of the high-energy electron are unlikely. In the analysis of an EDS spectrum such as that in Fig. 2.21, a power series in \( E \) is typically used to model the background. With two or more adjustable parameters, the background can be modeled well. Subtracting the background from the spectrum provides peaks that can be either integrated numerically (with the procedure of Fig. ??), or fit to analytical functions such as Gaussian functions. The peak areas can be treated individually,
and this would be acceptable in the simple case of Fig. 2.21. When overlaps of peaks occur, it is preferable to work with sets of peaks (such as $K\alpha$, $K\beta$, L-series, etc.) with the energies and relative intensities expected for each element (including the sensitivity of the EDS spectrometer). Either method provides a set of peak intensities, $\{I_j\}$, where $j$ denotes a particular chemical element. These $\{I_j\}$ are converted to a set of elemental concentrations, $\{c_j\}$, as described next.

In thin foil specimens, it is unlikely that an x-ray emitted from one atom will be absorbed by a second atom (cf., Fig. 2.16). Such double-scattering processes are neglected in the “thin-film approximation.” This simplifies enormously the task of determining the $\{c_j\}$ from the $\{I_j\}$. In the thin-film approximation, the ratio of x-ray peak intensities from the elements A and B, $I_A/I_B$, is simply proportional to the corresponding weight-fraction ratio, $c_A/c_B$:

$$\frac{c_A}{c_B} = k_{AB} \frac{I_A}{I_B},$$  \hspace{1cm} (2.58)

where $k_{AB}$ is a constant for a given accelerating voltage and a specific EDS spectrometer, and is independent of specimen thickness and composition. This constant $k_{AB}$ is often called a “Cliff–Lorimer factor.” It accounts for the efficiency of x-ray production for different accelerating voltages and the efficiency of the detector at the relevant x-ray energies.

A convenient feature of EDS is that for a large number of elements, the $k_{AB}$ factor for their $K\alpha$ x-ray peaks is approximately 1. The ratio of the peak intensities (or even peak heights) therefore gives a good approximation of the sample composition, making for a simple, semi-quantitative EDS analysis. This approximation holds for elements from about Mg to Zn. Below or above this range of atomic numbers the $k_{AB}$ factor gradually increases, but for elements of similar atomic number it is still reasonable to estimate their concentrations by comparing intensities of their $K\alpha$ peaks.

A normalization procedure:

$$\sum_j c_j = 1,$$  \hspace{1cm} (2.59)
is used to convert the ratios of the weight fractions to weight percentages (or, alternatively, atomic fractions to atomic percentages). That is, if $k_{AB}$ for elements A and B in a binary system is known, quantification is based on the measured ratio of $I_A$ and $I_B$ (2.58), and using (2.59):

$$c_A + c_B = 1 \quad (2.60)$$

For a ternary system with elements A, B and C, the following equations are used:

$$\frac{c_C}{c_A} = \frac{k_{CA} I_C}{I_A}, \quad (2.61)$$

$$\frac{c_C}{c_B} = \frac{k_{CB} I_C}{I_B}, \quad (2.62)$$

$$c_A + c_B + c_C = 1 \quad (2.63)$$

For a ternary alloy we have one more unknown, but one more independent peak ratio and another equation (2.62). In general, as we add more elements we can still use a set of linear equations like (2.61) and (2.62), plus (2.59) to complete the alloy chemistry.

The Cliff-Lorimer factors are mutually related. This is seen by dividing (2.62) by (2.61):

$$\frac{c_A c_C}{c_C c_B} = \frac{k_{CB} I_C I_A}{k_{CA} I_B I_C}. \quad (2.64)$$

By the definition in (2.58), $k_{CA} = 1/k_{AC}$, so:

$$\frac{c_A}{c_B} = k_{AC} \frac{k_{CB}}{k_{CA}} \frac{I_A}{I_B}. \quad (2.65)$$

Comparing (2.58) and (2.65), we obtain a general relationship between the Cliff-Lorimer factors:

$$k_{AB} = k_{AC} k_{CB} \quad (2.66)$$

Cliff-Lorimer factors, or “$k$-factors,” are often stored in a look-up table in the EDS software.

**k-Factor Determination**

Considerable effort is devoted to obtaining accurate Cliff-Lorimer factors, $k_{AB}$, since the accuracy of the EDS analysis depends on them. The $k$-factors are a combination of specimen and detector properties. Consider a $k_{AB}$ coefficient for $K_\alpha$ x-ray emission from elements A and B. The thin film approximation assumes both types of x-rays originate in the same region, and take direct paths through the specimen. We therefore expect the $k_{AB}$ coefficient to be the ratio:

$$k_{AB} = \frac{A_A \bar{\sigma}_B \bar{\theta}_B Q_{KB} \epsilon^{(K_\alpha^A - K_\alpha^B) y}}{A_B \bar{\sigma}_A \bar{\theta}_A Q_{KA}}, \quad (2.67)$$
where $A_i$ is the atomic weight of element $i$, (needed when the $k_{AB}$ are for determining mass fractions), $\alpha_i$ is its fluorescence yield, $a_i$ is its fraction of $K\alpha$ emission (for which $K\beta$ emission competes, but $a_i = 1$ for $Z < 19$), and $\mu_i^{\text{Be}}$ is the “effective” mass-absorption coefficient for the x-ray from element $i$ and the detector window of effective thickness $t$ (comprising, for example, the Be window, the Si dead layer, and the Au conductive film). The $Q_{Ki}$ are the $K$-shell ionization cross-sections (which could in principle be obtained from the total cross-section of (2.54), but better results are available).

There are essentially three ways to determine $k_{AB}$: 1) determine it experimentally using standards, 2) use values available in the literature, or 3) calculate it from first principles. The first method is the most reliable. Experimental $k_{AB}$ values are determined for a specific microscope, detector and operating conditions. Use of calculated and/or experimental $k_{AB}$ values from the literature is possible, but errors are expected owing to differences in the characteristics of the specimen, microscope, detector, and experimental geometry (including the tilt of the sample). The agreement between experimental and calculated $k_{AB}$ values is typically good to $5\%$ for $Z > 14$, and for these elements it is often sufficient to calculate $k_{AB}$ values for a given detector and accelerating voltage. The disagreement between theory and experiment for low $Z$ may be due to a combination of inadequate theory, absorption of low-energy x-rays within the specimen, contamination on the detector window, or the loss of light elements during electron irradiation. For routine analysis it is common to use the $k_{AB}$ values provided by the software of the EDS spectrometer system. Performing similar measurements on experimental standards of known composition can provide correction procedures to improve quantification for specimens of similar compositions.
Chapter 3

Inelastic X-Ray Scattering

Historically, inelastic x-ray scattering would be out of place for readers of this book. When x-ray wavelengths are of the order of atomic distances, their energies are far out of range of phonons, magnons, and their couplings to electronic excitations. This handy expression

$$\lambda[\text{Å}] = \frac{12.40}{E}[\text{keV}],$$

shows that x-rays with wavelengths $\lambda$ of atomic distances have energies $E$ of 10 keV or so. Nevertheless, since these x-rays can be monochromated by a factor of a several millions, their energy transfers can be used to study excitations of meV energies. Such monochromators and the third-generation synchrotrons to illuminate them are remarkable technical achievements.

These highly monochromatic x-rays have led to new inelastic x-ray spectroscopies, and some are directly complementary to the neutron methods that are the main topic of this book. The technical details do not translate from neutrons to x-rays quite so easily as the concepts involving the excitations. There is a huge body of research in inelastic x-ray scattering that does not have bearing on thermal excitations, except, perhaps, through the Debye-Waller factor that is usually considered a nuisance to be avoided. To work with inelastic x-ray scattering studies of phonons, it is necessary to have some knowledge of the vast range of modern x-ray spectroscopies if only because these methods invariably compete with the high resolution methods. A better justification is that these modern methods of x-ray scattering also provide key insights into the underpinnings of the electronic and atomic phenomena that underlie thermal excitations.

A deeper difference between the scattering of high energy (10 keV) x-rays and thermal neutrons (10 meV) is in the quantum mechanics of the scattering. Both are quantum mechanical, of course. X-rays probe the electronic structures of materials and molecules, and with coincidence methods it is possible to study electron correlations in state occupancies.
Energy resolution gives a time scale through the uncertainty relationship

$$\Delta t = \frac{\hbar}{\Delta E}$$,

$$\Delta t [s] = 6.58 \times 10^{-16} [\text{eV}]$$  \hspace{1cm} (3.2)

and two examples are illustrative:

- First consider an energy resolution of 658 eV, giving $$\Delta t = 10^{-18} \text{ s}$$. This is the time it takes for a near-relativistic electron to move across an atom. It is a lower bound on the time required for atomic electrons to rearrange around a positive hole that was left after a core electron was removed by ionization.

- Now consider an energy resolution of 0.66 meV, which would be a good resolution for phonon spectroscopy. Here $$\Delta t = 10^{-12} \text{ s}$$, which is approximately the frequency of a phonon.

In the first case the interesting quantum mechanics is found in the time sequence of how an emission from the atom might occur after, during, or even before the primary ionization. In the second case, quantum mechanics forbids the knowledge of when the excitation occurs in the phonon vibrational period. Another consequence is that x-ray scattering sees atom vibrations as snapshots of configurations of displaced atoms, whereas the slower neutron scattering probes atom vibrations over full vibrational cycles. In this slower regime, the momentum transfer of the scattering and the atomic displacements need to be handled with care because the momentum and position operators do not commute, leading to the quantum formalism of Chapter 7.

The present chapter provides some minimal explanations of x-ray spectroscopies, including methodologies that enable modern synchrotron experiments. Primary ionizations cause energy losses of the incident x-rays, which are the basis for absorption spectroscopies. Radiations emitted from the excited atoms give x-ray fluorescence spectroscopies, which have long been used for chemical analyses of materials. Modern synchrotron techniques offer improvements on these important tools, but now combinations of these two spectroscopies allow measurements of joint spectra with two energies, or measuring the temporal relations between these different x-ray processes. Fundamentals of these methods are given in the book by de Groot and Kotani [F. de Groot and A. Kotani, Core Level Spectroscopy of Solids (CRC Press, 2008).], and will only be sketched in the present chapter. Perhaps you should consider this chapter as a “traveler’s guide” to de Groot and Kotani. You should go there yourself if you are interested in this content.

### 3.1 X-Ray Fluorescence Spectrometries

Spectroscopy with ionizing x-rays or electrons starts with the primary ionization of an atom. This is the basis for some spectroscopies
3.1. X-RAY FLUORESCENCE SPECTROMETRIES

- in x-ray absorption spectroscopy (XAS), the energy of a monochromatic x-ray beam is tuned over a range of ionization energies, and the intensity of the transmitted beam shows features where ionized atoms remove x-rays from the beam.

- in electron energy loss spectrometry (EELS), the incident electron beam is kept at one energy, but the measured spectrum of the transmitted beam shows features at energies where the beam loses energy from ionizations.

There are a number of features at energies close to the ionization threshold that lend specialized names to spectroscopies, such as near-edge spectroscopy (NEXS) for studies of chemical information, and extended structure away from the edge (EXAFS) that gives information on the atomic environment of the ionized atom. There are variants of names for transmission or backscatter radiation detection, and surface or bulk analyses.

The number of spectroscopy techniques expands considerably when the secondary processes are included. After a primary ionization, spectroscopies can be based on

- the emission of an x-ray, which is a standard way to identify chemical elements in a material; x-ray fluorescence spectroscopy (XRF)

- the emission of an electron, which has a precise kinetic energy that is set by the binding energies of three different electrons at the atom (Auger spectroscopy).

These basic emission processes are standard ones for materials characterization. The XRF technique is used with a variety of methods for ionizing the atom, such as incident x-rays from an x-ray tube, electrons in an electron microscope, or \( \gamma \)-rays from a radioisotope source.

Combinations of these primary and secondary spectroscopies can be made by simultaneously measuring pairs of the different processes above. This cooks up a rich alphabet soup of modern spectroscopy techniques, and some are presented in Fig. 3.1. New phenomena probed by these combinations include the intermediate states of the ionized atom. The time delay between the ionization and emission can explore electronic lifetimes in the atom. There may be resonances within the atom, and the letter “R” is included in the acronyms of these spectroscopies (sometimes “N” if not). Many of these methods demand high x-ray high intensity because they measure small parts of the emission spectrum at precise energies and angles. The RIXS (resonant inelastic x-ray spectroscopy) technique has evolved with the brilliance of modern synchrotron sources, for example. Interpreting the RIXS data, which are affected strongly by an unseen internal resonant electron state, requires serious quantum computations of the atomic physics.
3.1.1 Synchrotron Radiation

Storage Rings

Synchrotron radiation is a practical source of x-rays for many experiments that are impractical with the conventional x-ray sources. High flux and collimation, energy tunability, and timing capabilities are some special features of synchrotron radiation sources. Facilities for synchrotron radiation experiments are available at several national or international laboratories. These facilities are centered around an electron storage ring with a circumference of about one kilometer. The electrons in the storage ring have energies of typically $7 \times 10^9$ eV, and travel close to the speed of light. The electron current is perhaps 100 mA, but the electrons are grouped into tight bunches of centimeter length, each with a fraction of this total current. The bunches have vertical and horizontal spreads of nanometers.

The electrons lose energy by generating synchrotron radiation as their trajectories are bent. These energy losses are primarily in the electron mass, not velocity (which stays close to the speed of light), so the bunches remain intact. The electrical power needed to replenish the energy of the electrons is provided by a radiofrequency electric field. This cyclic electric field accelerates the electron bunches by alternately attracting and repelling them as they move through a dedicated section of the storage ring. (Each bunch must be in phase with the radiofrequency field.) The ring is capable of holding a number of bunches equal to the radiofrequency times the cycle time around the ring. For example, with a 0.3 GHz radiofrequency, an electron speed of $3 \times 10^5$ km/s, and a ring circumference of 1 km, the number of “buckets” to hold the bunches is 1,000.

Although the energy of the electrons in the ring is restored by the high power radiofrequency system, electrons are lost by occasional collisions with gas atoms in the vacuum. The characteristic decay of the beam current over several hours requires that new electrons are injected into the bunches.

As the bunches pass through bending magnets or magnetic “insertion devices,” their accelerations cause x-ray emission. X-ray emission therefore oc-

---

1 Three premier facilities are the European Synchrotron Radiation Facility in Grenoble, France, the Advanced Photon Source at Argonne, Illinois, USA, and the Super Photon Ring 8-GeV, SPring-8 in Harima, Japan [1.4].
curs in pulsed bursts, or “flashes.” The flash duration depends primarily on the length of the electron bunch, and may be 0.1 ns. In a case where every fiftieth bucket is filled in our hypothetical ring, these flashes are separated in time by 167 ns. Some experiments based on fast timing are designed around this time structure of synchrotron radiation.

**Undulators**

Synchrotron radiation is generated by the dipole bending magnets used for controlling the electron orbit in the ring, but all modern “third generation” synchrotron radiation facilities derive their x-rays from “insertion devices.” These are magnet structures such as “wigglers” or “undulators,” made with rows of magnets along the path of the electron beam. The fields of these magnets alternate up and down, perpendicular to the direction of the electron beam. Synchrotron radiation is produced when the electrons accelerate under the Lorentz forces of the row of magnets. The mechanism of x-ray emission by electron acceleration is essentially the same as that of bremsstrahlung radiation.

Because the electron accelerations lie in a plane, the synchrotron x-rays are polarized with $\vec{E}$ in this same plane and perpendicular to the direction of the x-ray.

Magnetic fields in an undulator are positioned precisely so that the photon field is built by the constructive interference of radiation from a row of accelerations. The x-rays emerge from the undulator in a tight pattern analogous to a Bragg diffraction from a crystal, where the intensity of the x-ray beam in the forward direction increases as the square of the number of coherent magnetic periods (typically tens). Again in analogy with Bragg diffraction, there is a corresponding decrease in the angular spread of the photon beam.

The relativistic nature of the GeV electrons is central to undulator design and operation for two reasons:

**•** Suppose that the magnetic fields cause oscillation of the electron in the vertical direction, out of the plane of the paper of Fig. 3.2. As the electron velocity $\vec{v}$ to the right approaches the speed of light, there is a rapid decrease of wavelength in the forward direction. The energy of the photon is enhanced by the relativistic factor $\gamma = \sqrt{1 - (v/c)^2}^{-1}$, where $v$ is the electron velocity and $c$ is the speed of light.

**•** The second relativistic effect is the shortening of the undulator, which is moving with negative velocity in the frame of the stationary electron. This contraction brings the poles of the undulator closer together by the factor $\gamma$.

With these two factors of $\gamma$, in the forward direction along the electron path, the electron oscillation frequency is enhanced by the factor $2(1 - (v/c)^2)^{-1}$. This factor is about $10^8$ for electrons of several GeV energy. Typical spacings of the magnets are 3 cm, a distance traversed by light in $10^{-10}$ sec. The relativistic enhancement brings the frequency to $10^{18}$ Hz, which corresponds to an x-ray
energy, $h\nu$, of several keV. The relativistic Lorentz contraction along the forward direction further the radiation pattern. The x-ray beam emerging from an undulator may have an angular spread of microradians, diverging by only a millimeter over distances of tens of meters. A small beam divergence and a small effective source area for x-ray emission makes an undulator beam an excellent source of x-rays for operating a monochromator.

**Free Electron Laser**

The electrons moving through the undulator have a bunch length that spreads over a magnetic period of the undulator or so. The electrons along the bunch are therefore undergoing accelerations that are not in phase with each other. The next advance beyond an undulator is controlling the electron groupings so they form “microbunches” within the main bunch. Electrons within each microbunch undergo accelerations in phase with each other, a process that is underlies the “free electron x-ray laser.” The microbunching is sustained by the x-rays themselves, which move along with the electrons at nearly the same velocity. The slight slippage of the electron phase from the x-ray phase can be used to advantage to stabilize the microbunches. Stability also depends on the loss of x-ray photons from the microbunch, which reduces its energy. Today the design of free electron x-ray lasers requires very long (>100 m) arrays of aligned magnet structures, and only one beam can be produced at a time.

Here is a brief summary of the intensities from these synchrotron and laser devices for x-rays emitted in the forward direction. This intensity depends on the number of periods in the magnet structure, $N$, and the number of electrons in each bunch (or microbunch) $N_e$, as:

- Bending magnet has no structure within its magnet, so $N = 1$, and its intensity $I \propto N_e$.
- Wiggler has a magnetic structure with $N$ alternating fields, and its intensity $I \propto N N_e$.
- Undulator has a relativistically-tuned magnetic structure so each electron...
emits in phase with itself through the magnetic accelerations, and its intensity $I \propto N^2 N_e$.

- Free electron laser has microbunches from which all electrons emit coherently, and its intensity $I \propto N^2 N_e^2$.

**Brightness**

Various figures of merit describe how x-ray sources provide useful photons. The figure of merit for operating a monochromator is proportional to the intensity (photons/s) per area of emitter (cm$^{-2}$), but another factor also must be included. For a highly collimated x-ray beam, the monochromator crystal is small compared to the distance from the source. It is important that the x-ray beam be concentrated into a small solid angle so it can be utilized effectively. The full figure of merit for monochromator operation is “brightness” (often called “brilliance”), which is normalized by the solid angle of the beam. Brightness has units of [photons (s cm$^2$ sr)$^{-1}$]. The brightness of an undulator beam can be $10^9$ times that of a conventional x-ray tube. Finally, the x-ray intensity is not distributed uniformly over all energies. The term “spectral brilliance” is a figure of merit that specifies brightness per eV of energy in the x-ray spectrum. The spectral brilliance of x-ray sources has improved remarkably over the past 50 years, and Fig. 3.3 shows the brightness of some x-ray beamlines.

Undulators are tuneable to optimize their output within a broad energy range. Their power density is on the order of kW mm$^{-2}$, and much of this energy is deposited as heat in the first crystal that is hit by the undulator beam. There are technical challenges in extracting heat from the first crystal of this “high heat load monochromator.” It may be constructed for example, of water-cooled diamond, which has excellent thermal conductivity.

**Beamlines and User Programs**

The x-ray monochromators, goniometers and detectors needed for synchrotron radiation experiments are located in a “beamline,” which is along the forward direction from the insertion device. These components are typically mounted in lead-lined “hutches” that shield users from the lethal radiation levels produced by the undulator beam.

Synchrotron radiation user programs are typically organized around beamlines, each with its own capabilities and scientific staff. Work at a beamline requires success with a formal proposal for an experiment. This typically begins by making initial contact with the scientific staff at the beamline, who can often give a quick assessment of feasibility and originality. Successful beamtime proposals probably will not involve measurements that can be performed with conventional x-ray diffractometers. Radiation safety training, travel arrangements, operating schedules and scientific collaborations are issues for

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2Brightness is also a figure of merit for specialized beamlines that focus an x-ray beam into a narrow probe of micron dimensions.
Figure 3.3: Spectral brilliance of x-ray sources. X-ray tubes are at bottom, with synchrotron beamlines at higher brilliance. Notice the range of $10^{22}$ on the y-axis.
3.1. X-RAY FLUORESCENCE SPECTROMETRIES

experiments at synchrotron facilities. The style of research differs considerably from that with instruments in a university laboratory.

3.1.2 X-Ray Core Electron Spectroscopies

There are numerous types of inelastic spectroscopies involving x-ray excitations of electrons in atoms. Figure 3.4 depicts four essential methods involving incident and outgoing x-rays. All four show a primary excitation of a core electron by an incident x-ray.

Figure 3.4a depicts how a spectrum of transmitted x-rays can be used in x-ray absorption spectroscopy (XAS) if the energy of incident x-rays can be varied. Alternatively, x-ray photoelectron spectroscopy (XPS) can measure an energy spectrum of outgoing electrons with incident x-rays of a single energy. In XPS, the incident x-ray exceeds the binding energy of an electron in the material, and the difference in energy becomes the kinetic energy of the outgoing photoelectron. An energy spectrum of the photoelectrons is typically recorded over a range of hundreds of eV. Detailed features in the spectra are washed out if the photoelectron is scattered on its way out of the sample, and the characteristic distance of scattering is only a nm or so. XPS is therefore a technique for surface science, probing the top several atomic layers, whereas XAS is a method more suited for bulk analysis.

Figure 3.4b shows a configuration similar to panel a, with a lower energy of the incident x-ray, $\hbar \Omega$. Just above the Fermi energy, or just above the highest occupied molecular orbital in chemical parlance, there are usually many states for electrons that reflect the potential energy of the material. An electron band may be partly full, for example, and the unfilled states that lie just above the Fermi level offer final states for the outgoing photoelectron when its kinetic energy is low. The absorption spectrum is stronger when more of these states are available, so x-ray absorption near-edge spectroscopy (XANES) is a method for probing the electronic structure of solids in the range of 50-100 eV above the Fermi level.

Figures 3.4c,d have similarities to Figs. 3.4a,b, but a second electronic level is involved, emitting an x-ray of energy $\hbar \omega$. Two x-ray energies are recorded, $\hbar \Omega$ and $\hbar \omega$, giving a spectral intensity in a two-dimensional plot. The cross-sections for these spectra involve consideration of intermediate states, such as for the electron that transitions into the core hole and emits the x-ray of energy $\hbar \omega$. Be forewarned, however, that the process is fast, and time uncertainty allows ambiguity in the time sequence of photoelectron emission from absorption of the incident x-ray of energy $\hbar \Omega$ and the emission of the x-ray with energy $\hbar \omega$.

Figures 3.4d relates to Fig. 3.4c in the same way as panel b relates to a. In Fig. 3.4d the probability of the outgoing photoelectron is influenced strongly by the presence of unoccupied states just above the Fermi level. The two-x-ray process of Fig. 3.4d is the basis for resonant inelastic x-ray scattering (RIXS). Quantitative interpretations of RIXS spectra generally require considerations of the electronic structure of the material, and cross-section calculations with second-order perturbation theory. This is described in Section ??.
Figure 3.4: X-ray spectroscopies based on primary ionizations (a and b) and with secondary processes (c and d). (a) The spectra of transmitted x-rays are used in XAS (x-ray absorption spectroscopy), and the electron spectrum is recorded for XPS (x-ray photoelectron spectroscopy). (b) Excitation of a core electron by an incident x-ray, but the incident x-ray has just enough energy to bring the core electron above the Fermi level. The x-ray spectrum reflects details of the unoccupied states above the Fermi level, and is called XANES (x-ray absorption near-edge spectroscopy). (c) The emission of an x-ray of energy $\hbar \omega$ after ionization by an incident x-ray of energy $\hbar \Omega$ gives NEXS (normal x-ray emission spectroscopy). (d) The emission of an x-ray of energy $\hbar \omega$ after ionization by an incident x-ray of energy $\hbar \Omega$ gives RIXS (resonant inelastic x-ray emission spectroscopy) when the incident x-ray is near the absorption edge, as in panel b.
Finally, it should be mentioned that time is an additional dimension in modern x-ray spectroscopies. A short burst of energy from x-rays or a laser puts the material in a transient state that is probed by subsequent emission of x-ray or electron. Sometimes this can be stimulated emission. There are numerous possible time-dependent x-ray spectroscopies, and their development is underway now.

### 3.1.3 Chemical Shifts of Core Levels

The simple hydrogen atom, with one electron and one nucleus, has orthogonal orbitals that are calculated as Laguerre polynomials in $r$ and spherical harmonics in $\theta, \phi$. The orthogonality of these hydrogenic wavefunctions does not survive the addition of other electrons, which change the potential energy and the shapes of the wavefunctions. Core electron polarization is an important phenomenon in atoms with multiple electrons. In particular, adding or removing an outer valence electron of an atom causes slight changes in energy, and altered wavefunctions of all the inner electrons of the atom. Precise calculations of these core electron polarizations are possible today without too much difficulty with "all electron" codes.

We can make a classical estimate the energy shift of a core electron by a rearrangement of an outer shell of charge. From electrostatics we know that a negative charge $Q$ distributed over a sphere of radius $R$ raises the potential for an electron within this sphere by the amount $E = +Q/(4\pi\mu_0 R)$. This would shift the energy of a core electron, but it would be a constant shift if $Q$ and $R$ were constant. What is more interesting is if there is a change in valence, and the charge $Q$ moves to nearby atoms, expanding from $R$ to $R + \Delta r$. There is a change in the potential seen by the core electron of

\[
\Delta E = \frac{1}{4\pi\mu_0} \left[ \frac{1}{R + \Delta r} - \frac{1}{R} \right], \quad (3.3)
\]

\[
\Delta E = \frac{1}{R 4\pi\mu_0} \left[ 1 - \Delta r - 1 \right], \quad (3.4)
\]

\[
\Delta E = -\frac{\Delta r}{R} \frac{1}{4\pi\mu_0}. \quad (3.5)
\]

The core electron levels become more negative (stronger bonding) when the outer electrons expand outwards, such as when transferred to neighboring atoms. The effect can be a few eV, and is readily studied by core electron spectroscopies.

### 3.1.4 Core Hole Lifetimes

After ionization, the core hole is highly unstable. A hole in the inner K-shell is filled with another electron in a time of order $10^{-16}$ s. Figure 3.5 shows core hole lifetimes for different chemical elements, with contributions from the
more constant Auger electron emission, dominant for lighter elements, and x-ray emission, which is faster and dominant for heavier elements.

### 3.1.5 Many-Body Effects

**Plasmons**

During the lifetime of a core hole of $10^{-16}$ s, a photon travels a distance of approximately 3 microns, which is large on the scale of atoms and local electronic excitations. A consequence is that the unstable states of ionized atoms can interact with other electronic degrees of freedom, such as plasmons.

A free electron gas has a uniform charge distribution in equilibrium, but a jolt of a quick change in charge can cause it to oscillate with the plasmon frequency. Typically the plasmons are at an energy of 10 eV, and are damped in 10 oscillations or so as they transfer their energy to other electronic excitations. The plasmon lifetime may be of comparable to the lifetime of the core excitation, and plasmons may become part of the x-ray absorption that generates the core hole. A consequence is that new features appear in x-ray absorption spectra at the combined energy of the plasmon plus core level. These are a common example of a “many body effect” that alters the shape of x-ray absorption spectra.

Plasmon excitations resolve a conceptual problem with core electron ex-
citations in simple metals. At low temperatures, a simple metal has a sharp
interface in energy between occupied and unoccupied electron states across the
Fermi level. The simple metal therefore has excitations that require an infinites-
imal amount of energy. The number of these excitations can lead to divergences
in the spectral intensity of many-body effects near absorption edges, broadening
the shapes of features from core edges. Quantizing the excitations of the
free electron gas into plasmons moves their many-body effects away from the
core edge features, and into distinct peaks some 10 eV away from the edge.

Shake-Up Satellites
Semiconductors and insulators have few or no states just above the Fermi
level. The creation of excitons can occur in semiconductors as part of the
ionization process, and the exciton energy can add a feature to the spectrum of
the core energy called a “shake-up” satellite, presumably because the electronic
disturbance of the core excitation shakes up the electronic degrees of freedom
including exciton states.

It should be added that other electron states in the ionized atom can con-
tribute features to the core energy spectrum. For example, in NXES of Fig. 3.4c
an electronic transition of energy $\hbar \omega$ participates in the core excitation process.
This is not called a many-body effect in the main absorption spectrum, but is a
method of spectroscopy in its own right, as described in the next section.

3.1.6 First- and Second-Order Spectroscopies
The purpose of this section is to give a conceptual approach to understanding
how NXES and RIXS differ from XAS and XANES. A second electronic transition
of energy $\hbar \omega$ is intrinsic to NXES in Fig. 3.1c that is not shown for XAS
in Fig. 3.1a. Both processes compete for scattering incident x-rays of energy $\hbar \Omega$, and it is a role of theory to calculate the relative cross-sections for the two
processes. Cross sections are not calculated in the brief survey here. The goal
is to understand a quantum mechanical difference between the two of them
using first- and second-order perturbation theory.

Perturbation theory is developed with a small parameter $\lambda$, which is used to
scale first order effects $e^0$ and second order effects $e^2$. The second-order effects
tend to be much smaller than first-order effects, and even more so as $\lambda$ becomes
smaller. Start with an unperturbed Schrödinger equation with pure electron
states denoted with a superscript $(0)$

$$H_0 \psi_i^{(0)} = E_0 \psi_i^{(0)}.$$  \hspace{1cm} (3.6)

Add a perturbation to the Hamiltonian $\lambda \delta$, which will be from the oscillating
electric field of an x-ray. A general expression for the Schrödinger equation is now

$$(H_0 + \lambda \delta) \left( \psi_i^{(0)} + \lambda \psi_i^{(1)} + \lambda^2 \psi_i^{(2)} \right)$$

$$= (E_i^0 + \lambda E_i^1 + \lambda^2 E_i^2) \left( \psi_i^{(0)} + \lambda \psi_i^{(1)} + \lambda^2 \psi_i^{(2)} \right)$$ \hspace{1cm} (3.7)
where we stopped at second powers in \( \lambda \). The terms in Eq. 3.7 with \( \lambda^0 = 1 \) give Eq. 3.6, which is also obtained when \( \lambda = 0 \). Collecting all the terms in \( \lambda^1 \) in Eq. 3.7 (and dividing by \( \lambda \)) gives

\[
H_0 \psi_i^{(1)} + \delta \psi_i^{(0)} = E_i^0 \psi_i^{(1)} + E_i^1 \psi_i^{(0)},
\]

(3.8)

\[
H_0 |i^{(1)}⟩ + δ|i^{(0)}⟩ = E_i^0 |i^{(1)}⟩ + E_i^1 |i^{(0)}⟩.
\]

(3.9)

where the change to Dirac notation allow cleaner expressions with the inner product of Eq. 3.9 with \( ⟨i^0 | H_0 |i^{(1)}⟩ + ⟨i^0 | δ|i^{(0)}⟩ = E_i^0 ⟨i^0 | i^{(1)}⟩ + E_i^1 ⟨i^0 | i^{(0)}⟩ \).

(3.10)

The first terms on each side are inner products of orthogonal functions, and by normalization \( ⟨i^0 | i^{(0)}⟩ = 1 \)

\[
E_i^1 = ⟨i^0 | δ|i^0⟩.
\]

(3.11)

3.1.7 * ZAF Correction

In an EDS spectrum, the x-ray peaks from different elements have intensities that depend on: 1) the path and energy of the high-energy electron passing through the sample, 2) the ionization cross-sections of the elements, 3) the fluorescence yields, and 4) the probabilities that emitted x-rays are seen by the detector. The thin film approximation collects all these effects into a constant factor for each type of characteristic x-ray. In the thin-sample limit, all peaks in an EDS spectrum increase in intensity with increased sample thickness, but the ratios of peak intensities remain unchanged. This permits the use of (2.58) for samples of all thickness. For thicker samples, however, the peak intensity ratios are altered. In TEM, the generation of characteristic x-rays from different elements is not altered by changes in the incident beam as it passes through a sample of moderate thickness. The thickness effects originate with the scattering of the characteristic x-rays by the different elements in the sample. As the samples become thicker and the x-ray exit paths through the sample become longer, these inelastic x-ray scattering processes involve a larger fraction of the x-rays, altering the ratios of peak intensities. Correction for these inter-element interactions is performed by considering the atomic number, \( Z \), the absorption, \( A \), and fluorescence, \( F \), in procedures called “ZAF corrections.”

* X-Ray Absorption Within the Specimen

X-ray absorption follows Beer’s Law (2.58). Since x-rays are generated throughout the foil thickness, evaluating the average absorption generally requires an integration of (2.58) over the sample thickness. Fortunately, for thin foils we can linearize the exponential in (2.58) as: \( e^{-x} \approx 1 - x \), and take the average depth of x-ray emission as \( t/2 \), where \( t \) is the sample thickness. In this case absorption
Table 3.1: Limits to the thin foil approximation caused by absorption. Thickness limit is for a 3% error in the $k_{AB}$ factor [5.9]

<table>
<thead>
<tr>
<th>Material</th>
<th>Thickness [nm]</th>
<th>Absorbed x-ray(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al–7%Zn</td>
<td>94</td>
<td>Al $K_{\alpha}$</td>
</tr>
<tr>
<td>NiAl</td>
<td>9</td>
<td>Al $K_{\alpha}$</td>
</tr>
<tr>
<td>Ag$_2$Al</td>
<td>10</td>
<td>Al $K_{\alpha}$, Ag $L_{\alpha}$</td>
</tr>
<tr>
<td>FeS</td>
<td>50</td>
<td>S $K_{\alpha}$</td>
</tr>
<tr>
<td>FeP</td>
<td>34</td>
<td>P $K_{\alpha}$</td>
</tr>
<tr>
<td>Fe–5%Ni</td>
<td>89</td>
<td>Ni $K_{\alpha}$</td>
</tr>
<tr>
<td>CuAu</td>
<td>11</td>
<td>Cu $K_{\alpha}$, Au $M_{\alpha}$</td>
</tr>
<tr>
<td>MgO</td>
<td>25</td>
<td>Mg $K_{\alpha}$, O $K_{\alpha}$</td>
</tr>
<tr>
<td>Al$_2$O$_3$</td>
<td>14</td>
<td>Al $K_{\alpha}$, O $K_{\alpha}$</td>
</tr>
<tr>
<td>SiO$_2$</td>
<td>14</td>
<td>Si $K_{\alpha}$, O $K_{\alpha}$</td>
</tr>
<tr>
<td>SiC</td>
<td>3</td>
<td>Si $K_{\alpha}$, C $K_{\alpha}$</td>
</tr>
</tbody>
</table>

alters the x-ray intensity ratio $I_A/I_B$ from the ratio recorded for an infinitely-thin specimen, $I_{A0}/I_{B0}$:

$$
\frac{I_A}{I_B} \approx \frac{I_{A0}}{I_{B0}} \left( \frac{1}{1 - \frac{\mu_A}{\rho_A} t \frac{1}{2} \rho_A \csc \psi} \right),
$$

(3.12)

$$
\frac{I_A}{I_B} \approx \frac{I_{A0}}{I_{B0}} \left( 1 + (\mu_B - \mu_A) \frac{t}{2} \csc \psi \right).
$$

(3.13)

Equation (3.13) shows the importance of the difference in absorption coefficients for the x-rays of elements A and B – if they have similar $\mu$, the intensity ratios $I_A/I_B$ are unaffected. Table 3.1 shows thicknesses at which the thin-film approximation is no longer valid due to absorption effects in specific materials.

* Characteristic Fluorescence Correction

Characteristic x-rays from a heavier element can photoionize atoms of lighter elements, causing them to fluoresce. This enhances the number of x-rays detected from the light element, and suppresses the number from the heavier element. Fluorescence effects in thin foils are much weaker than in bulk samples (Fig. 2.16). Nevertheless when strong fluorescence does occur, e.g., Cr $K_{\alpha}$ fluorescence under Fe $K_{\alpha}$ radiation, quantitative microchemical analysis of TEM specimens may require a fluorescence correction (Fig. 3.6). Several

---

*To make an absorption correction, however, it is necessary to know the mean x-ray path length within the specimen, and this is difficult to determine from wedge-shaped or irregular specimens.*
fluence corrections for thin films have been developed, and a successful model \[5.13\] uses an enhancement factor, \(X_A\), for the element, A, that undergoes fluorescence:

\[
X_A = c_B \tilde{\omega}_{KB} \frac{r_A - 1}{r_A} A_A \frac{A_B}{\mu_{BA}} \frac{U_B \ln U_B}{U_A \ln U_A} 2^\left(0.923 - \ln(\mu_{BA})\right) \tag{3.14}
\]

where \(U_i\) is the overvoltage ratio (ratio of incident electron energy to K-edge energy) for element \(i\), \(A_i\) is its atomic weight, \(c_i\) is its weight fraction, \(r_i\) is its absorption-edge jump-ratio (the fractional countrate change across the EELS absorption edge), \(\tilde{\omega}_{K_i}\) is a fluorescence yield, and \(\mu_{BA}\) is the mass absorption coefficient of element B in element A. For a sample in which the element B causes fluorescence of element A, the measured composition is corrected by:

\[
\frac{c_B}{c_A} = k_{BA} \frac{I_B}{I_A} (1 + X_A) \tag{3.15}
\]

3.1.8 * Limits of Microanalysis

There are three quantifiable limits to microanalysis: 1) the absolute accuracy of quantification, 2) the minimum detectable mass (fraction), and 3) the spatial resolution. Of course there are other practical limits including contamination, insensitivity to low Z in EDS, and specimen preparation and geometry, but here we discuss the first two quantifiable limits. Limits on spatial resolution were discussed in Sects. ?? and ??.

The accuracy of quantification is limited by the counting statistics of the x-ray spectra. For strong peaks on a weak background, the standard deviation, \(\sigma\), is given by:

\[
\sigma = \sqrt{N} \tag{3.16}
\]
where $N$ is the number of counts in the peak after background subtraction. Once the standard deviation is known, different confidence limits can be set for the value of $N$, i.e. 68% confidence that $N$ will lie in $N \pm \sigma$, 95% in $N \pm 2\sigma$ and 99% in $N \pm 3\sigma$. The value of $3\sigma$, taken to be the 99% confidence level in the value of $I_A$, is often used to estimate the error in the peak intensity:

$$\text{Error}(\%) = \pm \frac{3\sigma}{N} \times 100 = \pm \frac{3}{\sqrt{N}} \times 100.$$  

(3.17)

The larger is $N$, the lower the error in the analysis. For a 1% accuracy at the 99% confidence level, one needs $10^5$ counts in a peak, or $10^4$ counts for 1% accuracy at the 68% confidence level. The error in $I_A/I_B$ is:

$$\text{Error}(\%) = \pm \left( \frac{3}{\sqrt{N_A}} + \frac{3}{\sqrt{N_B}} \right) \times 100.$$  

(3.18)

When using (2.58) for composition analysis, to the error of (3.18) we must add any error in $k_{AB}$, which is again the sum of the errors in $I_A$ and $I_B$ for the standard.

If Gaussian statistics are assumed, there is a simple statistical criterion that can be used to define the minimum mass fraction (MMF). A peak containing $I_B$ counts from element B in a matrix of A is considered statistically real and not a random fluctuation in the background intensity, $I_B$, when:

$$I_B \geq 3\sqrt{2I_B}.$$  

(3.19)

The MMF of B that can be detected in a binary material of elements A and B, $c_B$(MMF) in at.%, is obtained using (2.58) and (3.19):

$$c_B(\text{MMF}) = 3\sqrt{2I_B} \frac{c_A k_{BA}}{I_A - I_B}.$$  

(3.20)

In practice, a MMF of approximately 0.1 wt.% can be obtained in EDS if enough counts are collected. Similarly, the minimum detectable mass (MDM) is predicted to be around $10^{-20}$ g for a range of $Z$ from 10 to 40.

These statistical analyses give the accuracy for quantification of a single measurement. In many cases, it is possible to obtain only a limited number of counts in a spectrum owing to factors such as beam damage or specimen drift. In such situations, it is possible to reduce the error in quantification (or at least assess it) by combining the results from $n$ different measurements of the intensity ratio $I_A/I_B$. The total absolute error in $I_A/I_B$ at a given confidence value is obtained using the Student-$t$ distribution. In this approach, the error of the estimate $E$ is given by:

$$E < \frac{t_{\alpha/2}S}{\sqrt{N}},$$  

(3.21)

When the background is a substantial fraction of the peak height, this argument is invalid for reasons stated in Problem 1.9. For weak peaks it is more accurate to use the background counts over the width of the peak, $N_B$, to obtain $\sigma = \sqrt{N_B}$ for use in (3.16).
where \( t_{\alpha/2} \) is the Student-\( t \) value such that the normal curve area to its right equals \( \alpha/2 \) with a probability of \( 1 - \alpha \), \( S \) is the standard deviation for \( n \) measurements of the intensity \( N_i \), given by:

\[
S = \sqrt{\frac{\sum_{i=1}^{n} (N_i - \langle N_i \rangle)^2}{n - 1}},
\]

which contain on average \( \langle N_i \rangle \) counts. By increasing the number of measurements, one can reduce the error of measurement. In other words, if we estimate \( \mu \) by means of a random sample size of \( n \), we can assert with a probability of \( 1 - \alpha \) (where \( 1 - \alpha = 0.95 \) for a 95% confidence level for example) that the error in the measurement \( E = \langle N_i \rangle - \mu \) is less than \( (t_{\alpha/2}S)/\sqrt{n} \), at least for sufficiently large values of \( n \). Equation (3.21) can also be rearranged and solved for \( n \) to determine the number of measurements \( n \) that must be taken to achieve a mean \( \langle N_i \rangle \) which is in error by less than \( E \).

### Further Reading

The contents of the following are described in the Bibliography.


Problems

4.1

Find the thickness (in nm) of the Al sample in the EELS spectrum of Fig. 3.7. Assume 100 keV electrons.

4.2

A TEM specimen undergoes a type of radiation damage known as “knock-on damage” when a high-energy electron transfers enough energy to an atom to displace it from its crystallographic site. For a given electron energy, knock-on damage tends to be most severe for elements of low atomic number.

(a) In a direct ("head-on") collision between a high-energy electron and an atom, show that the energy transfer scales inversely with the atomic weight of the atom. (For simplicity, you may assume that the incident electron is scattered elastically by an angle of 180°.)

(b) If a Li atom requires 10 eV to leave its crystal site, calculate the threshold energy for an incident electron to induce knock-on damage. Do the same calculation for Al, Cu, and Au.

4.3

This problem presents two mathematical tricks for working with Dirac δ-functions. Calculations of electronic energies or scattering intensities often employ sums of Dirac δ-functions because δ-functions are handy for representing energy eigenvalues. For example, it is possible to write a distribution
function for an energy spectrum, \( n(E) \), as:

\[
n(E) = \frac{1}{N} \sum_{\alpha} \delta(E - \varepsilon_{\alpha}) .
\] (3.23)

The idea behind this equation is that if \( N \) is large so there are numerous states (or transitions), each of energy \( \varepsilon_{\alpha} \), the discrete sum on the right becomes a continuum. To integrate the number of states up to some energy \( E' \), each \( \delta \)-function on the right side contributes 1 to the sum when \( E' > \varepsilon_{\alpha} \). A direct calculation of this type can be clumsy, however. Two expressions for \( \delta \)-functions can be of assistance in mathematical work:

\[
\delta(E - \varepsilon_{\alpha}) = \lim_{\delta \varepsilon \to 0} \frac{1}{\pi} \Im \left( \frac{1}{E + i\delta \varepsilon - \varepsilon_{\alpha}} \right) ,
\] (3.24)

\[
\delta(E - \varepsilon_{\alpha}) = \frac{1}{2\pi} \int_{-\infty}^{\infty} e^{i(E-\varepsilon_{\alpha})t} dt .
\] (3.25)

Prove, or convince yourself, that these two equations are appropriate ways to represent a \( \delta \)-function.

4.4

Suppose that samples containing mixtures of elements A,B and A,C were used to obtain the Cliff–Lorimer constants, \( k_{AB} \) and \( k_{AC} \). Suppose the x-ray intensity of element C was less reliably correlated to an independent determination of composition of the samples, and the error in \( k_{AB} \) was estimated as 1 %, whereas the error in \( k_{AC} \) was estimated as 10 %.

(a) Estimate the error in a calculated \( k_{BC} \).

(b) Estimate the error in the absolute concentration of B for a material of A and B elements, with nominal compositions 10 % B, 50 % B, and 90 % B.

4.5

The EDS data in Figs. 3.8b–d below were obtained from the Al-Ag precipitate shown in a. The number of counts in the peaks and background beneath them in d are:

- Al K\( \alpha \): 14,986 in peak, 1,969 in background
- Ag K\( \alpha \): 10,633 in peak, 1,401 in background

Given that \( k_{\text{AgAl}} = 2.3 \) for the microscope conditions used, and that the thin-film approximation is valid, what is the composition of the precipitate?

4.6

In an EELS experiment, suppose we seek to measure the near-edge region from an element with a low concentration in the specimen. To optimize the ratio of
edge jump to background, is it better to use a large or small acceptance angle for the EELS spectrometer? Why?

(Hint: Assume the angular dependence of the background is that of a single absorption edge that lies at lower energy than the edge of the dilute element.)

(Further Hint: Look at the Bethe surface.)

4.7

Consider the probability of inelastic scattering, \( p_i \), and elastic scattering, \( p_e \), through a thin layer of material. We set \( p_e + p_i = p \), where \( p \) is the total probability of scattering from the incident beam.

(a) For thin samples of \( n \) layers, show that the x-ray mass absorption factor, \( \mu = (n/x)p \), where \( x \) is the thickness of one layer.

Subsequent layers have the same probabilities, so for \( n \) thin layers we expect:

\[
(p_e + p_i)^n = p^n .
\]  

(b) For thin samples, show that the ratio of double inelastic scattering, \( p_{2i} \), to single inelastic scattering, \( p_i \), is: \( p_{2i}/p_i = (n/2)p_i \).

(Hint: Perform a binomial expansion of (10.54) and consider the physical meaning of the individual terms.)
4.8

When a hole in the core shell of an atom decays by an Auger process, one electron falls into the core hole and a second electron carries energy from the atom. A proper treatment of the Auger effect accounts for the indistinguishability of the two electrons. For electrons of the same spin, this includes antisymmetrizing the two-electron wavefunction of the initial state:

\[ \psi_{\alpha\gamma} = \frac{1}{\sqrt{2}} \left[ \psi_{\alpha}(\vec{r}_1)\psi_{\gamma}(\vec{r}_2) - \psi_{\gamma}(\vec{r}_1)\psi_{\alpha}(\vec{r}_2) \right]. \]  

(3.27)

(a) Write integral expression(s) for the matrix element, \( \langle \beta k | H' | \alpha\gamma \rangle \), with \( H' = e/(|\vec{r}_1 - \vec{r}_2|) \), using (3.27) for \( \psi_{\alpha\gamma} \) and a similar expression for \( \psi_{\gamma\alpha}^* \).

(b) Show that the rate of the Auger transition,

\[ \Gamma = \frac{2\pi}{\hbar} \int \psi_{\gamma\alpha}^* \frac{e}{|\vec{r}_1 - \vec{r}_2|} \psi_{\alpha\gamma} d^3\vec{r}_1 d^3\vec{r}_2, \]  

(3.28)

involves the difference of two matrix elements, one for the transition \( |\alpha\gamma\rangle \to |\beta k\rangle \) and the other for the “exchange transition,” \( |\alpha\gamma\rangle \to |k\beta\rangle \).

(c) The inelastic scattering of a high-energy electron by a core electron transition involves two electrons, the perturbation \( H' = e/(|\vec{r}_1 - \vec{r}_2|) \), and the same calculation as in a and b (compare (2.24) and (3.28)). Is the exchange transition important for the scattering of a high-energy electron? Why?
Chapter 4

Correlation Function for Elastic Scattering – The Patterson Function

4.1 Overview and Definitions

In much of Chapter 1, scattering theory has been developed by calculating the amplitude of the wave scattered from crystals with excitations or disorder. The amplitude of the diffracted wave, $\psi$, is the sum of phase factors of wavelets emitted from individual atoms. For elastic scattering, which we consider presently, the phase information in $\psi(\vec{Q})$ includes details of atom positions, which can be obtained by inverse Fourier transformation, $F^{-1}\psi$. We then calculate the intensity $I(\vec{Q}) = \psi^*\psi$.

This Chapter 4 takes a different approach of calculating directly the diffracted intensity $I(\vec{Q})$, rather than calculating it as $\psi^*\psi$. In this new approach, the real space information is obtained with the Fourier inversion $F^{-1}I$, rather than $F^{-1}\psi$, but this sacrifices some information about atom positions. Nevertheless, the intensity is the actual quantity measured in a diffraction experiment, so this new approach offers a more rigorous understanding of what structural information is available from diffraction experiments. Furthermore, in cases of severely disordered materials, there may be no obvious way to obtain the atom positions needed for a calculation of $\psi(\vec{Q})$. For problems involving severe structural disorder, another advantage of direct manipulations of $I(\vec{Q})$ is that a convenient reference state proves to be a homogeneous distribution of scatterers, or uncorrelated scatterers as in an ideal gas. A powerful tool for calculating diffraction intensities from such materials (and regular crystals too) is the “Patterson function,” defined in Sect. 4.1.2 as an autocorrelation function of the scattering factor distribution.
Whereas the diffracted wave, \( \psi(\vec{Q}) \), is the Fourier transform of the scattering factor distribution, the diffracted intensity, \( I(\vec{Q}) \), is the Fourier transform of the Patterson function of the scattering factor distribution.

The Patterson function is a function in real space, with argument \( \vec{r} \). The Patterson function is a convolution, so the reader should be familiar with convolutions and the convolution theorem (Sect. A.1) before reading the present chapter. The presentation here of real-space correlation functions is good preparation for the discussion that follows on space-time correlation functions. We begin by proving the emphasized statement above. The subsequent section uses the Patterson function to explain diffraction phenomena involving displacements of atoms off of periodic positions owing to temperature.

### 4.1.1 Atom Centers at Points in Space

The most important results in this chapter are obtained by assuming the scatterers are points. At each point, \( \vec{r}_j \), resides the scattering strength of one entire atom, \( f_\vec{r}_j \) (or one unit cell). The actual shape of the atom is included later by convolution, and does not change the main results obtained with point atoms.

It proves convenient to consider a distribution of scatterers, \( f(\vec{r}) \), with a continuous variable, \( \vec{r} \), rather than a sum over discrete points, \( \{\vec{r}_j\} \). We change variables as:

\[
\psi(\vec{Q}) = \sum_{\vec{r}_j} f_\vec{r}_j e^{-i\vec{Q} \cdot \vec{r}_j} = \int_{-\infty}^{\infty} f(\vec{r}) e^{-i\vec{Q} \cdot \vec{r}} d^3\vec{r} . \tag{4.1}
\]

To equate a continuous integral to a discrete sum requires that \( f(\vec{r}) \) is not a smooth function of position. Over most of space \( f(\vec{r}) \) is zero, but at atom centers such as \( \vec{r} = \vec{r}_i \), \( f(\vec{r}_i) \) is a Dirac delta function times a constant, \( f_\vec{r}_i \):

\[
f(\vec{r}_i) = f_\vec{r}_i \delta(\vec{r} - \vec{r}_i) . \tag{4.2}
\]

Recall the important property of the Dirac delta function:

\[
y(x') = \int_{-\infty}^{\infty} \delta(x - x') y(x) \, dx . \tag{4.3}
\]

Equation (4.3) requires that \( \delta(x - x') \) is zero everywhere, except at the point \( x = x' \). At this point the delta function is infinitely high, but of unit area, so the integral of (4.3) picks out only the value of \( y(x) \) at \( x' \). To extend (4.2) to include many atom centers, we take the sum over \( \vec{r}_i \):

\[
f(\vec{r}) = \sum_{\vec{r}_i} f_\vec{r}_i \delta(\vec{r} - \vec{r}_i) , \tag{4.4}
\]

so we satisfy the equality in (4.1) between points in space, \( \{\vec{r}_j\} \), and a continuous function of \( \vec{r} \). We include the shape of the atomic form factor, \( f_a(\vec{r}) \), in Sect. 4.2.2.
4.1. OVERVIEW AND DEFINITIONS

4.1.2 Definition of the Patterson Function

We define the “Patterson function,” \( P(\vec{r}) \):

\[
P(\vec{r}) \equiv \int_{-\infty}^{\infty} f^*(\vec{r}') f(\vec{r} + \vec{r}') \, d^3\vec{r}'. \tag{4.5}
\]

Equation (4.5) is a convolution. Since the function \( f(\vec{r}) \) is not inverted in the usual way for a convolution, we write:

\[
P(\vec{r}) = f^*(\vec{r}) * f(-\vec{r}), \tag{4.6}
\]

This is a specific type of convolution known as an “autocorrelation function,” sometimes denoted with a special symbol:

\[
P(\vec{r}) = f(\vec{r}) \odot f(\vec{r}). \tag{4.7}
\]

The most important feature of the Patterson function is that its Fourier transform is the diffracted intensity in kinematical theory. To show this, we use (4.1) to write \( I(\vec{Q}) = \psi^* \psi \) as:

\[
I(\vec{Q}) = \int_{-\infty}^{\infty} f^*(\vec{r}') e^{i\vec{Q} \cdot \vec{r}'} \, d^3\vec{r}' \int_{-\infty}^{\infty} f(\vec{r}'') e^{-i\vec{Q} \cdot \vec{r}''} \, d^3\vec{r}''. \tag{4.8}
\]

Since \( \vec{r}' \) and \( \vec{r}'' \) are independent variables:

\[
I(\vec{Q}) = \int_{-\infty}^{\infty} \left( \int_{-\infty}^{\infty} f^*(\vec{r}') f(\vec{r}'' - \vec{r}') \, d^3\vec{r}' \right) d^3\vec{r}''. \tag{4.9}
\]

Define \( \vec{r} \equiv \vec{r}'' - \vec{r} \), and change variables \( \vec{r}'' \rightarrow \vec{r} + \vec{r} \). In so doing, the limits of integration for \( \vec{r} \) are shifted by \( -\vec{r} \), but this is not of concern for integrations performed over all of space:

\[
I(\vec{Q}) = \int_{-\infty}^{\infty} \left( \int_{-\infty}^{\infty} f^*(\vec{r}) f(\vec{r} + \vec{r}) \, d^3\vec{r} \right) e^{-i\vec{Q} \cdot \vec{r}} \, d^3\vec{r}'. \tag{4.10}
\]

\[
I(\vec{Q}) = \int_{-\infty}^{\infty} \left( \int_{-\infty}^{\infty} f^*(\vec{r}) f(\vec{r} + \vec{r}) \, d^3\vec{r} \right) e^{-i\vec{Q} \cdot \vec{r}} \, d^3\vec{r}. \tag{4.11}
\]

Using the definition of (4.5), we rewrite (4.11):

\[
I(\vec{Q}) = \int_{-\infty}^{\infty} P(\vec{r}) e^{-i\vec{Q} \cdot \vec{r}} \, d^3\vec{r}. \tag{4.12}
\]
Equation (4.12) shows that the diffracted intensity is the Fourier transform of the Patterson function:

$$I(\vec{Q}) = FP(\vec{r})$$ \hspace{1cm} (4.13)

and by the inverse transformation we must have:

$$P(\vec{r}) = F^{-1}I(\vec{Q})$$ \hspace{1cm} (4.14)

For comparison, the diffracted wave, $\psi(\vec{Q})$ of (4.1), is the Fourier transform of the scattering factor distribution, $f(\vec{r})$. We therefore have another relationship between $I(\vec{Q})$ and $f(\vec{r})$:

$$I(\vec{Q}) = \psi^*(\vec{Q}) \psi(\vec{Q})$$ \hspace{1cm} (4.15)

$$I(\vec{Q}) = \left(Ff(\vec{r})\right)^* Ff(\vec{r}) = |Ff(\vec{r})|^2$$ \hspace{1cm} (4.16)

Comparing (4.13) and (4.16):

$$FP(\vec{r}) = |Ff(\vec{r})|^2$$ \hspace{1cm} (4.17)

Equation (4.17) is consistent with the convolution theorem of Sect. A.1 – a convolution in real space (the Patterson function of (4.5)) corresponds to a multiplication in Fourier space (right-hand side of (4.17)). Note how (4.16) shows the effects of the flip and the complex conjugation of $f(\vec{r})$ in the convolution of (4.5):

$$F[f^*(\vec{r}) * f(-\vec{r})] = \left(Ff(\vec{r})\right)^* Ff(\vec{r}) = |f(\vec{Q})|^2$$ \hspace{1cm} (4.18)

as compared to:

$$F[f(\vec{r}) * f(\vec{r})] = Ff(\vec{r}) Ff(\vec{r}) = (f(\vec{Q}))^2$$ \hspace{1cm} (4.19)

### 4.2 Properties of Patterson Functions

#### 4.2.1 Friedel's Law

It is instructive to illustrate the steps in constructing a Patterson function (4.5). The steps in any convolution are shift, multiply, and integrate, and are shown in Fig. 4.1. Figure 4.1a shows the overlap of a function shifted by the distance $\vec{r}$ against the original position shown as a dashed curve. To obtain the Patterson function in Fig. 4.1b, at each shift the function was multiplied by its shifted counterpart, then integrated.

Note that the peaks of the Patterson function in Fig. 4.1b are broader than the peaks in the scattering factor distribution of Fig. 4.1a. Since the peaks in Fig. 4.1a are Gaussian functions of equal width, the peaks in the Patterson function
are broadened by a factor of $\sqrt{2}$. Second, the periodicity of the Patterson function is one lattice constant, $a$. This is expected, since the overlap of the peaks in the function of Fig. 4.1a is maximized each time the shift equals an integral number of lattice constants. The intensities of these primary maxima are proportional to $A^2 + B^2$. There are secondary maxima that occur at shifts of $\pm 0.3a$ when the large peak overlaps the small peak. The intensities of these secondary maxima are proportional to $AB$. The Patterson function has a peak at each distance corresponding to a separation between the peaks in Fig. 4.1a.

The Patterson function, $P(\vec{r})$ of Fig. 4.1b, has a higher symmetry than the $f(\vec{r})$ of Fig. 4.1a. Identical secondary peaks occur in $P(\vec{r})$ when the large peak is shifted to the right by $+0.3a$ and overlaps the small peak, or when the small peak is shifted to the left by $-0.3a$ and overlaps the large peak. For this reason, even when $f(\vec{r})$ has no center of inversion, $P(\vec{r})$ has inversion symmetry. The Patterson function is unchanged if the original function is inverted.¹ Equation (4.14) shows that the measured x-ray diffraction intensity provides the Patterson function, not the scattering factor distribution. We therefore have “Friedel’s law”:

Diffraction experiments cannot distinguish between an atom arrangement and the atom arrangement when it is inverted.

This is sometimes called the “phase problem” in structure determination, since the phase of the diffracted wave $\psi(\vec{Q})$ is not measured, only its intensity,

¹You can obtain the same $P(\vec{r})$ by taking the mirror image of the $f(\vec{r})$ in Fig. 4.1a (with the small peak to the immediate left of the large peak), and repeating the construction.
4.2.2 Perfect Crystals

In working problems with Patterson functions, it is often convenient to write the scattering factor distribution for an entire crystal, \( f(\vec{r}) \), in the following way:

\[
 f(\vec{r}) = \sum_{\vec{R}_n} f_{\text{at}}(\vec{r}) \delta(\vec{r} - \vec{R}_n). \tag{4.20}
\]

Here \( f_{\text{at}}(\vec{r}) \) is the form factor of one atom. In (4.20) the form factor of the atom is convoluted with a sum of delta functions, each centered at a different atom site, \( \vec{R}_n \). We evaluate (4.20) by first writing explicitly the convolution:

\[
 f(\vec{r}) = \int_{-\infty}^{\infty} f_{\text{at}}(\vec{r}') \delta(\vec{r} - (\vec{r}' - \vec{R}_n)) d^3\vec{r}'. \tag{4.21}
\]

Rearranging the operations on independent variables:

\[
 f(\vec{r}) = \sum_{\vec{R}_n} \int_{-\infty}^{\infty} f_{\text{at}}(\vec{r}') \delta(\vec{r}' - \vec{R}_n) d^3\vec{r}'. \tag{4.22}
\]

The integral of (4.22) serves to pick out the value of \( f_{\text{at}}(\vec{r}') \) at the location of the delta function, cf., (4.3). By shifting the delta function continuously by \( \vec{r}' \), the shape of \( f_{\text{at}}(\vec{r}) \) is generated around the center of each delta function. These centers are each atom site, \( \vec{R}_n \), so after the integration of (4.22):

\[
 f(\vec{r}) = \sum_{\vec{R}_n} f_{\text{at}}(\vec{r} - \vec{R}_n). \tag{4.23}
\]

Please compare (4.20) and (4.23).

The Patterson function of an infinite one-dimensional perfect crystal, \( P_0(x) \), is:

\[
 P_0(x) = f^*(x) * f(-x), \tag{4.24}
\]

which we write using (4.20) for \( N \) atoms:

\[
 P_0(x) = \left( \sum_{n'=-\infty}^{+\infty} \delta(x - n'a) \right) \ast \left( \sum_{n''=-\infty}^{+\infty} \delta(n''a - x) \right). \tag{4.25}
\]

Convolutions are commutative and associative, so we rearrange (4.25):

\[
 P_0(x) = \left( \sum_{n'=-\infty}^{+\infty} \delta(x - n'a) \right) \ast \left( \sum_{n''=-\infty}^{+\infty} \delta(n''a - x) \right). \tag{4.26}
\]
4.2. PROPERTIES OF PATTERSON FUNCTIONS

Recall that a convolution of two functions requires a shift, overlap, multiplication, and integration. Because the δ-functions are infinitesimally narrow, there is zero overlap of the two series of δ-functions unless the shift, x, satisfies the condition \( x = na \), where \( n \) is an integer. Therefore:

\[
\left( \sum_{n'=-\infty}^{\infty} \delta(x - n'a) \right) \ast \left( \sum_{n'=-\infty}^{\infty} \delta(x - n'a) \right) = N' \left( \sum_{n=-\infty}^{\infty} \delta(x - na) \right).
\]

Here \( N' = \infty \), which is as expected for an infinite number of overlaps of an infinite chain of atoms. For a chain of \( N \) atoms, the Patterson function is:

\[
P_0(x) = \sum_{n=-\infty}^{\infty} \delta(x - na) \ast \left( \sum_{n=-\infty}^{\infty} \delta(x - na) \right),
\]

The Fourier transformation of \( P_0(x) \) provides the diffracted intensity, \( I(Q) \). By the convolution theorem of Sect. A.1, the two convolutions and one multiplication of (4.28) become, after Fourier transformation, two multiplications and one convolution. Using (4.18):

\[
I(Q) = N (f_{at}(Q) \ast f_{at}(-Q)) \ast \left( \sum_{n=-\infty}^{\infty} \delta(x - na) \right),
\]

The Fourier transform of the δ-function series is:

\[
F \left[ \sum_{n'=-\infty}^{\infty} \delta(x - n'a) \right] = \int_{-\infty}^{\infty} e^{-iQx} \sum_{n'=-\infty}^{\infty} \delta(x - n'a) \, dx.
\]

The condition \( Qa = 2\pi h \) (where \( h \) is an integer) must be satisfied, or the integration over an infinite range of \( x \) is zero. The \( k \)-space Fourier transform is therefore zero except when \( Q = 2\pi h / a \) precisely, so:

\[
F \left[ \sum_{n'=-\infty}^{\infty} \delta(x - n'a) \right] = \sum_{h=-\infty}^{\infty} \delta(Q - 2\pi h / a) = N \sum_{g} \delta(Q - g).
\]

Here again \( N \) is the number of terms in the sum in (4.31). In a formal problem, \( N \) becomes a mathematical infinity, but it is useful to keep the \( N \) because it shows the proportionality to the size of the crystal. The diffraction intensity of (4.29) is:

\[
I(Q) = N^2 |f_{at}(Q)|^2 \left[ \sum_{h=-\infty}^{\infty} \delta(Q - 2\pi h / a) \right].
\]

Equation (4.32) is a familiar result in a new form. The series of δ-functions gives the centers of the Bragg peaks from the crystal. These peaks are still sharp, but are attenuated at large \( Q \) by the atomic form factor intensity, \( |f_{at}(Q)|^2 \).
CHAPTER 4. CORRELATION FUNCTION FOR ELASTIC SCATTERING – THE PATTERSON FUNCTION

Figure 4.2: Overlap of periodic delta functions, $\sum_n \delta(x-na)$, with a random function of zero mean, $\Delta f(x)$. Since the deviation function $\Delta f(\vec{r})$ has zero mean and is non-periodic, the periodic delta functions overlap $\Delta f(-\vec{r})$ at as many positive values as negative values, demonstrating (4.38).

4.3 Deviations from Periodicity

4.3.1 The Deviation Patterson Function

In many cases of interest, a scattering factor distribution, $f(\vec{r})$, can be expressed as the sum of a perfectly periodic function, $f(\vec{r})$, plus a deviation function, $\Delta f(\vec{r})$, which provides the random or semi-random deviations from perfect periodicity. We know that the perfectly periodic function, $f(\vec{r})$, provides sharp Bragg diffractions, but how does the deviation function, $\Delta f(\vec{r})$, affect the diffracted intensity? To find out, we calculate the Patterson function of $f(\vec{r})$:

$$f(\vec{r}) = \langle f(\vec{r}) \rangle + \Delta f(\vec{r}) \quad (4.33)$$

$$P(\vec{r}) = f^*(\vec{r}) f(-\vec{r}) \quad (4.34)$$

$$P(\vec{r}) = \langle f^*(\vec{r}) \langle f(-\vec{r}) \rangle + \langle f^*(\vec{r}) \Delta f(-\vec{r})
+ \Delta f^*(\vec{r}) \langle f(-\vec{r}) \rangle + \Delta f^*(\vec{r}) \Delta f(-\vec{r}) \quad (4.35)$$

Look at the second term in (4.35). We rewrite it with the aid of (4.17):

$$\langle f^*(\vec{r}) \Delta f(-\vec{r}) = \langle f^*(\vec{r}) \rangle \sum_{\vec{R}_n} \delta(\vec{r} - \vec{R}_n) \Delta f(-\vec{r}) \quad (4.36)$$

Convolutions are associative, so we can group the second and third factors in (4.36), and consider the new convolution:

$$\sum_{\vec{R}_n} \delta(\vec{r} - \vec{R}_n) \Delta f(-\vec{r}) = \sum_{\vec{R}_n} \Delta f(-\vec{R}_n) \quad (4.37)$$

where we used (4.3) in the same way as for (4.22)–(4.23). We assume that the deviation function, $\Delta f(\vec{R}_n)$, has zero mean value.\footnote{This does not restrict generality because any non-zero mean could have been transferred into $\langle f(\vec{r}) \rangle$ in (4.33).} Therefore:

$$\sum_{\vec{R}_n} \delta(\vec{r} - \vec{R}_n) \Delta f(-\vec{r}) = 0 \quad (4.38)$$
The second term for $P(\vec{r})$ in (4.35) is therefore zero (see also Fig. 4.2). Because $\vec{R}_n$ has precise periodicity over an infinite distance, (4.38) also holds true when $\Delta f(\vec{r})$ has short-range structure. By the same argument, the third term in (4.35) is also zero. Equation (4.35) becomes:

$$P(\vec{r}) = \langle f^*(\vec{r}) \rangle \ast \langle f(-\vec{r}) \rangle + \Delta f^*(\vec{r}) \ast \Delta f(-\vec{r}).$$ (4.39)

The Patterson function for an alloy with disorder is reduced to two parts defined as the two terms in (4.39): 1) a Patterson function from the average crystal, $P_{avge}(\vec{r})$, and 2) a Patterson function from the deviation crystal, $P_{devs}(\vec{r})$:

$$P(\vec{r}) = P_{avge}(\vec{r}) + P_{devs}(\vec{r}).$$ (4.40)

The diffracted intensity is the Fourier transform of the Patterson function of the alloy:

$$I(\vec{Q}) = \mathcal{F} \left[ P_{avge}(\vec{r}) + P_{devs}(\vec{r}) \right],$$ (4.41)

and since Fourier transforms are distributive:

$$I(\vec{Q}) = \mathcal{F} \left[ P_{avge}(\vec{r}) \right] + \mathcal{F} \left[ P_{devs}(\vec{r}) \right].$$ (4.42)

Equation (4.42) shows that the diffraction patterns from the average crystal, $\langle f(\vec{r}) \rangle$, and the deviation crystal, $\Delta f(\vec{r})$, are additive. In terms of the diffracted waves from these average and deviation crystals (cf., (4.17)):

$$I(\vec{Q}) = |\mathcal{F} \langle f(\vec{r}) \rangle|^2 + |\mathcal{F} \Delta f(\vec{r})|^2.$$ (4.43)

We are familiar with the first term in (4.43), $|\mathcal{F} \langle f(\vec{r}) \rangle|^2$, which gives the sharp Bragg diffractions from the average crystal.

The second term in (4.43), $|\mathcal{F} \Delta f(\vec{r})|^2$, is new. It is often a broad, diffuse intensity, as we show next. We will also show that with increasing disorder and larger $\Delta f(\vec{r})$, the sharp Bragg diffractions become weaker, and the diffuse intensity becomes stronger. Two important sources of $\Delta f(\vec{r})$ in a crystalline alloy are atomic displacement disorder and chemical disorder. Atomic displacement disorder comprises small deviations of atoms from the sites of a perfect crystal. These displacements may be static, or dynamic as in the case of thermal motion. Chemical disorder exists when there is randomness in the species of atoms that occupy the sites of a crystal. We consider these two types of disorder in sequence.

### 4.3.2 Coherent and Incoherent Scattering

The same argument of Section 4.3 can be used to understand the balance of coherent and incoherent neutron scattering. The idea is that the scattering length $b_i$ from atom $i$ contains a part that is the same for all nuclei, $\langle b \rangle$, and
a deviation part, $\delta b_i$. Since this $\delta b_i$ is a deviation from the average, it is both positive and negative. When averaged over all atoms, the sum of the deviations is zero, i.e.,

$$\langle \delta b_i \rangle = 0 .$$  \hspace{1cm} (4.44)

The scattering length at atom $i$ is

$$b_i = \langle b \rangle + \delta b_i .$$  \hspace{1cm} (4.45)

The scattered neutron wavefunction is constructed as usual

$$\psi(\vec{Q}) = \sum_i b_i e^{i \vec{Q} \cdot \vec{r}_i} ,$$  \hspace{1cm} (4.46)

$$\psi(\vec{Q}) = \sum_i [\langle b \rangle + \delta b_i] e^{i \vec{Q} \cdot \vec{r}_i} .$$  \hspace{1cm} (4.47)

The intensity, $\psi^* \psi$, is

$$I(\vec{Q}) = \sum_i (b_i)^2 e^{-i \vec{Q} \cdot \vec{r}_i} \sum_j (b_j)^2 e^{i \vec{Q} \cdot \vec{r}_j} ,$$  \hspace{1cm} (4.48)

$$I(\vec{Q}) = \sum_i \sum_j \left[ (b_i)^2 + (\delta b_i + \delta b_j) (b_i + \delta b_i \delta b_j) \right] e^{i \vec{Q} \cdot (\vec{r}_j - \vec{r}_i)} .$$  \hspace{1cm} (4.49)

We define the pair distance $\vec{r}_k \equiv \vec{r}_j - \vec{r}_i$. Averaged over all pairs of atoms, the middle term gives zero. The last term $\delta b_i \delta b_j$ is also zero, except for the special case when $i = j$, which denotes the deviation at the same atom (so $\vec{r}_k = 0$)

$$I(\vec{Q}) = N \sum_k (b_k)^2 e^{i \vec{Q} \cdot \vec{r}_k} + \sum_i \delta b_i^2 e^{i \vec{Q} \cdot \vec{r}_i} ,$$  \hspace{1cm} (4.50)

$$I(\vec{Q}) = N \sum_k (b_k)^2 e^{i \vec{Q} \cdot \vec{r}_k} + \sum_i \delta b_i^2 .$$  \hspace{1cm} (4.51)

Equation (4.51) separates the intensities of coherent scattering and incoherent scattering. The first term depends on the interference of waves scattered by the average atom. It gives a series of Bragg peaks. This is wave-like behavior from coherent scattering. The second term is independent of $\vec{Q}$ and does not depend on wave interference. The intensity of the second term increases in proportion to the number of atoms in the material, but the flat shape of the intensity in $\vec{Q}$ is the same for one atom or for $N$ atoms. This is particle-like behavior from incoherent scattering.

Sometimes the second term in (4.51) is rewritten as follows

$$\langle b_i^2 \rangle = \left( \langle b \rangle + \delta b_i \right)^2 ,$$  \hspace{1cm} (4.52)

$$\langle b_i^2 \rangle = \langle b \rangle^2 + 2\langle b \rangle \delta b_i + \delta b_i^2 ,$$  \hspace{1cm} (4.53)

$$\langle b_i^2 \rangle = \langle b \rangle^2 + 2\langle b \rangle \delta b_i + \langle \delta b_i^2 \rangle ,$$  \hspace{1cm} (4.54)

$$\langle b_i^2 \rangle = \langle b \rangle^2 + \langle \delta b_i^2 \rangle ,$$  \hspace{1cm} (4.55)

$$\langle \delta b_i^2 \rangle = \langle b_i^2 \rangle - \langle b \rangle^2 .$$  \hspace{1cm} (4.56)
4.4 Uncorrelated Displacements

Atomic displacement disorder exists when atoms do not sit precisely on the periodic sites of a crystal. Atomic size differences in an alloy cause static displacements from lattice sites, and thermal vibrations cause dynamic displacement disorder. Both cause diffuse scattering. Here we consider a simple type of displacement disorder where each atom has a small, random shift, \( \delta \), off its site of a periodic lattice as shown in Fig. 4.3.

For now we assume there are no correlations between the displacements, \( \delta_i \), of neighboring atoms.\(^3\) The Patterson function, \( f(x) * f(-x) \), for this displacement distribution is shown in Fig. 4.4a. To understand this Patterson function, consider the overlap of the atom center distribution with itself after a shift of \( x = na + \xi \), where \( a \) is the lattice parameter, \( n \) is an integer, and \( \xi \) is a small distance (typically \( \xi < a \)). With no correlation between the displacements of neighboring atoms, the probability of overlap of two atom centers is the same for a shift of the crystal by many lattice constants, \( na + \xi \), as it is for a shift of one lattice constant, \( 1a + \xi \). The important exception occurs around \( x = 0 \), i.e., when \( n = 0 \). All the atom centers overlap perfectly with themselves when \( \xi \) is exactly zero, but even for the smallest shift, \( \xi \neq 0 \), there are zero overlaps of atom centers.

The best way to work with the Patterson function in Fig. 4.4a is to break it into periodic and non-periodic parts (4.40), as shown in the two plots in Fig.

\(^3\)For example, we assume that if one atom is displaced to the left, its neighbor to the right is equally likely to be displaced to the left or to the right.
Figure 4.4: (a) Patterson function for the random displacements of Fig. 4.3 and (4.40). (b) The Patterson function at top is the sum of $P_{\text{avg}}(x)$ and $P_{\text{devs}}(x)$. (c) $P_{\text{devs}}(x)$ is the sum of $P_{\text{devs1}}(x)$ and $P_{\text{devs2}}(x)$. 
4.4. UNCORRELATED DISPLACEMENTS

Figure 4.5: The Fourier transform of the Patterson functions of Fig. 4.4. Fourier transform of $P_{avge}(x)$ (top), Fourier transforms of the two components of $P_{devs}(x)$ (middle). The sum of all three components (bottom) is the diffraction intensity from our linear crystal with Gaussian displacement disorder.

4.4b. The diffracted intensity from our crystal with displacement disorder is obtained from (4.42) as the sum of the Fourier transforms of these two functions, $P_{avge}(x)$ and $P_{devs}(x)$. The Fourier transform of $P_{avge}(x)$ is the well-known series of Bragg peaks. These peaks are suppressed at large values of $Q$ owing to the breadths of the peaks in $P_{avge}(x)$ caused by displacement disorder (see top of Fig. 4.5). This suppression of the Bragg peaks at large $Q$ is similar to the suppression caused by the atomic form factor, which also broadens the scattering centers of the atoms.

The Fourier transform of $P_{devs}(x)$ is new for us. To understand its contribution to the diffraction intensity, we split $P_{devs}(x)$ into two parts, $P_{devs1}(x)$ and $P_{devs2}(x)$ (Fig. 4.4c). The first, $P_{devs1}(x)$, is a Dirac delta function, whose Fourier transform is a constant in $k$-space ($\mathcal{F}[P_{devs1}(x)]$ in Fig. 4.5). The second part, $P_{devs2}(x)$, is a short, broadened function with negative sign. (In Sect. 4.5 we will consider it to be a Gaussian function.) Its Fourier transform, $\mathcal{F}[P_{devs2}(x)]$, is also shown in Fig. 4.5. The areas of these two parts, $P_{devs1}(x)$ and $P_{devs2}(x)$, are equal, since both arise from the same total number of atom-atom overlaps (equal to the number of atoms, $N$). This has an important consequence for the diffracted intensity at $Q = 0$:

$$I(Q=0) = \int_{-\infty}^{\infty} P_{devs}(x) e^{-i0x} \, dx = \int_{-\infty}^{\infty} P_{devs}(x) \, dx,$$

which is simply the area of the Patterson function, $P_{devs}(x)$. Since $P_{devs1}(x)$ and $P_{devs2}(x)$ have equal and opposite areas, at $Q = 0$ there is zero diffuse scattering from atomic displacement disorder.

The $\mathcal{F}[P_{devs2}(x)]$ has a negative sign that decreases in magnitude with $Q$. The diffuse scattering therefore increases with $Q$, as the flat contribution originating from $\mathcal{F}[P_{devs1}(x)]$ increasingly dominates over $\mathcal{F}[P_{devs2}(x)]$. The function
$P_{\text{devs}_2}(x)$, incidentally, has the same shape as the individual peaks in $P_{\text{ave}}(x)$. In this case the $Q$-dependence of the rolloff of the Bragg peaks is the same as the $Q$-dependence of the diffuse scattering. The effects of displacement disorder increase with the characteristic size of the displacements, $\delta_j$. The larger the characteristic $\delta$, the faster the rolloff of the Bragg peaks with $Q$, and the greater the intensity of the diffuse scattering.

### 4.5 Temperature

During thermal vibrations, the distances between atoms undergo rapid changes with time. It is useful, however, to think of each x-ray scattering event as taking an instantaneous snapshot of the atom positions. The diffraction data are averages of many different instantaneous atom configurations. Over a large crystal, however, each instantaneous snapshot looks approximately the same. The same argument of the previous section on atomic displacement disorder is then appropriate for understanding the diffraction effects caused by thermal disorder in atom positions. This section uses a simple model of atom vibrations to calculate two effects of temperature:

- the Debye–Waller factor that causes the Bragg peaks to lose intensity,
- the thermal diffuse scattering, which is where the “lost” intensity reappears.\(^4\)

A detailed analysis of thermal vibrations is not simple, because it should be performed with knowledge of the polarizations and numbers of all phonon modes. A complete analysis considers the contribution of each phonon to the relative separation of each atom-atom pair in the solid. In phonons with long wavelengths, for example, neighboring pairs of atoms tend to move together.\(^5\) In contrast, high frequency phonons affect strongly the mutual displacements of neighboring atoms. In addition, it is important to know how the atom motions within each phonon are oriented with respect to $\vec{Q}$ — atom motions nearly perpendicular to $\vec{Q}$ have weak effects on the scattering. Calculating the Patterson function from the densities of phonons with all polarizations is a problem for computers, and is beyond the scope of this book.

Thermal vibrations broaden the Patterson function of the scattering factor distribution. To develop a simple analytical model, we assume each atom center has a thermal spread around its crystal site that is a Gaussian function of characteristic width, $\sigma$. (A plausibility argument for a Gaussian function is provided in Appendix A.11.) For any $n^{th}$ neighbor pair of atoms, we expect the vibrations of both atoms to affect the Patterson function. Suppose we place a

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\(^4\)The total coherent cross-section remains constant.

\(^5\)Another aspect of the problem is that a crystal has fewer long-wavelength than short-wavelength vibrational modes. However, the lower energy of the long-wavelength modes means that their occupancy is higher at all temperatures, especially low temperatures.
4.5. TEMPERATURE

Figure 4.6: (a) The thermal spread of centers for atom 2. (b) Weights for the centers of the thermal distribution of the atom 1. (c) The distribution of all thermal separations of atom 1 with respect to atom 2.

stationary atom 1 at a point in space. When an atom 2 vibrates with respect to atom 1, there is a probability distribution for their interatomic separation, $x$:

$$p_{\text{atom}2}(x) = \frac{1}{\sqrt{\pi \sigma}} e^{-x^2/\sigma^2}.$$  \hfill (4.60)

This function is shown schematically in Fig. 4.6a. Now let atom 1 vibrate. For every interatomic separation provided by the thermal motion of the atom 2, make a thermal distribution for the position of atom 1. To obtain the distribution of separations between atoms 1 and 2, the displacement distribution of atom 2 serves to weight the displacement distribution of atom 1. The various weights are shown in Fig. 4.6b, and the weighted sum of the net thermal distribution of atom 1 with respect to atom 2 is shown schematically in Fig. 4.6c as $p \ast p(x)$. The procedure we followed was in fact a convolution: the distribution of atom 1 was shifted, $p_{\text{atom}1}(x - x')$, multiplied by $p_{\text{atom}2}(x')$, and summed (integrated) over all values of $x'$:

$$P_{\text{therm}}(x) = \int_{-\infty}^{\infty} p_{\text{atom}2}(x') p_{\text{atom}1}(x - x') \, dx'.$$  \hfill (4.61)

For $n^{\text{th}}$ neighbor pairs of atoms, the Patterson function of the thermal spread, $P_{\text{therm}}(x)$, is the convolution of the Gaussian thermal spread functions of both atoms (cf., (8.24)):

$$P_{\text{therm}}(x) = \left( \frac{1}{\sqrt{\pi \sigma}} e^{-x^2/\sigma^2} \right) \ast \left( \frac{1}{\sqrt{\pi \sigma}} e^{-x^2/\sigma^2} \right) \text{ when } n \neq 0.$$  \hfill (4.62)

$$P_{\text{therm}}(x) = \frac{1}{\sqrt{2\pi \sigma}} e^{-x^2/(2\sigma^2)} \text{ when } n \neq 0.$$  \hfill (4.63)
A detailed analysis treats closer pairs of atoms differently from more distant pairs, but here we ignore this difference except for the case of \( n = 0 \). In the special case of \( n = 0 \), we are considering the autocorrelation function between the positions of the individual atoms with themselves. Each atom sees itself as being at rest, so the Patterson function for the thermal spread is:

\[
P_{\text{therm}}(x) = N\delta(x) \quad \text{when} \quad n = 0 . \tag{4.64}
\]

We now obtain the Patterson function for the entire crystal by convoluting the thermal spread function, \( P_{\text{therm}}(x) \), with the Patterson function of the perfect crystal (4.28). The Patterson function, \( P(x) \), for the crystal with thermal displacement disorder is the following modification of (4.28). Note the special treatment of the \( n = 0 \) term, which provides the \( \delta \)-function:

\[
P(x) = N\left[ f_{\text{at}}^*(x) \ast f_{\text{at}}(-x) \right] + \left[ \delta(x) + \sum_{n \neq 0, \infty} \delta(x - na) \ast \left( \frac{1}{\sqrt{2\pi}\sigma} e^{-x^2/(2\sigma^2)} \right) \right]. \tag{4.65}
\]

We rewrite the sum in (4.65) by adding and subtracting the \( n = 0 \) term (the same trick used in Fig. 4.4 to give an uninterrupted infinite series for \( P_{\text{ave}}(x) \)):

\[
P(x) = N\left[ f_{\text{at}}^*(x) \ast f_{\text{at}}(-x) \right] \left[ \delta(x) - \frac{1}{\sqrt{2\pi}\sigma} e^{-x^2/(2\sigma^2)} \right] \\
+ \left( \sum_{n \neq 0, \infty} \delta(x - na) \ast \left( \frac{1}{\sqrt{2\pi}\sigma} e^{-x^2/(2\sigma^2)} \right) \right] . \tag{4.66}
\]

The diffracted intensity is the Fourier transform of the Patterson function of (4.66). The transformation from (4.66) to (4.67) follows that from (4.28) to (4.32), plus the fact that the Fourier transform of a Gaussian is a Gaussian:

\[
I(Q) = N\left| f_{\text{at}}(Q) \right|^2 \left[ 1 - e^{-\sigma^2(Q)^2/2} \right] \\
+ e^{-\sigma^2(Q)^2/2} \sum_n \delta(Q - 2\pi n a) \right] . \tag{4.67}
\]

The last term in the square brackets is the expected set of sharp Bragg peaks, but attenuated at larger values of \( Q \) by the “Debye–Waller factor,” \( D(Q) \):

\[
D(\sigma, Q) = e^{-\sigma^2(Q)^2/2} . \tag{4.68}
\]

The Debye–Waller factor suppresses the intensity of Bragg peaks at high \( Q \), as does the size of the atom through the factor \( |f_{\text{at}}(Q)|^2 \) of Sect. 3.3.2, so the Debye–Waller factor can be considered a “thermal fattening of the atoms.” The intensity lost from the Bragg peaks reappears\(^6\) as the first term in brackets in

\(^6\)Never forget that the total cross-section for coherent scattering is constant.
4.5. TEMPERATURE

(4.67), $1 - e^{-x^2(Q)^2/2}$, which is the “thermal diffuse scattering.” The thermal diffuse scattering has no distinct peaks, but usually has gradual modulations that increase with $Q$ as shown in Fig. 4.5.

The Debye–Waller factor can provide quantitative information about the mean-squared displacement, $\langle x^2 \rangle$, during thermal motion of the atoms. The larger is $\langle x^2 \rangle$, the smaller the Debye–Waller factor (and the larger the suppression of the Bragg diffractions). We first relate $\langle x^2 \rangle$ to the $\sigma^2$ in the thermal spread function of the individual atoms. This is the second moment of the Gaussian function:

$$\langle x^2 \rangle = \int_{-\infty}^{\infty} x^2 \frac{1}{\sqrt{\pi} \sigma} e^{-x^2/\sigma^2} \, dx = \frac{1}{2} \sigma^2,$$  

(4.69)

so from (4.68) and (5.20):

$$D(\sigma, Q) = e^{-\langle x^2 \rangle(Q)^2} = e^{-\langle x^2 \rangle(4\pi \sin \theta/\lambda)^2}.$$  

(4.70)

At modest temperatures and small $Q$ we can often linearize the exponential to predict a suppression of the Bragg peaks that is quadratic in $Q$:

$$D(\sigma, Q) \approx 1 - \langle x^2 \rangle \left(\frac{4\pi \sin \theta}{\lambda}\right)^2.$$  

(4.71)

Physically, the Debye–Waller factor accounts for the loss of constructive interference in diffraction when the mean-squared atomic displacements become comparable to the x-ray wavelength. The Debye–Waller factor always suppresses the intensity of Bragg peaks.

Equations (4.70) or (4.71) can be used to determine $\langle x^2 \rangle$ from experimental data on diffraction intensities.\(^7\) Conversely, it is often important to predict the Debye–Waller factor for a material at a known temperature. In essence, $\langle x^2 \rangle$ is proportional to the potential energy of a harmonic oscillator, and scales linearly with temperature, $T$. Although $\langle x^2 \rangle$ can be calculated easily for the single oscillator in the Einstein model, it is more handy to express the Debye–Waller factor in terms of a Debye temperature, $\theta_D$, since tabulations of $\theta_D$ are conveniently available. For the Debye model the Debye–Waller factor has been worked out, and at temperatures comparable to the Debye temperature or higher, the Debye–Waller factor is:

$$D(T, Q) \approx \exp \left[ \frac{-12\hbar^2 T}{mk_b \theta_D^2} \left(\frac{\sin \theta}{\lambda}\right)^2 \right],$$  

(4.72)

$$D(T, \theta) \approx 1 - \frac{22,800 T}{m \theta_D^2} \left(\frac{\sin \theta}{\lambda}\right)^2.$$  

(4.73)

\(^7\)Note that $\langle x^2 \rangle$ is along the direction of $Q$. In an isotropic material $\langle x^2 \rangle$ would equal 1/3 of the mean-squared atomic displacement.
Here the units of mass are the atomic weight (e.g., 55.847 for Fe), $T$ and $\theta_D$ are in Kelvin, and $\lambda$ is in Å. For use in (4.70) and (4.71), in the Debye model:

$$\left\langle x^2 \right\rangle = 144.38 \frac{T}{m \theta_D^2}. \quad (4.74)$$

Although the Debye–Waller factor pertains to the thermal spread of distances between pairs of atoms, a Debye–Waller factor is often assigned to the scattering from a single atom. With this approximation, the atomic form factor, $f$, of each atom is replaced with $f \exp(-M)$. The Debye–Waller factor for the intensity is therefore $\exp(-2M)$. Also defined is the parameter $B$, related to $\left\langle x^2 \right\rangle$.

Standard relationships are:

$$2M = \left\langle x^2 \right\rangle \left(\frac{4\pi \sin \theta}{\lambda}\right)^2, \quad (4.75)$$

$$M = B \left(\frac{\sin \theta}{\lambda}\right)^2. \quad (4.76)$$

In the case of an alloy, it is typical to assign different Debye–Waller factors for each type of atom, A or B, written as $e^{-M_A}$ and $e^{-M_B}$:

$$\psi(\vec{Q}) = \sum_r \left[ e^{-M_A} f_A \delta_A(\vec{r}) + e^{-M_B} f_B \delta_B(\vec{r}) \right] e^{i\vec{Q}\cdot\vec{r}}. \quad (4.77)$$

Here the $\delta$-functions are Kroneker delta functions indicating the presence of an A or B atom at $\vec{r}$.

At temperatures below approximately half the Debye temperature, and especially below a quarter of the Debye temperature, (4.72) is no longer reliable for calculating the Debye–Waller factor. Two quantum effects are important at low temperatures. First, owing to Bose–Einstein phonon population statistics, the higher frequency phonons are not excited in simple proportion to the ratio $kT/\varepsilon$, where $\varepsilon$ is the phonon energy. Second, at temperatures below about half the Debye temperature, the “zero-point” vibrations of the solid account for an increasingly large fraction of the atom displacements. Owing to zero-point vibrations, the thermal diffuse scattering can never be eliminated, even by cooling to arbitrarily low temperature.

The derivation of (4.67) was clean because we assumed the same Gaussian thermal spread for all interatomic correlations. For long wavelength phonons, however, adjacent atoms tend to move together in a group. In general, the nearest-neighbor pair correlations are less broadened than the correlations for more distant neighbor pairs. If atoms tend to move in groups, as in acoustic modes, the displacement has long-range modulations, and the thermal diffuse scattering intensity is concentrated near the reciprocal lattice points. The detailed shape of the thermal diffuse scattering depends on the lattice dynamics of the crystal vibrations [9.2]. With a Born–von Kármán model of lattice dynamics, for example, it is possible to calculate the projected components of the atom movements normal to the diffracting planes, and obtain a more accurate $P_{\text{therm}}(x)$ of (4.63). Alternatively, the phonon spectrum of the crystal can
be deduced from measurements of the thermal diffuse scattering, at least in principle. In practice, such measurements require careful correction for other sources of diffuse intensity (such as atomic size and displacement effects).
Chapter 5

Memory Functions and Incoherent Inelastic Scattering

5.1 Overview and Definitions

Chapter 4 showed how the measured diffraction intensity \( I(Q) \) (as opposed to the wave \( \psi(Q) \)) can be inverted to obtain the Patterson function \( P(x) \). It does not give the positions of atoms, but correlations between pairs of atoms. Formally, this is all the information that can be obtained by a diffraction experiment.

In the present chapter, consider the spectral power density \( I(\omega) \) (or with energy \( I(\epsilon) \) because \( \epsilon = \hbar \omega \)). The \( I(\omega) \) can be inverted by Fourier transformation to obtain a function of time, but this does not give the time dynamics of individual atoms (which are individual atom velocities). It gives, rather, the correlations of velocities of one atom between pairs of times. This sacrifices some information about atom dynamics. Nevertheless, the intensity is the actual quantity measured in an incoherent inelastic scattering experiment, so this new approach offers a more rigorous understanding of dynamics information from experiment. Furthermore, the absolute time is usually not important because the system has the same properties at all times.\(^1\) A powerful tool for calculating inelastic intensities is the “Memory function,” \( M(\tau) \), defined in Sect. 5.1.3 as an autocorrelation function of a scattering factor of one atom as it moves. What is missing from this approach is considerations of correlated motions of the atom with its neighbors. The correlations of pairs of atoms in both space and time are described by Van Hove functions, which are presented in the next chapter.

\(^1\) An important and obvious exception is a pump-probe experiment, where the clock starts at the sharp pump event.
of the Memory function of the velocity of the atom. The Memory function is a function of time, with argument $\tau$. Here $\tau$ is a difference in times. For example, $\tau = t_1 - t_2$, but without a particular marker in time we might write it as $\tau = t_1 - t_0$, or $\tau = t_1 - 0$, or even $\tau = t$ (although this last form may let us forget that $\tau$ is used for correlations between two times). The Memory function is a convolution, so the reader should be familiar with the convolution theorem before reading the present chapter. The text follows closely the form of Sect. 4. We prove the emphasized statement above in Sect. 5.1.3. The later sections use the Memory function to explain scattering phenomena involving errors in the velocity of an atom as it moves with a near-periodic waveform.

5.1.1 Atom Centers at Snapshots in Time

The most important results in this chapter are obtained by assuming the scatterers are points. At each instant, $t_j$, resides the scattering strength of one entire atom, $f_{t_j}$ (or one unit cell). The actual shape of the atom is included later by convolution, and does not change the main results.

It proves convenient to consider a distribution of the scattering centers, $f(t)$, with a continuous variable, $t$, rather than a sum over discrete times, $\{t_j\}$. We change variables as:

$$
\psi(\omega) = \sum_{t_j} f_{t_j} e^{-i\omega t_j} = \int_{-\infty}^{\infty} f(t) e^{-i\omega t} dt .
$$

(5.1)

To equate a continuous integral to a discrete sum requires that $f(t)$ is not a smooth function of time. At any time, $f(t)$ is generally zero, but when the atom center crosses a point at $x$, at times such as $t = t_j$, $f(t_j)$ is a Dirac delta function times a constant, $f_{t_j}$:

$$
f(t_j) = f_{t_j} \delta(t - t_j) .
$$

(5.2)

Recall the important property of the Dirac delta function:

$$
y(t') = \int_{-\infty}^{\infty} \delta(t - t') y(t) dt .
$$

(5.3)

Equation (5.3) requires that $\delta(t - t')$ is generally zero, except at the instant $t = t'$. At this instant the delta function is infinitely high, but of unit area, so the integral of (5.3) picks out only the value of $y(t)$ at $t'$. To extend (5.2) to include many atom centers, we take the sum over $t_j$:

$$
f(t) = \sum_{t_j} f_{t_j} \delta(t - t_j) .
$$

(5.4)
so we satisfy relation (5.1) between instants in time, \(\{t_i\}\), and a continuous function of \(t\). We later include the shape of the atomic form factor, \(f_{at}(t)\), in Sect. 5.2.1. The \(f_{at}(t)\) does not mean that the physical scattering amplitude of an atom is changing with time, but that the atom takes some time to cross a point in space as it moves with velocity \(v\).

### 5.1.2 Patterson Function of Moving Atoms

It is perhaps instructive to start with the Patterson function of the previous chapter, and use it to assess the scattered intensity when the atoms are moving. From (4.12)

\[
I(Q) = \int_{-\infty}^{\infty} e^{iQx} P(x) dx . \tag{5.5}
\]

Consider a single atom that is moving while it is scattering a wave. Its \(P(x)\) will be zero when it no longer overlaps itself at the same position. After time \(\tau\),

\[
P_{\tau}(x) = (f_0^* \delta(x)) \ast (f_0 \ast \delta(x + \chi)) . \tag{5.6}
\]

There is a nonzero contribution to \(P_{\tau}(x)\) for a shift \(\chi = -vt\), when the shift \(\chi\) matches the atom translation over the time \(\tau\):

\[
P_{\tau}(x) = |f_0|^2 \ast \delta(v \tau) . \tag{5.7}
\]

\[
P_{\tau}(x) = |f_0|^2 \ast \delta(x - vt) . \tag{5.8}
\]

We can now switch the argument of the Patterson function from position \(x\) to the velocity or to the time – either \(v\) or \(t\) can be used in what follows. Since we will work with time in the next section, we consider here the velocity.

There is an instant in time \(\tau\) when the atom is at the point \(x\).

\[
M_x(v) = |f_0|^2 \ast \delta(v - x/\tau) . \tag{5.9}
\]

This (5.9) is from the correlation function

\[
M_x(v) = |f_0|^2 \ast \delta t = 0(v) \ast \delta t = \tau(v - x/\tau) , \tag{5.10}
\]

with these last two equations following in reverse order as (5.8) to (5.7). Assuming the velocities are the same in the arguments of the \(\delta\)-functions in (5.10), there will be a nonzero contribution as in (5.9).

Now take an average over a long time. The velocity profile of an individual atom is \(v(t)\). If this \(v(t) = v\), a constant, there will be a large contribution to \(M_x(v)\), as is evident from (5.10) and (5.9). In general, \(v(t)\) is not constant, and the average for \(N\) identical atoms is

\[
M_x(\tau) = N|f_0|^2 \int_{t=-\infty}^{\infty} v(t) v(t - \tau) dt . \tag{5.11}
\]
This correlation function is a Patterson function for moving atoms. Consistent with (5.5), the scattered intensity is

\[ I(\omega) = \int_{-\infty}^{\infty} e^{i\omega t} v(t) v(t - \tau) \, dt, \]  

(5.12)

which is the Fourier transform of a velocity-velocity correlation function. Normalization by \( \overline{v^2} \) is typical, where \( \overline{v} \) is the average magnitude of the velocity.

### 5.1.3 Definition of the Memory Function

We define the “Memory function,” \( M(\tau) \):

\[ M(\tau) \equiv \int_{-\infty}^{\infty} f^*(t') f(t + t') \, dt'. \]  

(5.13)

Equation (5.13) is a convolution. Since the function \( f(t) \) is not inverted in the usual way for a convolution, we write:

\[ M(\tau) = f^*(t) \ast f(-t), \]  

(5.14)

This is a specific type of convolution known as an “autocorrelation function,” sometimes denoted with a special symbol:

\[ M(\tau) = f(t) \ast \ast f(t). \]  

(5.15)

The most important feature of the Memory function is that its Fourier transform is the inelastic intensity, for scatterings when the energy transfer creates a single excitation. This is the Wiener–Khinchin Theorem,\(^2\) sometimes stated as “the spectral power density is the Fourier transform of the time correlation function.”

To show this, we use (5.1) to write \( I(\omega) = \psi^* \psi \) as:

\[ I(\omega) = \int_{-\infty}^{\infty} f^*(t') e^{i\omega t'} \, dt' \int_{-\infty}^{\infty} f(t'') e^{-i\omega t''} \, dt''. \]  

(5.16)

Since \( t' \) and \( t'' \) are independent variables:

\[ I(\omega) = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} f^*(t') f(t'') e^{-i\omega (t'' - t')} \, dt dt'. \]  

(5.17)

Define \( t \equiv t'' - t' \), and change variables \( t'' \rightarrow t + t' \). In so doing, the limits of integration for \( t \) are shifted by \( -t' \), but this is not of concern for integrations.

\(^2\)After Norbert Wiener and Aleksandr Khinchin.
5.1. OVERVIEW AND DEFINITIONS

performed over all of time:

\[
I(\omega) = \int_{-\infty}^{\infty} \left( \int_{-\infty}^{\infty} f^*(t') f(t + t') e^{-i\omega t} dt' \right) dt, \quad (5.18)
\]

\[
I(\omega) = \int_{-\infty}^{\infty} \left( \int_{-\infty}^{\infty} \left( f^*(t') f(t+t') \right) e^{-i\omega t} dt' \right). \quad (5.19)
\]

Using the definition of (5.13), we rewrite (5.19):

\[
I(\omega) = \int_{-\infty}^{\infty} M(\tau) e^{-i\omega \tau} d\tau. \quad (5.20)
\]

Equation (5.20) shows that the inelastic intensity is the Fourier transform of the Memory function:

\[
I(\omega) = FM(\tau), \quad (5.21)
\]

and by the inverse transformation we must have:

\[
M(\tau) = F^{-1} I(\omega). \quad (5.22)
\]

These equivalent equations (5.22) and (5.23) are the Wiener–Khinchin theorem.

For comparison, the scattered wave, \(\psi(\omega)\) of (5.1), is the Fourier transform of the scattering factor distribution, \(f(t)\). We therefore have another relationship between \(I(\omega)\) and \(f(t)\):

\[
I(\omega) = \psi^*(\omega) \psi(\omega), \quad (5.23)
\]

\[
I(\omega) = (Ff(t))^* Ff(t) = |Ff(t)|^2. \quad (5.24)
\]

Comparing (5.21) and (5.24):

\[
FM(\tau) = |Ff(t)|^2. \quad (5.25)
\]

Equation (5.25) is consistent with the convolution theorem that states “a convolution in real time (the Memory function of (5.13)) corresponds to a multiplication in frequency space (right-hand side of (5.25)).” Note how (5.24) shows the effects of the flip and the complex conjugation of \(f(t)\) in the convolution of (5.13):

\[
F[ f^*(t) \ast f(-t) ] = (Ff(t))^* Ff(t) = |f(\omega)|^2, \quad (5.26)
\]

as compared to:

\[
F[ f(t) \ast f(t) ] = Ff(t) Ff(t) = (f(\omega))^2. \quad (5.27)
\]
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Figure 5.1: (a) Periodic pulse with a smaller echo occurring slightly later, shown unshifted in time (top), with unshifted pattern shown as dashed curves below it. The other solid curves are time-delayed patterns shown for shifts up to $0.45\tau_0$. (b) The Memory function, obtained by integrating the product of the solid and dashed curves for all shifts, \( r \).

5.2 Properties of Memory Functions

It is instructive to illustrate the steps in constructing a Memory function (5.13). The steps in any convolution are shift, multiply, and integrate, and are shown in Fig. 5.1. Figure 5.1a shows the overlap of a function delayed by the time \( t \) against the original position shown as a dashed curve. To obtain the Memory function in Fig. 5.1b, at each shift the function was multiplied by its shifted counterpart, then integrated.

The peaks of the Memory function in Fig. 5.1b are broader than the peaks in the scattering factor distribution of Fig. 5.1a. Since the peaks in Fig. 5.1a are Gaussian functions of equal width, the peaks in the Memory function are broadened by a factor of $\sqrt{2}$. Second, the periodicity of the Memory function is one vibrational cycle, $\tau_0 = 1/\nu$, where \( \nu \) is a pure vibrational frequency (and $2\pi\nu = \omega$). This is expected, since the overlap of the peaks in the function of Fig. 5.1a is maximized each time delay equal to an integral number of cycles. The intensities of these primary maxima are proportional to $A^2 + B^2$. There are
secondary maxima that occur at shifts of \( \pm 0.3\tau_0 \) when the large peak overlaps the small peak. The intensities of these secondary maxima are proportional to \( AB \). The Memory function has a peak at each distance corresponding to a separation between the peaks in Fig. 5.1a.

The Memory function, \( M(\tau) \) of Fig. 5.1b, has a higher symmetry than the \( f(t) \) of Fig. 5.1a. Identical secondary peaks occur in \( M(\tau) \) when the large peak is shifted to the right by \( +0.3\tau_0 \) and overlaps the small peak, or when the small peak is shifted to the left by \( -0.3\tau_0 \) and overlaps the large peak. For this reason, even when \( f(t) \) has no center of inversion, \( M(\tau) \) has inversion symmetry. The Memory function is unchanged if the original function is inverted. Equation (5.22) shows that the measured inelastic intensity provides the Memory function, not the scattering factor distribution. We therefore have the time analog of Friedel’s law:

\[
\text{Inelastic experiments cannot distinguish the time ordering of an event and its echo, or the echo and the event.}
\]

This is essentially the same as the “phase problem” in structure determination, since the phase of the scattered wave \( \psi(\omega) \) is not measured, only its intensity, \( \psi^*\psi \).

5.2.1 Perfect Periodicity

In working problems with Memory functions, it is often convenient to write the scattering factor distribution for the entire time of the periodicity, \( f(t) \), in the following way:

\[
f(t) = f_{at}(t) \ast \sum_{t_n} \delta(t - t_n).
\]  

(5.28)

Here \( f_{at}(t) \) is the form factor of one atom. It increases the time duration for an atom to pass a point in space, which increases with the size of the atom and inversely with its velocity. In (5.28) the form factor of the atom is convoluted with a sum of delta functions, each at a different time, \( t_n \). (These may be times when a vibrating atom crosses its neutral point, for example.) We evaluate (5.28) by first writing explicitly the convolution:

\[
f(t) = \int_{-\infty}^{\infty} f_{at}(t') \sum_{t_n} \delta(t - (t' - t_n)) dt'.
\]  

(5.29)

Rearranging the operations on independent variables:

\[
f(t) = \sum_{t_n} \int_{-\infty}^{\infty} f_{at}(t') \delta(t - (t' - t_n)) dt'.
\]  

(5.30)

\[3\] You can obtain the same \( M(\tau) \) by taking the mirror image of the \( f(t) \) in Fig. 5.1a (with the small peak to the immediate left of the large peak), and repeating the construction.
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The integral of (5.30) serves to pick out the value of \( f_{at}(t') \) at the location of the delta function, cf., (5.3). By shifting the delta function continuously by \( t' \), the shape of \( f_{at}(t) \) is generated around the center of each delta function. The centers are at the, \( t_n \), so after the integration of (5.30):

\[
f(t) = \sum_{t_n} f_{at}(t - t_n).
\]

Please compare (5.28) and (5.31).

The Memory function of an infinite perfect periodicity, \( M_0(\tau) \), is:

\[
M_0(\tau) = f'(-t) * f(-t),
\]

which we write using (5.28) for \( N \) atoms:

\[
M_0(\tau) = \left( f_{at}(t) * \sum_{n' = -\infty}^{\infty} \delta(t - n'\tau_0) \right) * \left( f_{at}(-t) * \sum_{n'' = +\infty}^{\infty} \delta(n''\tau_0 - t) \right).
\]

Here \( \tau_0 \) is the precise periodicity of the wave of frequency \( \nu \). Convolutions are commutative and associative, so we rearrange (5.33):

\[
M_0(\tau) = \left( f_{at}(t) * f_{at}(-t) \right) * \left( \sum_{n' = -\infty}^{+\infty} \delta(t - n'\tau_0) \right) * \left( \sum_{n'' = +\infty}^{\infty} \delta(n''\tau_0 - t) \right).
\]

Recall that a convolution of two functions requires a shift, overlap, multiplication, and integration. Because the \( \delta \)-functions are infinitesimally narrow, there is zero overlap of the two series of \( \delta \)-functions unless the shift, \( t \), satisfies the condition \( t = n\tau_0 \), where \( n \) is an integer. Therefore:

\[
\left( \sum_{n' = -\infty}^{\infty} \delta(t - n'\tau_0) \right) * \left( \sum_{n'' = -\infty}^{\infty} \delta(t - n''\tau_0) \right) = N' \left( \sum_{n = -\infty}^{\infty} \delta(t - n\tau_0) \right).
\]

Here \( N' = \infty \), which is as expected for an infinite number of overlaps of an infinite chain of atoms. For a chain of \( N \) atoms, the Memory function is:

\[
M_0(\tau) = N \left[ \sum_{n' = -\infty}^{\infty} \delta(t - n'\tau_0) \right] * \left( \sum_{n'' = -\infty}^{\infty} \delta(t - n''\tau_0) \right).
\]

The Fourier transformation of \( M_0(\tau) \) provides the scattered intensity, \( I(\omega) \). By the convolution theorem, the two convolutions and one multiplication of (5.36) become, after Fourier transformation, two multiplications and one convolution. Using (5.26):

\[
I(\omega) = N |f_{at}(\omega)|^2 \ast F \left[ \sum_{n' = -\infty}^{\infty} \delta(t - n'\tau_0) \right],
\]

where \( F \) is the Fourier Transform.
5.3. **DEVIATIONS FROM PERIODICITY**

The Fourier transform of the \( \delta \)-function series is:

\[
F\left[ \sum_{n'=-\infty}^{\infty} \delta(t-n'\tau_0) \right] = \int_{-\infty}^{\infty} e^{-i\omega t} \sum_{n'=-\infty}^{\infty} \delta(t-n'\tau_0) \, dx . \tag{5.38}
\]

The condition \( \omega\tau_0 = 2\pi h \) (where \( h \) is an integer) must be satisfied, or the integration over an infinite range of \( x \) is zero. The \( \omega \)-space Fourier transform is therefore zero except when \( \omega = 2\pi h/\tau_0 \) precisely, so:

\[
F\left[ \sum_{n'=-\infty}^{\infty} \delta(t-n'\tau_0) \right] = N \sum_{l=-\infty}^{\infty} \delta(\omega-2\pi h/\tau_0) . \tag{5.39}
\]

Here again \( N \) is the number of terms in the sum in (5.39). In a formal problem, \( N \) becomes a mathematical infinity, but it is useful to keep the \( N \) because it shows the proportionality to the duration of the periodicity. The inelastic intensity of (5.37) is:

\[
I(\omega) = N^2|f_{al}(\omega)|^2 \left[ \sum_{l=-\infty}^{\infty} \delta(\omega-2\pi h/\tau_0) \right] . \tag{5.40}
\]

Equation (5.40) is a series of \( \delta \)-functions at the primary frequency \( \nu = 1/\tau_0 \), and all of its harmonics. The harmonic structure is extensive in \( \omega \) if the \( f(t) \) is a series of sharp \( \delta \)-functions. In general, these sharp peaks will be attenuated at large \( \omega \) by the atomic form factor, \( |f_{al}(\omega)|^2 \). If the \( f_{al}(t) \) creates a sine wave when convolved with the \( f(t) \), of course only the fundamental \( \nu \) will appear in the spectrum of \( I(\omega) \).

### 5.3 Deviations from Periodicity

In many cases of interest, a scattering factor distribution, \( f(t) \), can be expressed as the sum of a perfectly periodic function, \( \langle f(t) \rangle \), plus a deviation function, \( \Delta f(t) \), which provides the random or semi-random deviations from perfect periodicity. We know that the perfectly periodic function, \( \langle f(t) \rangle \), gives intensity at a fundamental frequency and its harmonics, but how does the deviation function, \( \Delta f(t) \), affect the scattered intensity? To find out, we calculate the Memory function of \( f(t) \):

\[
f(t) = \langle f(t) \rangle + \Delta f(t) , \tag{5.41}
\]

\[
M(\tau) \equiv f^*(t) * f(-t) , \tag{5.42}
\]

\[
M(\tau) = \langle f^*(t) \rangle * \langle f(-t) \rangle + \langle f^*(t) \rangle * \Delta f(-t) + \Delta f^*(t) * \langle f(-t) \rangle + \Delta f^*(t) * \Delta f(-t) . \tag{5.43}
\]

Look at the second term in (5.43). We rewrite it with the aid of (5.25):

\[
\langle f^*(t) \rangle * \Delta f(-t) = \left[ \langle f_{a1}(t) \rangle * \sum_{t_a} \delta(t-t_a) \right] * \Delta f(-t) . \tag{5.44}
\]
Convolutions are associative, so we can group the second and third factors in (5.44), and consider the new convolution:

$$\sum_{t_n} \delta(t - t_n) \ast \Delta f(-t) = \sum_{t_n} \Delta f(-t_n) ,$$

(5.45)

where we used (5.3) in the same way as for (5.30)–(5.31). We assume that the deviation function, $\Delta f(-t_n)$, has zero mean value.\(^4\) Therefore:

$$\sum_{t_n} \delta(t - t_n) \ast \Delta f(-t) = 0 .$$

(5.46)

The second term for $M(\tau)$ in (5.43) is therefore zero (see also Fig. 5.2). Because $t_n$ has precise periodicity over an infinite distance, (5.46) also holds true when $\Delta f(t)$ has short-range structure. By the same argument, the third term in (5.43) is also zero. Equation (5.43) becomes:

$$M(\tau) = \langle f^\ast(t) \rangle \ast \langle f(-t) \rangle + \Delta f^\ast(t) \ast \Delta f(-t) .$$

(5.47)

The Memory function for a periodicity with disorder is reduced to two parts defined as the two terms in (5.47): 1) a Memory function from the average periodicity, $M_{\text{avge}}(\tau)$, and 2) a Memory function from the deviations from periodicity, $M_{\text{devs}}(\tau)$:

$$M(\tau) = M_{\text{avge}}(\tau) + M_{\text{devs}}(\tau) .$$

(5.48)

The scattered intensity is the Fourier transform of the total Memory function:

$$I(\omega) = \mathcal{F}[M_{\text{avge}}(\tau) + M_{\text{devs}}(\tau)] ,$$

(5.49)

and since Fourier transforms are distributive:

$$I(\omega) = \mathcal{F}[M_{\text{avge}}(\tau)] + \mathcal{F}[M_{\text{devs}}(\tau)] .$$

(5.50)

\(^4\)This does not restrict generality because any non-zero mean could have been transferred into $\langle f(t) \rangle$ in (5.41).
Equation (5.50) shows that the inelastic scattering from the average periodicity, \( f(t) \), and the deviation periodicity, \( \Delta f(t) \), are additive. In terms of the scattered waves from these average and deviations (cf., (5.25)):

\[
I(\omega) = |F(f(t))|^2 + |F(\Delta f(t))|^2.
\]  
(5.51)

We are familiar with the first term in (5.51), \( |F(f(t))|^2 \), which gives well-defined peaks in \( I(\omega) \). The second term in (5.51), \( |F(\Delta f(t))|^2 \), gives a broad, diffuse intensity, as we show next. We will also show that with increasing disorder and larger \( \Delta f(t) \), the sharp periodicities in the inelastic spectrum become weaker, and the diffuse intensity becomes stronger. Two important sources of \( \Delta f(t) \) are phase disorder and amplitude disorder. Phase disorder comprises small deviations of atom motions from perfect periodicity. Amplitude disorder exists when there is randomness in the heights of the peaks in \( f(t) \). For simplicity, we assume that these two types of disorder can be treated separately.

### 5.3.1 Amplitude Noise

The argument of Section 5.3 can be used to understand amplitude noise in inelastic scattering. The idea is that the scattering length \( b_i \) from atom \( i \) contains a part that is the same for all nuclei, \( \langle b \rangle \), and a deviation part, \( \delta b_i \). Since this \( \delta b_i \) is a deviation from the average, it is both positive and negative. When averaged over all atoms, the sum of the deviations is zero, i.e.,

\[
\langle \delta b_i \rangle = 0.
\]  
(5.52)

The scattering length at atom \( i \) is

\[
b_i = \langle b \rangle + \delta b_i(t).
\]  
(5.53)

The scattered neutron wavefunction is constructed as usual

\[
\psi(\omega) = \sum_i b_i e^{i\omega t_i},
\]  
(5.54)

\[
\psi(\omega) = \sum_i [\langle b \rangle + \delta b_i(t)] e^{i\omega t_i}.
\]  
(5.55)

The intensity, \( \psi^* \psi \), is

\[
I(\omega) = \sum_i [\langle b \rangle + \delta b_i(t)] e^{-i\omega t_i} \sum_j [\langle b \rangle + \delta b_j(t)] e^{i\omega t_j},
\]  
(5.56)

\[
I(\omega) = \sum_i \sum_j [\langle b \rangle^2 + \langle b \rangle \delta b_i(t) + \delta b_i(t) \langle b \rangle + \delta b_i(t) \delta b_j(t)] e^{i\omega (t_j - t_i)}.  
\]  
(5.57)

We define the time delay \( t_k \equiv t_j - t_i \). Averaged over all pairs of times, the two middle terms give zero. The last term \( \delta b_i(t) \delta b_j(t) \) is also zero, except for the
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special case when \( i = j \), which denotes the same time (so \( t_k = 0 \))

\[
I(\omega) = N \sum_k (\langle b \rangle^2 e^{i\omega t_k} + \sum_i \delta b_i^2 e^{i\omega t_k}) ,
\]

\[
I(\omega) = N \sum_k (\langle b \rangle^2 e^{i\omega t_k} + \sum_i \delta b_i^2) .
\]  

(5.58)

(5.59)

Equation (5.59) separates the intensities of sharp spectral peaks from the diffuse background in \( \omega \). The first term depends on the pattern of waves scattered by the periodic motion. It gives a series of frequency peaks. The second term is independent of \( \omega \). The intensity of the second term increases in proportion to the number of atoms in the material, but the flat shape of the intensity in \( \omega \) is the same for one atom or for \( N \) atoms.

Sometimes the second term in (5.59) is rewritten after the following rearrangement:

\[
\langle b^2 \rangle = \left\langle (\langle b \rangle + \delta b)^2 \right\rangle ,
\]

\[
\langle b^2 \rangle = \langle b^2 \rangle + 2\langle b \rangle \delta b + \delta b_i^2 ,
\]

\[
\langle b^2 \rangle = \langle b^2 \rangle + 2\langle b \rangle \delta b_i + \langle \delta b_i^2 \rangle ,
\]

\[
\langle b^2 \rangle = \langle b^2 \rangle + \langle \delta b_i^2 \rangle ,
\]

\[
\langle \delta b_i^2 \rangle = \langle b^2 \rangle - \langle b \rangle^2 .
\]  

(5.60)

(5.61)

(5.62)

(5.63)

(5.64)

Substituting (5.64) into (5.59), recognizing that the last term in (5.59) can be expressed as the average of \( \delta b_i^2 \) over all \( N \) atoms

\[
I(\omega) = N \sum_k (\langle b \rangle^2 e^{i\omega t_k} + N\langle \delta b^2 \rangle - \langle b \rangle^2) .
\]  

(5.65)

We see that the scattered intensity includes a contribution that depends on the time structuring, or time differences \( \{t_i\} \), and a featureless diffuse scattering. Written as a continuous variable, the average pair distribution is the average Memory function \( M_{\text{avge}}(\tau) \), so

\[
I(\omega) = N\langle b^2 \rangle \int_{-\infty}^{\infty} M_{\text{avge}}(\tau) e^{i\omega \tau} d\tau + N\langle \delta b^2 \rangle - \langle b \rangle^2 .
\]  

(5.66)

5.4 Phase Noise on a Pure Frequency

Suppose there is an underlying periodicity in time of frequency \( \nu \), perhaps imposed externally as a pure frequency. Phase disorder exists when atoms do not follow precisely this periodic waveform. The result is a scattered intensity that is diffuse in \( \omega \). Here we consider a simple type of phase disorder where each atom has a small, random shift, \( \xi \), off its waveform of perfect periodicity, as shown in Fig. 5.3.
5.4. PHASE NOISE ON A PURE FREQUENCY

For now we assume there are no correlations between the timing errors, $\xi_j$.

The Memory function, $f(t) * f(-t)$, for this displacement distribution is shown in Fig. 5.4a. To understand this Memory function, consider the overlap of the atom center distribution with itself after a shift of $t = n\tau_0 + \xi$, where $\tau_0$ is the cycle time, $n$ is an integer, and $\xi$ is a small delay or advance (typically $\xi \ll \tau_0$). With no correlation between the delays or advances, the probability of overlap of the atom centers at the same velocity is the same for a delay of many cycles, $n\tau_0 + \xi$, as it is for a delay of one cycle, $1\tau_0 + \xi$. The important exception occurs around $t = 0$, i.e., when $n = 0$. All the atom centers overlap perfectly with themselves when $n$ is zero, where even for the smallest shift, $\xi \neq 0$, there are zero overlaps of atom centers.

The best way to work with the Memory function in Fig. 5.4a is to break it into periodic and non-periodic parts (5.48), as shown in the two plots in Fig. 5.4b. The scattered intensity from our waveform with phase noise is obtained from (5.50) as the sum of the Fourier transforms of these two functions, $M_{\text{avge}}(\tau)$ and $M_{\text{dev}}(\tau)$. The Fourier transform of $M_{\text{avge}}(\tau)$ is a series of sharp peaks in the frequency spectrum. These peaks are suppressed at large values of $\omega$ owing to the breadths of the peaks in $M_{\text{avge}}(\tau)$ caused by phase errors (see top of Fig. 5.5). This suppression of the frequency peaks at large $\omega$ is similar to the suppression caused by the atomic form factor, which spreads the time when an atom passes a fixed point in $x$.

The Fourier transform of $M_{\text{dev}}(\tau)$ is new for us. To understand its contribution to the inelastic intensity, we split $M_{\text{dev}}(\tau)$ into two parts, $M_{\text{dev1}}(\tau)$ and $M_{\text{dev2}}(\tau)$ (Fig. 5.4c). The first, $M_{\text{dev1}}(\tau)$, is a Dirac delta function, whose Fourier transform is a constant in $\omega$-space ($F[M_{\text{dev1}}(\tau)]$) in Fig. 5.5). The second part, $M_{\text{dev2}}(\tau)$, is a short, broadened function with negative sign. (We could consider it to be a Gaussian function in $\omega$, following Sect. 4.5.) Its Fourier transform, $F[M_{\text{dev2}}(\tau)]$, is also shown in Fig. 5.5. The areas of these two parts, $M_{\text{dev1}}(\tau)$ and $M_{\text{dev2}}(\tau)$, are equal, since both arise from the same number, $N$, of atom-atom overlaps. This has an important consequence for the scattered intensity at $\omega = 0$:

$$I(\omega = 0) = \int_{-\infty}^{\infty} M_{\text{dev}}(\tau) e^{-i0t} \, dx = \int_{-\infty}^{\infty} M_{\text{dev}}(\tau) \, dx = 0.$$  \hspace{1cm} (5.67)

For example, if one cycle is delayed, the subsequent cycle could be either advanced or delayed.
Chapter 5. Memory Functions and Incoherent Inelastic Scattering

Figure 5.4: (a) Memory function for the random displacements of Fig. 5.3 and (5.48). (b) The Memory function at top is the sum of \( M_{\text{avge}}(\tau) \) and \( M_{\text{devs}}(\tau) \). (c) \( M_{\text{devs}}(\tau) \) is the sum of \( M_{\text{devs1}}(\tau) \) and \( M_{\text{devs2}}(\tau) \).

Figure 5.5: The Fourier transform of the Memory functions of Fig. 5.4. Fourier transform of \( M_{\text{avge}}(\tau) \) (top), Fourier transforms of the two components of \( M_{\text{devs}}(\tau) \) (middle). The sum of all three components (bottom) is the inelastic intensity from our periodic displacements plus Gaussian errors in time. This is the effect of “phase noise” on a periodic waveform.
The $F[M_{devs2}](\tau)$ has a negative sign that decreases in magnitude with $\omega$. The diffuse scattering therefore increases with $\omega$, as the flat contribution originating from $F[M_{devs1}](\tau)$ increasingly dominates over $F[M_{devs2}](\tau)$. The function $M_{devs2}(\tau)$, incidentally, has the same shape as the individual peaks in $M_{avege}(\tau)$. In this case the $\omega$-dependence of the rolloff of the frequency peaks is the same as the $\omega$-dependence of the diffuse scattering. The effects of displacement disorder increase with the characteristic size of the displacements, $\xi_j$. The larger the characteristic $\xi$, the faster the rolloff of the frequency peaks with $\omega$, and the greater the diffuse intensity in $\omega$.

5.5 Cumulative Phase Noise

5.5.1 Model of Phase Noise

The previous section described a type of phase noise that was on top of a strict, underlying periodicity. The atom vibrations were locked to this periodicity, except for a bit of jitter. Every cycle, there was a slight delay or advance of the phase of the vibrating atom, but over long times this phase error did not accumulate. Such might be the case for a system of atoms driven by a pure frequency that is applied from an external source.

Here we consider a scatterer without knowledge of an underlying periodicity, but there is still a tendency of the scattering atom to execute a particular vibrational frequency. Over long times, however, we allow the phase to drift, so the phase noise accumulates, and can exceed $\tau_0$ after a while. With random advances and delays in phase at each cycle, the problem becomes a bit like a random walk in time, where the phase drift increases as $\sqrt{n\gamma}$, where $n$ is the number of cycles, and $\gamma$ is an average shift in time from the phase error.

We seek $M(\tau)$, a Memory function for this model. We assume:

1) a statistical independence of errors in advances or delays in the phase of different cycles, and

2) a Gaussian probability distribution for adjacent the distribution of these timing errors.

In this model, the probability distribution for the timing error in the 1st cycle is $P_1(\tau)$, centered about an average separation $\tau_0$:

$$
P_1(\tau) = \frac{1}{\sqrt{\pi \gamma}} e^{-(\tau - \tau_0)^2/\gamma^2}.
$$

(5.68)

Figure 5.6 depicts the time separations between the original and the 2nd cycle. The 1st cycle has a Gaussian distribution, $P_1(\tau)$, with respect to $t = 0$ (Fig. 5.6a). The event at the 2nd cycle is separated by an average time of $\tau_0$ from the 1st, but with a distribution of times. Three arrows are drawn from the time of the 1st cycle in Fig. 5.6b. This gives a spread, $P_1(\tau)$, which is characteristic of the possible times of the 2nd cycle event with respect to the 1st on the left. This extra spread makes the time of the 2nd cycle even more uncertain than the time of the 1st.
Figure 5.6: Construction of the Memory function for a model of cumulative phase errors. (a) Probability distribution for times of 1st cycle. (b) The time of the 2nd cycle has a distribution about each point that is separated from 1st by times $\tau_0$. (c) Composite distribution of 2nd cycle timing errors, accounting for uncertainty in 1st plus 2nd cycle times.

The probability distribution for the 2nd cycle, $P_2(\tau)$, is obtained as $P_1(\tau) \ast P_1(\tau)$, but this distribution is centered about $2\tau_0$. Its width, the width of two convoluted Gaussians, is $\sqrt{2}\gamma$:

$$P_2(\tau) = \frac{1}{\sqrt{2\pi} \gamma} e^{-\left(t - 2\tau_0\right)^2 / (2\gamma^2)}.$$  (5.69)

(Except for the offset in $t$, this step is identical to the analysis of the model of timing error discussed with Fig. 5.4, and it may be useful to examine both figures to look at the differences. The two models now depart, however.) In an analogous way, the probability distribution for the time of the 3rd cycle is obtained as $P_3(\tau) = P_1(\tau) \ast P_1(\tau) \ast P_1(\tau)$, giving a Gaussian centered at $t = 3\tau_0$ with width $\sqrt{3}\gamma$:

$$P_3(\tau) = \frac{1}{\sqrt{3\pi} \gamma} e^{-\left(t - 3\tau_0\right)^2 / (3\gamma^2)}.$$  (5.70)

Examining (5.68), (5.69), (5.70), by induction we obtain $P_n(\tau)$ for the timing distribution for any subsequent cycle, $n$. The total autocorrelation function, $M(\tau)$, is the sum of all distributions of pair separations:

$$M(\tau) = \sum_{n=-\infty}^{\infty} \frac{1}{\sqrt{|n| \pi} \gamma} e^{-\left(t - n\tau_0\right)^2 / (|n|\gamma^2)}.$$  (5.71)

A graph of this Memory function is shown in Fig. 5.7.

The case $n = 0$ always deserves special consideration. For zero time shift ($t = 0$), there is no timing error associated with the overlap of the atom with
5.5. CUMULATIVE PHASE NOISE

Let’s consider the memory function of (5.71) for cumulative phase noise with \( \gamma = a/3 \).

(Conveniently, the \( n = 0 \) term in (5.71) is in fact a \( \delta \)-function.) The time between an atom and its \( n \)th cycle at \( |n|\tau_0 \) is increasingly more uncertain as \( n \) is larger (growing as \( \sqrt{n} \)).

The scattered intensity, \( I(\omega) \), is proportional to the Fourier transform of \( M(\tau) \) of (5.71):

\[
F[M(\tau)] = \int_{-\infty}^{\infty} e^{-i\omega t} \left( \sum_{n=-\infty}^{\infty} \frac{1}{\sqrt{|n| \pi \gamma}} e^{-\left(t-\pi n\tau_0\right)^2/(|n|\gamma^2)} \right) dx .
\]  

(5.72)

When we substitute \( t' = t - n\tau_0 \) to simplify the Gaussians, the phase factor becomes a product of two factors, \( \exp(-i\omega t') \exp(-i\omega n\tau_0) \), where the second factor is independent of \( t' \):

\[
F[M(\tau)] = \sum_{n=-\infty}^{\infty} e^{-i\omega n\tau_0} \int_{-\infty}^{\infty} e^{-i\omega t'} \frac{1}{\sqrt{|n| \pi \gamma}} e^{-t'^2/(|n|\gamma^2)} dx' .
\]  

(5.73)

The Fourier transform of a Gaussian is a Gaussian, and neglecting a constant prefactor:

\[
I(\omega) = \sum_{n=-\infty}^{\infty} \frac{\gamma^2}{4} e^{-\omega^2|n|\gamma^2/4} e^{-i\omega n\tau_0} .
\]  

(5.74)

This sum can be evaluated by first rearranging (5.74) into two geometric series, where the first series includes the terms from \(-\infty\) to \(-1\) and the second from \(0\) to \(+\infty\). Examining the sign of the \( n \) and \(|n|\) in the first sum:

\[
I(\omega) = \sum_{n=1}^{\infty} e^{-(\omega^2\gamma^2/4+i\omega n\tau_0)n} + \sum_{n=0}^{\infty} e^{-(\omega^2\gamma^2/4-i\omega n\tau_0)n} .
\]  

(5.75)

---

This is a handy result. A shift by a constant, \( b \), in time, \( t' = t - b \), amounts to a multiplication by the factor \exp(-i\omega b) \) in \( \omega \)-space.
These two geometric series have the forms:

\[
\sum_{n=0}^{\infty} y^n - 1 = \frac{1}{1 - y} - 1, \quad (5.76)
\]

\[
\sum_{n=0}^{\infty} x^n = \frac{1}{1 - x}. \quad (5.77)
\]

So (5.75) becomes:

\[
I(\omega) = \frac{1}{1 - \exp\left(-\frac{\omega^2 \gamma^2}{4} + i\omega \tau_0\right)} - 1
\]

\[
+ \frac{1}{1 - \exp\left(-\frac{\omega^2 \gamma^2}{4} - i\omega \tau_0\right)}. \quad (5.78)
\]

Using algebra to create common denominators and combine terms, it can be shown that:

\[
I(\omega) = \frac{1 - e^{-\omega^2 \gamma^2/2}}{1 + e^{-\omega^2 \gamma^2/2} - 2e^{-\omega^2 \gamma^2/4} \cos(\omega \tau_0)}.
\quad (5.79)
\]

Figure 5.8 presents graphs of the \( I(\omega) \) of (5.79). There is a \( \delta \)-function at the origin, surrounded by a set of broad peaks spaced at intervals of \( \omega \approx 2\pi/\tau_0 \). These peaks are particularly broad and weak at larger values of \( \omega \). The curves in Fig. 5.8 are labeled with the characteristic width, \( \gamma \), of the Gaussian function in (5.68). Notice the large, nonlinear sensitivity of the peaks to \( \gamma \). As \( \gamma \) decreases to zero, the scattered intensity of (5.79) tends to a sequence of sharp spectral peaks (lower part of Fig. 5.8). The delta function at the origin of our Memory function (the \( n = 0 \) term in (5.71)) causes some flat intensity at large \( \omega \), seen easily in the top part of Fig. 5.8. Finally, we note that the scattered intensity of (5.79) does not include effects of the atomic form factor, which would suppress the scattered intensity at large \( \omega \).
Figure 5.8: Scattered intensities for Memory function of Fig. 5.7 (5.79). The vertical scale of the lower graph is 100× that of the upper graph.
Chapter 6

Coherent Inelastic Scattering

Elastic scattering involves momentum transfers and positions \( \{ \vec{Q}, \vec{r} \} \), which are complementary variables in quantum mechanics: \( \vec{Q} \leftrightarrow \vec{r} \). Inelastic scattering is an extension into energy and time \( \{ \vec{Q}, E, \vec{r}, t \} \), which provides pairs of complementary variables: \( (\vec{Q}, E) \leftrightarrow (\vec{r}, t) \). The amplitude of the scattered wave, \( \psi \), is the sum of phase factors of wavelets emitted from individual atoms, but now we allow for a time variation. The phase information in \( \psi(\vec{Q}, E) \) includes details of atom positions and their motions. Recall the case for elastic scattering, where we obtained by inverse Fourier transformation:

\[
 f(\vec{r}) = F^{-1}_Q \psi(\vec{Q}).
\]

For inelastic scattering the analogous result is:

\[
 f(\vec{r}, t) = F^{-1}_Q F^{-1}_E \psi(\vec{Q}, E).
\]

An inelastic experiment measures an intensity and not a wave amplitude. The experimental information on \( \{ \vec{r}, t \} \) can be obtained directly from the scattered intensity \( I(\vec{Q}, E) \) by Fourier inversion \( F^{-1}_Q F^{-1}_E I(\vec{Q}, E) \). As for the case with the Patterson function for diffraction experiments, when we Fourier transform the intensity instead of the wave itself, there is a similar loss of information about atom positions and the phases of their motions. Nevertheless, the intensity is the actual quantity measured in a scattering experiment and we must make do with it. Inelastic scattering has an important analog to the Patterson function of elastic scattering, the “Van Hove function,” defined in Sect. 6.1.1 as an autocorrelation function of the scattering factor distribution in space and time.

Whereas the inelastically scattered wave, \( \psi(\vec{Q}, E) \), is the double Fourier transform of the moving scattering factor distribution, the scattered intensity, \( I(\vec{Q}, E) \), is the double Fourier transform of the Van Hove correlation function of the scattering factor distribution.

After proving this emphasized statement, subsequent sections use the Van Hove function to explain scattering phenomena involving various dynamical excitations. Then starting anew with Fermi’s Golden Rule, inelastic scattering is
calculated in a more precise and general way. The Van Hove space-time corre-
lation function is developed by taking careful account of the non-commutivity
of the position and momentum operators. A proper treatment of magnetic scat-
tering is then presented. These latter sections are parallel to similar sections in
the books by Squires and Lovesey, which are recommended to all practitioners
of inelastic neutron scattering, especially persons inclined towards theory.

6.1 Correlation Function for Inelastic Scattering –
The Van Hove Function

6.1.1 Concept of Van Hove Function

Preliminary: Atom Centers at Points in Space and Time

As was the case for elastic scattering, most of the essential results for inelastic
scattering can be obtained by assuming the scatterers are points. The point
scatterer emits a wavelet from position \( \vec{r}_j \) at time \( t_k \). This \( \vec{r}_j \) and \( t_k \) provide
the phase of the wavelet from the point emitter relative to wavelets from other
point emitters. This phase is \( \vec{Q} \cdot \vec{r}_j - \omega t_k \). The amplitude of the scattering is given
by the scattering strength of the point emitter, \( f(\vec{r}_j, t_k) \), which is an amplitude
at a point in space and an instant in time. For a nucleus we may consider
\( f \) as being at a point, although for magnetic spin distributions the shape of electron
orbitals may be included later by convolution. For the distribution in time,
we will usually consider a Fourier series with different frequencies, or energies
\( E = \hbar \omega \).

It proves convenient to consider a distribution of scatterers, \( f(\vec{r}, t) \), with
continuous variables, \( \vec{r} \) and \( t \), rather than a sum over discrete points, \( \{ \vec{r}_j \} \), and
snapshots in time \( t_k \). We change variables as:

\[
\psi(\vec{Q}, E) = \sum_{\vec{r}_j} \sum_{t_k} f_{\vec{r}_j, t_k} e^{-i(\vec{Q} \cdot \vec{r}_j - \omega t_k)} = \int_{-\infty}^{+\infty} f(\vec{r}, t) e^{-i(\vec{Q} \cdot \vec{r} - \omega t)} d^3 \vec{r} dt . \tag{6.1}
\]

To equate a continuous integral to a discrete sum requires that \( f(\vec{r}, t) \) is not a
smooth function of position or time. Over most of space and time, \( f(\vec{r}, t) \) is zero,
but when the scattering amplitude exists at \( \vec{r} = \vec{r}_j \) and \( t = t_k \), \( f(\vec{r}_j, t_k) \) is a Dirac
delta function times a constant, \( f_{\vec{r}_j, t_k} \):

\[
f(\vec{r}_j, t_k) = f_{\vec{r}_j, t_k} \delta(\vec{r} - \vec{r}_j) \delta(t - t_k) . \tag{6.2}
\]

To extend (6.2) to include many atom centers, we take the sum over \( \vec{r}_j \) and \( t_k \):

\[
f(\vec{r}, t) = \sum_{\vec{r}_j} \sum_{t_k} f_{\vec{r}_j, t_k} \delta(\vec{r} - \vec{r}_j) \delta(t - t_k) , \tag{6.3}
\]

so we satisfy the equality in (6.1) between points in space and time, \( \{\vec{r}_j, t_k\} \), and
continuous functions of \( \vec{r}, t \).
6.1. Correlation Function for Inelastic Scattering – The Van Hove Function

Definition of the Van Hove Function

We define the “Van Hove function,” \( G(\vec{r}, t) \):

\[
G(\vec{r}, t) = \int_{-\infty}^{+\infty} f^*(\vec{r}, t') f(\vec{r} + \vec{r}', t + t') \, d^3\vec{r}' \, dt'. \tag{6.4}
\]

Equation (6.4) is a double autocorrelation function – a space-time correlation function with limits of integration over all space and all time.

A most important feature of the Van Hove function is that its Fourier transform is the scattered intensity in kinematical theory. To show this, we use (6.1) to write \( I(\vec{Q}, E) = \psi^*(\vec{Q}, E) \times \psi(\vec{Q}, E) \) as:

\[
I(\vec{Q}, E) = \int_{-\infty}^{+\infty} f^*(\vec{r}, t') \, d^3\vec{r}' \times \int_{-\infty}^{+\infty} f(\vec{r}', t'') \, d^3\vec{r}'' \tag{6.5}
\]

Since \( \vec{r} \) and \( \vec{r}' \) are independent variables, as are \( t' \) and \( t'' \):

\[
I(\vec{Q}, E) = \int_{-\infty}^{+\infty} \left( \int_{-\infty}^{+\infty} f^*(\vec{r}, t') f(\vec{r}', t'') \right) \times e^{-i(\vec{Q} \cdot \vec{r} - \omega t')} \, d^3\vec{r} \, d^3\vec{r}' \, dt' \tag{6.6}
\]

Define \( \vec{r}' = \vec{r}'' - \vec{r} \) and \( t = t'' - t' \), and change variables \( \vec{r}'' \to \vec{r} + \vec{r} \) and \( t'' \to t + t' \). In so doing, the limits of integration for \( \vec{r} \) are shifted by \( -\vec{r} \) and \( -t' \), but this is not of concern for integrations performed over all of space and all of time:

\[
I(\vec{Q}, E) = \int_{-\infty}^{+\infty} \left( \int_{-\infty}^{+\infty} f^*(\vec{r}, t') f(\vec{r} + \vec{r}', t + t') \, e^{-i(\vec{Q} \cdot \vec{r} - \omega t)} \, d^3\vec{r} \, dt' \right) \, d^3\vec{r}'. \tag{6.7}
\]

Using the definition of (6.4), we rewrite (6.8):

\[
I(\vec{Q}, E) = \int_{-\infty}^{+\infty} G(\vec{r}, t) \, e^{-i(\vec{Q} \cdot \vec{r} - \omega t)} \, d^3\vec{r} \, dt. \tag{6.9}
\]
Equation (6.9) shows that the scattered intensity is the Fourier transform of the Van Hove function:

\[ I(\mathbf{Q}, E) = \mathcal{F}_x \mathcal{F}_t G(\mathbf{r}, t) , \]

(6.10)

and by the inverse transformation we must have:

\[ G(\mathbf{r}, t) = \mathcal{F}_x \mathcal{F}_t I(\mathbf{Q}, E) . \]

(6.11)

For comparison, the scattered wave, \( \psi(\mathbf{Q}, E) \) of (6.1), is the Fourier transform of the scattering factor distribution, \( f(\mathbf{r}, t) \). We therefore have another relationship between \( I(\mathbf{Q}, E) \) and \( f(\mathbf{r}, t) \):

\[ I(\mathbf{Q}, E) = \psi^*(\mathbf{Q}, E) \psi(\mathbf{Q}, E) , \]

(6.12)

\[ I(\mathbf{Q}, E) = (\mathcal{F}_x \mathcal{F}_t f(\mathbf{r}, t))^* \mathcal{F}_x \mathcal{F}_t f(\mathbf{r}, t) = |\mathcal{F}_x \mathcal{F}_t f(\mathbf{r}, t)|^2 . \]

(6.13)

Comparing (6.10) and (6.13):

\[ \mathcal{F}_x \mathcal{F}_t G(\mathbf{r}, t) = |\mathcal{F}_x \mathcal{F}_t f(\mathbf{r}, t)|^2 . \]

(6.14)

Equation (6.14) is consistent with the convolution theorem of Sect. A.1 – a (double) convolution in real space (the Van Hove function of (6.4)) corresponds to a multiplication in Fourier space (right-hand side of (6.14)).

### 6.1.2 Examples of Van Hove Functions

In this section we examine the scattering from a simple form of the Van Hove function from 6.9 with one spatial dimension:

\[ I(Q, E) = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} G(x, t) e^{-iQx - \omega_0 t} \, dx \, dt . \]

(6.15)

First assume coherent scattering so there are predictable phase relationships between different scattering centers. Section 6.2 develops further the cases where incoherent averaging of the space and time correlations produce a “self-correlation function” and the “Patterson function.” We later treat the incoherent case by following the “self-correlation function” of the individual scatterers.

#### Traveling Wave

Suppose an elementary excitation in a solid provides a periodic modulation of the scattering factor in space and time:

\[ f(x, t) = f_0 e^{i(Qx - \omega_0 t)} . \]

(6.16)
We use (6.4) to obtain its Van Hove function:

\[
G(x, t) = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} f_0^* e^{-i(qx' - \omega_0 t')} f_0 e^{i[q(x + x') - \omega_0(t + t')]} \, dx' dt',
\]

(6.17)

\[
G(x, t) = \int_{-\infty}^{\infty} e^{i(qx - \omega_0 t)} \, dx = e^{i(qx - \omega_0 t)}.
\]

(6.18)

\[
G(x, t) = |f_0|^2 e^{i(qx - \omega_0 t)}.
\]

(6.19)

If we ignore the prefactor $|f_0|^2$, we find $G(x, t) = f(x, t)$. In this case where $G(x, t)$ has the form of a wave (6.19), (6.15) becomes:

\[
I(Q, E) = |f_0|^2 \int_{-\infty}^{\infty} e^{i(qx - \omega_0 t)} \, dx dt,
\]

(6.20)

\[
I(Q, E) = |f_0|^2 \int_{-\infty}^{\infty} e^{i(Qx - \omega_0 t)} \, dx dt,
\]

(6.21)

\[
I(Q, E) = |f_0|^2 \delta(q - Q - g) \delta(\omega - \omega_0).
\]

(6.22)

Equation (6.22) shows that the energy transfer, $\hbar \omega$, must match that of the energy of the excitation in the material, $\hbar \omega_0$. Furthermore, the momentum transfer must match that of the wave, modulo a reciprocal lattice vector, $g$ (which provides a factor of $e^{i gx} = 1$ in the integrand).

An elementary excitation in a solid with unique \{q, \omega_0\} provides intensity at a single point in energy, and at a distance of Q away from each reciprocal lattice point. This of course assumes coherent scattering – incoherent scattering places a restriction on $\omega$ only.

**Dispersive Excitations**

Figure 6.1 presents simple sine waves of different wavelengths, but the same velocity. Such is the case for long-wavelength sound waves in solids. This is the “hydrodynamic limit,” where the wavelengths are so large that the material behaves as a continuum (i.e., atomic features are not important). Three such waves are presented in Fig. 6.1a, but these are not special ones, and we expect that all intermediate wavelengths are possible. Each wave can be understood individually with the same analysis of the traveling wave as in Sect. 6.1.2.

The simple wave of Eq. 6.16 had a scattered intensity that was a delta function at one frequency and one wavevector, Eq. 6.22. If we allow a continuous range of wavevectors, we will have a continuous range of energies of the waves. There will be a continuous set of delta functions on a plot of the scattered intensity in Fig. 6.1b. These are related as $\omega = v k$, where $v$ is the wave
velocity. Note also that $v = d\omega/dk$. Each traveling wave contributes a point to Fig. 6.1b, but for numerous wavelengths these points will make a straight line in the first quadrant of positive $\omega$ and $k$.

There are also excitations where the scattering factor distribution moves to the left in Fig. 6.1a, so we expect a mirror symmetry in $-k$. Also, we expect both creations and annihilations of these excitations, so in the scattered intensity we expect mirror symmetry in $-\omega$, too. The resulting intensity appears as straight lines are in all four quadrants of Fig. 6.1b.

Local Excitations

It is instructive to develop the Van Hove function and the scattered intensity with a pictorial approach, as in Fig. 6.2. The scattering factor distribution is shown in Fig. 6.2a as it undergoes a full cycle of oscillation. The construction of the Van Hove function $G(x,t)$ parallels that of the Patterson function in Fig. 4.1. The detailed steps of shifting and integrating in Fig. 4.1 are not shown here. Suffice to say that situation along the $x$-dimension is quite analogous to that of Fig. 4.1, but a different instantaneous structure exists at each different time. For the case where $t = 0$, for example, it is necessary to average the shifting and integrating of Fig. 4.1 for all nine times shown in Fig. 6.2a. Notice that although

---

1 The slopes of these straight lines $\omega(k)$ are expected to decrease as $k$ gets large, comparable to a reciprocal lattice vector, and atomic-scale effects become important. Nevertheless, it is useful to compare sound velocities obtained from macroscopic measurements to the slopes of phonon dispersions from inelastic scattering. These two velocities generally agree.

2 The convention is that times $t_k$ with larger subscripts are later times. The pattern is repeated after 8 time steps.
some of the scattering factor distributions have zero intensity for separations that are at odd multiples of \( a \), the result for \( G(x, t) \) always has intensity at these locations (although weaker). The time dimension of \( G(x, t) \) is obtained in a similar way. Consider the first time interval for time differences of \( t_1 \). For this case it is necessary to match all pairs of the scattering factor distributions that differ by one in their vertical stacking in Fig. 6.2a. These eight cases are then overlapped and shifted along the \( x \)-axis as before, and their results are averaged. The other time intervals require taking pairs separated further in time, but always it was eight cases whose shift and overlap are evaluated for constructing Fig. 6.2b.

To see how some of the trends in \( G(x, t) \) in Fig. 6.2b come from the \( f(x, t) \) in Fig. 6.2a, first identify the shifts in \( x \) and \( t \) that give optimal overlaps.\(^3\) There is always optimal overlap at the origin, i.e., \( (x, t) = (0, 0) \). Other strong overlaps occur for shifts of \( x \) by \( 2na \) (\( n \) is an integer), and \( t = 0 \). There is also a strong overlap for shifts of \( t \) by \( mt_8 \) (\( m \) is an integer) and \( x = 0 \). The \( G(x, t) \) in in Fig. 6.2b has maxima at these special shifts, and for any combination \( (x = 2na, t = mt_8) \). There is a second set of maxima, however, where the shifts in \( x \) and \( t \) are correlated as \( (x = 2(n + 1)a, t = (m + 1/2)t_8) \). Peaks in \( G(x, t) \) in in Fig. 6.2b can be found for these combinations, too.

The final step is to obtain the \( I(Q, \omega) \) of Fig. 6.2c. In this case the situation is fairly simple. There is only one time frequency in Fig. 6.2a, equal to \( 2\pi/t_8 \). If we allow creation and annihilation of such an excitation, we expect intensity at frequencies \( \omega = \pm 2\pi/t_8 \). The spatial periodicity of the problem includes a superlattice periodicity, so peaks appear at intervals of \( \pm n\pi/a \), where \( n \) is either even or odd.

Propagating Excitations

The next example in Fig. 6.3a is essentially the scattering factor distribution shown in Fig. 4.1, but now moves continuously to the right with increasing time. Its Van Hove function for zero time therefore has the same inversion symmetry along \( x \) as shown in Fig. 4.1, which was used to demonstrate Friedel’s law. In assessing the overlap of the scattering factor distributions, for all cases of zero time shift (i.e., the overlap of the scattering factor with itself before it moves any further), the largest intensity occurs at lattice translation vectors. For most other time shifts \( t \), the evaluation of \( G(x, t) \) sums the overlaps of pairs of \( f(x, t_k) \) in Fig. 6.3a that are separated in time. The best overlaps of these time-shifted \( f(x, t_k) \) generally do not occur at lattice translations. Figure 6.3b shows that the shape of \( G(x, t) \) is constant, but moves to the right with increasing time shift.

Peaks in the scattered intensity \( I(Q, \omega) \) are shown in Fig. 6.3c. Along the time dimension of Figs. 6.3a,b, the pattern repeats itself with a periodicity of eight time snapshots in Figs. 6.3a. The inelastic part of the scattering therefore occurs at \( \pm 2\pi/t_8 \), assuming that excitations of the type shown in Fig. 6.3a can

\(^3\)We assume that the space and time periodicities repeat outside the range shown in Fig. 6.2.
Figure 6.2: (a) Variations of a scattering function $f(x,t)$ in space and time. Nine snapshots are shown vertically for nine $t_i$.  (b) The Van Hove function, $G(x,t)$, obtained by overlapping all pairs of scattering factor distributions in (a), and summing the resulting product of overlaps for all pairs separated by the same number of time intervals.  (c) The scattered intensity, $I(Q,\omega)$. It is assumed possible to both create and annihilate an excitation as shown in (a), hence points at $\pm \omega$. 
6.1. CORRELATION FUNCTION FOR INELASTIC SCATTERING – THE VAN HOVE FUNCTION

Figure 6.3: (a) Variations of a scattering function in space and time. Nine snapshots in time are shown vertically. (b) The Van Hove function, obtained by overlapping all pairs of scattering factor distributions in (a), and summing the resulting product of overlaps for all pairs separated by the same number of time intervals. It was assumed that the scattering factor distribution moves to the right continuously between the time snapshots shown in (a). (c) The scattered intensity. It is assumed possible to both create and annihilate an excitation as shown in (a), hence points at ±ω₀.
be both created and annihilated. There is another point worth noting about the time dependence in Fig. 6.3c. Along the time dimension we encounter sets of three δ-functions for each fundamental periodicity of the wave. The basic periodicity of the δ-functions, and their grouping into threes, requires that we have higher-order Fourier components in the frequency domain—formally, we expect a whole series of excitations of energies: \( \pm n \hbar \omega_0 = \pm n \hbar 2\pi/t_8 \). The incident energy of the wave may be insufficient to excite some of these with higher \( n \), however. This is a bit different from the situation of elastic scattering in \( k \)-space, where it is common to measure a series of Bragg peaks of multiple orders, for example.

**Disordered Excitations**

Another example is presented in Fig. 6.4. This example is a space-time analog of displacement disorder presented in Sect. 4.4. Here we assume that the scattering factor distribution is initially periodic, but is set in motion. Each atom is first displaced to the right, but each atom oscillates with a slightly different frequency. We assume that the frequencies have a narrow spread \( \Delta \omega \) about a central frequency \( \omega_0 \). Likewise, we assume that the amplitudes of the displacements are not large, so each arrow does not travel far from its lattice site. Eventually the oscillation damps away, and the scatterers are back to their initial periodic configuration. It is easiest to first analyze the situation for long times. Here the scatterers are in periodic positions, and have stopped moving. As the time interval becomes long with respect to the damping time, most of the spatial correlations will involve correlations in a precisely periodic structure. For long times, \( G(x, t) \) will exhibit a set of peaks at equal intervals of \( x = a \).

At short time separations, the individual arrows in Fig. 6.4b have not displaced much. At the shortest times after the neutron impact, the scatterers have nearly the same displacements since they move nearly in phase. It is incorrect, however, to assume that the peaks in \( G(x, t) \) are displaced to the right. Recall that it is an average over many different time-separated snapshots of \( f(x, t) \) in Fig. 6.4a, many of which are moving to the left. For small \( t \), \( G(x, t) \) is therefore similar to that of a perfect crystal in its spatial periodicities, with sharp peaks at each lattice translation.

At intermediate times, however, this regularity is lost, even between neighboring arrows in this example. Figure 6.4c shows the intensity from this \( G(x, t) \). For \( \omega = 0 \) the situation is as in Fig. 4.5, but for \( \omega = 2\pi/t_8 \) the intensity is broadened along the \( \omega \) axis, owing the spread in frequencies. It is also expected that the disorder along the time domain will produce extra incoherent scattering between the sharp peaks in \( G(x, t) \).

The initial coherence of the arrows is lost at intermediate times (assuming that the dephasing time \( 2\pi/\Delta \omega \) is less than the damping time). If we assume that there are no short-range spatial correlations, we have precisely the situation considered in Sect. 4.4. Figures 4.4 and 4.5 show the situation for long times, or equivalently for \( \omega = 0 \) in the \( I(Q, \omega) \) of Fig. 6.4.
Figure 6.4: (a) Variations of a scattering function in space and time. Nine snapshots in time are shown vertically. (b) The Van Hove function, obtained by overlapping all pairs of scattering factor distributions in (a), and summing the resulting product of overlaps for all pairs separated by the same number of time intervals. (c) The scattered intensity. It is assumed possible to both create and annihilate an excitation as shown in (a), hence points at $\pm \omega_0$. 
6.2 Autocorrelation Functions

The Van Hove function is worthy of deep respect, because it includes all the information available from all four types of scattering:
(coherent, inelastic), (coherent, elastic),
(incoherent, inelastic), (incoherent, elastic).
Subsets of this total information are often important, perhaps because the scattering is primarily incoherent, or is primarily elastic, for example. In particular, two important, although less general, correlation functions are:

- The Patterson function, \( P(\vec{r}) \), contains the spatial information obtained from diffraction measurements with elastic coherent scattering.
- The “self-correlation” function, \( G_s(t) \), contains the time information obtained from measurements of incoherent inelastic scattering. It involves an averaging over \( Q \).

6.2.1 Autocorrelation Functions from Van Hove Functions

Concept of Patterson and Self-Correlation Functions

Figure 6.5 shows the essential ideas. A “snapshot in time” is obtained by horizontal sampling across the figure. In a real experiment the samplings will be at many different times, but here we see that all such snapshots have the same periodic structure. As we know from Sect. 4.1.2, the diffraction intensity for this time snapshot is obtained from the Patterson function, whose periodicity in space gives the Bragg diffractions. In this particular case of a sine wave, however, we have Bragg peaks only at \( Q = 0 \) and \( Q = \pm g \). If we had a series of \( \delta \)-functions with the same periodicity, however, we would have a series of Bragg peaks to arbitrarily large orders of \( Q = ng \). To obtain the Patterson function, the horizontal lines in Fig. 6.5a, each at a particular \( t_k \), are convoluted with themselves, and the results are averaged for all \( t_k \). The \( f(x, t_k) \) for different \( t_k \) in Fig. 6.5a do not interact with each other until after the convolution is performed. Specifically:

\[
P(x) = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} f(x', t') f(x + x', t') \, dx' \, dt',
\]

where we use the same value of \( t' \) in both arguments of the scattering factor. The \( P(x) \) includes an averaging over time, and does not consider correlations in time. A \( P(\vec{r}) \) is shown in Fig. 6.5 as a horizontal line at a specific time. This information is probed by elastic scattering experiments, which do not allow for measuring the time variations and hence energy transfers.

Likewise we can take a “snapshot in space” by sampling the moving wave, at a specific \( \vec{r}_j \) for example (although the results for all other values of \( \vec{r}_j \) are the same in this example). Notice how the wave in Fig. 6.5 shows a time periodicity
6.2. AUTOCORRELATION FUNCTIONS

Figure 6.5: A simple sine wave, moving to the right with increasing time. Three methods of sampling the intensity of the wave in space and time are shown as straight lines, as explained in the text. The relevant correlation functions for each sampling are shown for each line.

along a vertical line, labeled as \( G_s(t) \). The time periodicity of our moving wave is, of course, \( \omega \), in the wave function: \( e^{i Q x - \omega t} \). The restriction to a single value of \( Q \) eliminates the possibility of obtaining any spatial information on the excitation, but the \( \omega \)-dependence is retained. We obtain the time correlations for a specific position:

\[
G_s(t) = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} f^*(x', t') f(x', t' + t) \, dx' \, dt',
\]

(6.24)

where we use the same value of \( x' \) in both arguments of the scattering factor. The \( G_s(t) \) includes an averaging over position, but does not consider correlations in position. This information is probed by incoherent inelastic scattering experiments, which do not allow for measuring the \( Q \)-dependences and hence spatial information.

More information comes by identifying correlations in both time and space, however, as is indicated by the diagonal line in Fig. 6.5. This requires the full Van Hove function, \( G(\vec{r}, t) \). This \( G(\vec{r}, t) \) cannot be obtained by simply summing the correlation functions of different space or time snapshots, i.e., summing \( G_s(t) \) or \( P(\vec{r}) \). Nevertheless, it is instructive to evaluate the Patterson function and self-correlation function for each of the three examples shown in Figs. 6.2, 6.3, 6.4 (see Problem 3).

Local Excitations

Along the horizontal axis of \( \omega = 0 \) in Fig. 6.2c, we obtain a diffraction pattern with strong fundamentals at \( Q = \pm n\pi/a \), where \( n \) is an even integer, and somewhat weaker superlattice diffractions for odd \( n \). The same result is true for two other values of \( \omega = \pm 2\pi/t_8 \). This periodic structure in \( Q \) is evident from the Van Hove function \( G(x, t) \) in Fig. 6.2b. The spatial correlation function of (6.23), \( P(x) = \int f^*(x', t_k) f(x' + x, t_k) \, dx' \, dt_k \) has spatial periodicities that are similar at any time \( t_k \), although different weights for the superlattice diffractions. The average over all \( t_k \) is of course what makes the final result for \( I(Q, \omega = 0) \) in Fig. 6.2c.
Now look at the correlation functions along lines parallel to the time axis for fixed \( x' \) in \( G(x', t) \) of Fig. 6.2b. The time periodicity of \( G(x', t) \) in Fig. 6.2b differs in phase for even and odd \( n \) in \( x' = \pm na \). All time periodicities are the same, however, and include only the frequencies \(-2\pi/t_8, 0, -2\pi/t_8\). The frequency spectrum of \( G(x', t) \), measured along \( x' = 0 \), is generally the same as the inelastic scattering along any other \( x' = na \). If we average the intensities along these individual \( x' \), we lose the information on phases of scattering between the scatterers at different \( x' \). The mixed constructive and destructive interference between these different scatterers results in intensities between individual atoms only. The result is a “self-correlation” function of (6.24), or a case of incoherent inelastic scattering, \( G_s(t) = \int f^*(x_j, t') f(x_j + x, t' + t) dx_j dt' \).

In this particular case, the incoherent average of these (obtained by summing the intensities at all \( Q \)) looks generally the same as the spectrum along a slice along any particular \( Q \).

The Patterson and self-correlation functions of Fig. 6.3 can be understood in much the same way. The largest horizontal periodicity in Fig. 6.3b is \( a \), so this Patterson function gives diffraction peaks at multiples of \( 2\pi/a \). The self-correlation function likewise has the simple periodicities of \(-2\pi/t_8, 0, -2\pi/t_8\), neglecting the detailed structure in cuts along the time axis that would give higher-order Fourier components.

**Acoustic Excitations**

More interesting are the dispersions of different excitations that have the same wave speed (presented in Fig. 6.1b). All these excitations would be measured in an incoherent inelastic scattering experiment, since the averaging over all \( x' \) of \( G(x', t) \) would preserve all time periodicities. An incoherent inelastic scattering experiment would collapse all the inelastic intensity onto a single \( \omega \)-axis, and lose the relationship between \( \omega \) and \( Q \) that is so evident from the coherent inelastic scattering shown in Fig. 6.1b. For dispersive excitations, the self-correlation function has considerably less information compared to the full Van Hove function. On the other hand, if one is interested in obtaining an accurate density of states (i.e., a total energy spectrum), irrespective of the values of \( Q \), the self-correlation function may be advantageous. It does not require accurate measurements of intensities over all values of \( Q \) and \( \omega \).

**Disordered Excitations**

Finally we consider the Patterson function and self-correlation function of Fig. 6.4. The scatterers oscillate about their lattice sites, and the Patterson function is therefore as expected of a perfect crystal. For the time average, there is displacement disorder in the arrangement, however, so we expect a diffuse

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Incidentally, if the arrows were to oscillate through zero in Fig. 6.2a, we would have no value of \( \omega = 0 \), and no elastic scattering. This corresponds to half the scatterings occurring out-of-phase with the other half.
6.2. AUTOCORRELATION FUNCTIONS

background, increasing with $Q$ as shown in Fig. 4.5. The self-correlation function for incoherent inelastic scattering is the same for all $x$, since the scatterers are assumed independent in their motions, without positional correlations that may cause neighbors to move together in phase. In this case we expect the same time structure for $G_s(t)$ and $G(x,t)$, and hence inelastic incoherent scattering will have the same energy spectrum as the coherent inelastic scattering for all $Q$. Both will show a broadening in frequency of the excitation around $\omega_0$. The broadening arises from two effects. First is the frequency spread of the oscillators, postulated to be $\Delta \omega$. The second effect is the damping of the oscillations. The damping time of $\tau$ provides a broadening in energy of $\hbar/\tau$, known as “lifetime broadening.” In the present problem we have assumed that the broadening of the excitation at $\omega_0$ is dominated by the frequency spread $\Delta \omega$, since the dephasing time was assumed shorter than the damping time.

6.2.2 Inelastic Scattering from Diffusion

Atom Correlations from Diffusion Equation

Here we pick a simple case for uncorrelated atom movements in diffusion. We begin with the Gaussian concentration profile, $c(x,t)$, which is a standard solution of the 1D diffusion equation

$$c(x,t) = \frac{c'}{\sqrt{4\pi Dt}} e^{-\frac{x^2}{4Dt}}.$$  \hspace{1cm} (6.25)

The Van Hove function of Eq. 6.4 is

$$G_{\text{diff}}(x,t) = \int_{-\infty}^{+\infty} c(x',0) c(x' + x,t) \, dx'.$$ \hspace{1cm} (6.26)

An important point is that there are no space or time correlations between different atoms in diffusion – two different atoms do not know about each other’s jumps, and all atoms follow the same concentration profile.\textsuperscript{5} Ensemble averaging is therefore unnecessary.

$$G_{\text{diff}}(x,t) = \int_{-\infty}^{+\infty} c(x',0) c(x' + x,t) \, dx'.$$ \hspace{1cm} (6.27)

As $t \to 0$, the function $c(x', t \to 0)$ of Eq. 6.25 becomes a Dirac delta function of unit area

$$G_{\text{diff}}(x,t) = \int_{-\infty}^{+\infty} \delta(x') c(x',t) \, dx'.$$ \hspace{1cm} (6.28)

\textsuperscript{5}This is unlike phonon scattering, for example, where the motion of one atom follows the motion of another as a vibrational wave moves from one atom to the other.
so the spatial integral simply returns the function \(c(x, t)\) of Eq. 6.25

\[
G_{\text{diff}}(x, t) = \frac{c'}{\sqrt{4\pi D t}} e^{-\frac{x^2}{4Dt}}. \tag{6.29}
\]

We calculate the \(S(Q, \omega)\) by using Eq. 6.29 in the Fourier transform relationship of Eq. 6.9

\[
S(Q, \omega) = \frac{c'}{2\pi \hbar} \int_{-\infty}^{+\infty} \int_{-\infty}^{+\infty} \frac{e^{-\frac{t^2}{4Dt}}}{\sqrt{4\pi D t}} e^{-(Qx-\omega t)} \, dx \, dt, \tag{6.30}
\]

\[
S(Q, \omega) = \frac{c'}{2\pi \hbar} \int_{0}^{+\infty} \int_{-\infty}^{+\infty} e^{-Q^2 Dt} e^{i\omega t} \, dt \, dx, \tag{6.31}
\]

\[
S(Q, \omega) = \frac{c'}{2\pi \hbar} \frac{DQ^2}{(DQ^2)^2 + \omega^2}, \tag{6.32}
\]

where we used a standard result that the Fourier transform of a Gaussian is a Gaussian to obtain Eq. 6.31, and that the Fourier transform of a decaying exponential is a Lorentzian function to obtain Eq. 6.32. The Lorentzian function is a symmetrical, peaked function with a maximum at \(\omega = 0\), or \(\Delta E = 0\). Its half-width at half-maximum is at \(\omega = DQ^2\), or

\[
\Delta E = \hbar DQ^2. \tag{6.33}
\]

**Physical interpretation – quasielastic scattering**

We can interpret Eq. 6.32 in terms of how inelastic scattering is sensitive to the atom jumps in diffusion. The energy width of the inelastic intensity of Eq. 6.33 corresponds to a time \(\Delta \tau\) by the uncertainty relationship

\[
\Delta \tau = \frac{\hbar}{\Delta E} = \frac{1}{DQ^2}. \tag{6.34}
\]

An important result attributed to Einstein relates the diffusion coefficient \(D\) to the atom jump frequency \(\Gamma\) and the jump length \(a\). A factor of \(1/6\) is appropriate for 3D diffusion

\[
D = \frac{\Gamma a^2}{6}, \tag{6.35}
\]

where \(\Gamma\) is the atom jump frequency and \(a\) is its jump length. The time uncertainty becomes

\[
\Delta \tau = \frac{1}{\left(\frac{1}{6}\right)^{\frac{3}{2}}}, \tag{6.36}
\]

\[
\Delta \tau = \frac{0.30}{\Gamma}, \tag{6.37}
\]
for a representative $Q$ of $2\pi/a$. In other words, without diffusion ($D = 0$) the inelastic spectrum would be a sharp peak at $E = 0$, corresponding to elastic scattering only. When the atoms jump frequently, there is time-broadening of the energy $\Delta E = \hbar \Gamma$ at a characteristic $Q$. This is typical of wave scattering that is interrupted by a jump of the scattering center, which shortens the coherence time for the emitted wave, broadening it in energy. This process of scattering by moving atoms is often called “quasielastic scattering” because it can be considered as elastic scattering (no overall energy shift), but with an energy broadening as the moving atoms interrupt the coherence of the elastic line.

### 6.2.3 Relationships Between Intensities, Correlation Functions, Waves, and Scattering Lengths

Much of scattering science involves relationships between the wavefunction $\psi(Q,\omega)$ and physical scattering lengths $f(r,t)$ in the sample. Many important functional relationships are obtained by:

- Fourier transformation $F$
- autocorrelation $\ast$
- multiplication $||^2$
- averaging $\langle \rangle$

A map of the important physical functions and how they are transformed into one another is presented in Fig. 6.6. It is a map of the universe of scattering functions.

When using Fig. 6.6, reverse transformations are possible in all cases, but information is lost by averaging over an argument of a function. For example, $I(\omega)$ can be transformed to $S(Q,\omega)$ in the incoherent approximation, but dispersive information about $\omega(Q)$ cannot be obtained. (Of course, if the scattering is incoherent, there is no dispersive information to begin with.) Averaging over $\omega$ is not shown in Fig. 6.6 because a more common manipulation in scattering science is to identify the elastic scattering within a set of inelastic data at $E = 0$. Finally, the functions $f(r,\omega)$ and $\psi(Q,t)$ are not shown. They may prove useful as intermediate steps in some calculations, but they mix the phase information about the wave and the scatterer.

Some definitions are:

- $S(Q,\omega)$ is the “scattering law.”
- $Y(Q, \tau)$ is the momentum-time correlation function. It is called the “intermediate scattering function.”
- $\Gamma(R, \omega)$ is the dynamical pair distribution function.
- $G(R, \tau)$ is the Van Hove space-time correlation function.
• $P(R)$ is the Patterson function.
• $M(\tau)$ is the “memory function,” a time correlation function for dynamics at a site (sometimes called $G_s(\tau)$, a self-correlation function).
• $I(Q)$ is a diffraction pattern.
• $I(\omega)$ is an inelastic spectrum.
• $f(r, t)$ is a scattering length density.
• $\psi(Q, \omega)$ is a wavefunction.

Two correlation functions in Fig. 6.6 have special uses. The function $\Gamma(R, \omega)$, the dynamical pair distribution function, is used for projections of excitations onto specific sites. More precisely, it gives the vibrational spectrum between a particular pair of atoms. The function $Y(Q, \tau)$ is useful for studies of transient phenomena that may occur after an impulsive perturbation.

### 6.3 Essence of Coherent Inelastic Neutron Scattering

#### 6.3.1 Spherical Waves from Point Scatterers in Motion

An intuitive shortcut from (1.58) to (1.65) is to regard $k^2 + \vec{\kappa}^2$ as a scattering operator that generates a scattered wavelet proportional to $U(\vec{r}')\Psi(\vec{r}')$. The scattered wavelet must have the properties of (1.50) for its amplitude and phase versus distance. The scattered wavelet amplitude emitted from a small volume, $d^3\vec{r}'$, centered about $\vec{r}'$ is:

$$d\Psi_{sc}(\vec{r}', \vec{r}) = U(\vec{r}')\Psi(\vec{r}') \frac{e^{i\vec{\kappa} \cdot (\vec{r}' - \vec{r}) - \omega_0 t}}{|\vec{r}' - \vec{r}|} d^3\vec{r}'. \quad (6.38)$$

This is an expression for a spherical wave at $\vec{r}$ originating from the small volume $d^3\vec{r}'$ surrounding $\vec{r}'$. It is isotropic – note that the exponential factor $k_0 |\vec{r}' - \vec{r}|$ is a product of scalars.

We obtain the total scattered wave by integrating around all volume of the scatterer. The incident plane wave, $\propto e^{i\vec{\kappa} \cdot \vec{r}}$ (1.68), helps sets the phase of the scattering at each volume interval. The phase of the outgoing wave also depends on the orientation of the outgoing $\vec{k}_i$ with respect to the position of the scattering point, $\vec{r}$. The relative phase from each scattering point depends on the change in wavevector, $Q \equiv \vec{k}_i - \vec{k}_f$, as $e^{iQ \cdot \vec{r}}$.

$$\Psi_{sc}(Q, \vec{r}) = -\frac{e^{i(k_0|\vec{r}' - \omega_0 t|)}}{|\vec{r}|} \frac{m}{2\pi\hbar^2} \int \mathcal{V}(\vec{r}') e^{iQ \cdot \vec{r}' \cdot d^3\vec{r}'}. \quad (6.39)$$
Figure 6.6: Interrelationships between Correlation Functions, Scattering Length Densities, Intensities, Wavefunctions
In arriving at (6.38) we have repeated, in an intuitive way, the steps to (1.74) and (1.75). We note that the momentum transfer from the neutron to the sample is \( \hbar \vec{Q} \).

The trick now is to replace the potential, \( V(\vec{r}) \), with a suitable potential for neutron scattering. For nuclear scattering, relevant to phonon measurements, we use the “Fermi pseudopotential,” which places all the potential at a point nucleus:

\[
V_{\text{nuc}}(\vec{r}) = 4\pi \frac{\hbar^2}{2m} \delta(\vec{r}) ,
\]

(6.40)

where \( b \) is a simple constant (perhaps a complex number). For thermal neutrons, the \( \delta \)-function is an appropriate description of the shape of a nucleus.

Fermi pseudopotentials are placed at crystal translation vectors, \( \{ \vec{R}_j \} \), accounting for all \( N \) nuclei in the crystal:

\[
V_{\alpha}(\vec{r}) = 4\pi \frac{\hbar^2}{2m_\alpha} \sum_{j=1}^{N} b_j \delta(\vec{r} - \vec{R}_j) .
\]

(6.41)

This sum of \( \delta \)-functions transforms the integration of Eq. 6.39 into a sum over points at \( \vec{R}_j \). For laboratory distances (large compared to the size of the scatterer), \( \vec{r} \gg \vec{R}_j \), the outgoing spherical wave can be treated as a plane wave, allowing rearrangement

\[
\Psi_{\text{sc}}(\vec{r}, t') = \frac{e^{i(k_f \cdot \vec{r} - \omega_f t')}}{|\vec{r}|} \sum_j b_j e^{i(\vec{k}_i - \vec{k}_f) \cdot \vec{R}_j} e^{i(\omega_f - \omega_i)t} .
\]

(6.42)

The factor \( e^{i(k_f \cdot \vec{r} - \omega_f t')} \) gives the precise phase at the time and place of neutron detection. It has modulus unity, and we ignore it because we measure the intensity of scattering. We use Eq. 8.44 for \( \vec{R}_j \),

\[
\vec{R}_j = \vec{r}_l + \vec{r}_\kappa + \vec{u}_{\kappa}(t) ,
\]

(6.43)

where the three terms are the lattice vector, basis vector of the unit cell (containing \( R \) atoms), and the displacement vector. We allow the scattered wave to emanate from the nucleus at later times

\[
\Psi_{\text{sc}}(\vec{r}) = \frac{1}{r} \sum_{j, \kappa} b_{\kappa} e^{i(\vec{k}_i - \vec{k}_f) \cdot \vec{u}_{\kappa}(0)} e^{-i\vec{k}_f \cdot \vec{u}_{\kappa}(t)} e^{i(\omega_f - \omega_i)t} ,
\]

(6.44)

where we note that because all unit cells are identical, \( b \) depends on the basis vector only.

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6Typically of a nuclear scale of \( 10^{-12} \) cm or so, sometimes with an imaginary component to account for absorption. For magnetic scattering, however, the \( \delta \)-function should be convoluted with a real-space form factor for the magnetic electrons. This could be done at the end of the calculation by multiplying the \( k \)-space result with a form factor.
6.3. ESSENCE OF COHERENT INELASTIC NEUTRON SCATTERING

It is easiest to understand the case when the scattering is a fast process, such as for x-ray scattering, where an x-ray traverses the crystal more rapidly than typical vibrational periods of atoms. Incoherent neutron scattering is also fast, where the thermal neutron is scattered from individual atomic nuclei without concern for phase relationships between nuclei. This motivates a first approximation; ignore the time dependence of the potential by setting $\vec{u}_l(\kappa, t) = \vec{u}_l(\kappa, 0)$, and take the time average

$$\Psi_{sc}(\vec{Q}, \vec{r}) = \frac{1}{\tau} \sum_{l, \kappa} b_{\kappa} e^{i\vec{Q} \cdot (\vec{r}_l + \vec{r}_\kappa)} e^{i\vec{Q} \cdot \vec{u}_l(0)} \frac{1}{\tau} \int_0^\tau e^{i(\omega_\kappa - \omega_l) t} dt,$$

(6.45)

$$\Psi_{sc}(\vec{Q}, \vec{r}) = \frac{1}{\tau} \sum_{l, \kappa} b_{\kappa} e^{i\vec{Q} \cdot (\vec{r}_l + \vec{r}_\kappa)} e^{i\vec{Q} \cdot \vec{u}_l(0)} \delta(\omega_\kappa - \omega_l).$$

(6.46)

The $\delta$-function enforces no change of frequency or energy, so Eq. 6.46 pertains to elastic scattering, where the modes of atom vibrations are unchanged. We need to perform an ensemble average over the different values of $\vec{u}_l(0)$, which evaluates as

$$\langle e^{i\vec{Q} \cdot \vec{u}(0)} \rangle_N = e^{-\frac{1}{2} \langle \vec{Q} \cdot \vec{u}(0) \rangle^2} \equiv e^{-W}. \quad (6.47)$$

Equation 6.46 for elastic scattering becomes

$$\Psi_{elsc}(\vec{Q}, \vec{r}) = e^{-W} \frac{1}{\tau} \sum_{l, \kappa} b_{\kappa} e^{i\vec{Q} \cdot \vec{r}_\kappa} \sum_{l} e^{i\vec{Q} \cdot \vec{r}_l} \equiv e^{-W} F(\vec{Q}) S(\vec{Q}).$$

(6.48)

Here we separated the sums over basis and lattice vectors into a structure factor, $F(\vec{Q})$, and a shape factor, $S(\vec{Q})$. Equation 6.48 can be used to calculate the neutron diffraction pattern from a crystal, giving the average static structure. The factor $e^{-W}$ is the “Debye–Waller factor,” which becomes appreciably less than 1 when the atom displacement $\vec{u}_l$ is comparable to $1/\vec{Q} \sim \lambda/2\pi$, the “wavelength” associated with the scattering angle and energy. This is consistent with the origin of the Debye–Waller factor from the destructive interference of scattered wavelets that are emitted from different spatial positions of the displaced nuclei. Note that the ensemble average of Eq. 6.47 does not reveal the energy of an atom vibration, only how displaced atom positions alter the phase.

Next consider a process of scattering that is slow compared to the atomic vibrational period. Such is the case for coherent scattering of neutrons by

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7An approximate demonstration of this is by expanding $exp(i\vec{Q} \cdot \vec{u}) = 1 + i\vec{Q} \cdot \vec{u} + ...$ and taking either an ensemble average or a time average. The average of the second term, $i\vec{Q} \cdot \vec{u}$ is zero owing to positive and negative excursions of $\vec{u}$, leaving the quadratic term, consistent with Eq. 6.47. The average of the subsequent cubic term also vanishes, conveniently, but unfortunately the quartic term has the wrong sign. Nevertheless, it can be proved that Eq. 6.47 is exact if the probability distribution of $\vec{u}$ is Gaussian about the origin, and this is often reasonable for lattice dynamics.
phonons, and for nuclear resonant scattering of photons by phonons. For clarity, we pick a single phonon with wavevector $\vec{k}$. The form of $\vec{u}_{\kappa \vec{k}}(t)$ is obtained later in Eq. 8.56, but is presented here (mostly for interest, with normalization for a quantum vibration)

$$
\vec{u}_{\kappa \vec{k}}(t) = \sqrt{\frac{2\hbar}{N m_\kappa \alpha_\kappa}} e^{i (\vec{k} \cdot \vec{r}_l - \omega \vec{k} t)}.
$$  (6.49)

Equation 6.49 shows that $\vec{u}_{\kappa \vec{k}}(t) = \vec{u}_{\kappa \vec{k}}(0)e^{-i \omega \vec{k} t}$. We need to take a time average of the displacement factors in Eq. 6.44

$$
\langle e^{i (\vec{k}_i \cdot \vec{r}_l(0))} e^{-i (\vec{k}_f \cdot \vec{r}_l(0))} \rangle_t = \langle e^{i (\vec{k}_i \cdot \vec{u}_{\kappa \vec{k}}(0))} e^{-i (\vec{k}_f \cdot \vec{u}_{\kappa \vec{k}}(0))} e^{-i \omega \vec{k} t} \rangle_t
$$  (6.50)

Coherent phonon scattering involves correlations of displacements $\vec{u}_{\kappa \vec{k}}(t)$ at different atom positions at different times. This Van Hove space-time correlation function (1) is a rich approach to studying dispersive excitations in solids. The proper quantum mechanical treatment of time and temperature averages is taken up in Section 7, but Eq. 6.51 below is the proper result for incoherent scattering. Furthermore, it is possible to average the coherent scattering over all $\vec{Q}$ to recover this incoherent result (2; 3; 4), and the incoherent approximation serves us well in the following analysis of multiphonon scattering. The loss of phase information by averaging the coherent scattering over all $\vec{Q}$ brings us back to consideration of a fast scattering process.

### 6.3.2 One-Phonon and Multiphonon Scattering

Equation 6.48 is the elastic scattering only. It does not conserve the nuclear scattering cross section. Nuclear interactions do not necessarily depend on the displacements of atoms, so we need a compensating factor $e^{+\langle (\vec{Q} \cdot \vec{u}_{\kappa \vec{k}})^2 \rangle}$ to conserve the cross-section and account for the rest of the scattering intensity. Expanding this new factor, we calculate the intensity as $\Psi_{sc}^* \Psi_{sc}$, and for clarity we normalize the intensity

$$
\langle \Psi_{sc}^* \Psi_{sc} \rangle_N = \frac{1}{R} \sum_k e^{-\langle (\vec{Q} \cdot \vec{u}_{\kappa \vec{k}})^2 \rangle}
\times \left( 1 + \langle (\vec{Q} \cdot \vec{u}_{\kappa \vec{k}})^2 \rangle + \frac{1}{2} \langle (\vec{Q} \cdot \vec{u}_{\kappa \vec{k}})^2 \rangle^2 + ... \right)
$$  (6.51)

The first and second terms in parentheses of Eq. 6.51 are the fractions of elastic (0-phonon) and 1-phonon scattering. The third term with the $\frac{1}{2} \langle (\vec{Q} \cdot \vec{u}_{\kappa \vec{k}})^2 \rangle^2$ accounts for 2-phonon scattering processes. Only the 1-phonon scattering proves useful for obtaining a phonon DOS, but it is important to have an understanding of the higher-order terms (called “multiphonon scattering”) for designing an experiment, or performing quantitative corrections of experimental data.
6.3. ESSENCE OF COHERENT INELASTIC NEUTRON SCATTERING

The incoherent approximation proves more robust than it may seem to deserve. Energy and momentum conservation suppress the possibilities for cross-terms between scattering by 0-phonon and 1-phonon processes. The frequencies of the incident and scattered waves are mismatched. Integrated over many periods of the neutron wavefunction, these mismatched waves will not add constructively or destructively, so there is no coherent interaction between these two scattering processes.\(^8\)

A more detailed treatment of 1-phonon scattering includes the thermal factors for phonon populations, giving the double-differential scattering cross section\(^9\)

\[
\frac{d^2\sigma}{d\Omega dE} = \frac{k_i}{k_f} \frac{(2\pi)^3}{2V_0} \sum_{\vec{g},\vec{k}} \frac{n(\vec{k}) + \frac{1}{2} \pm \frac{1}{2}}{\omega_{\vec{k}}^2} \times \left| \sum_{\kappa} \frac{b_\kappa}{\sqrt{m_\kappa}} e^{-i(\vec{Q} \cdot \vec{u}_\kappa)} [\vec{Q} \cdot \vec{e}_\kappa(\vec{k})] e^{i\vec{Q} \cdot \vec{r}_\kappa} \right|^2 \times \delta(Q + \vec{k} \pm \vec{g}) \delta(\omega - \omega_{\vec{k}}),
\]

where the phonon occupancy factor, \(n(\vec{k})\), is the Planck distribution of Eq. 8.16 at the temperature of interest and \(\epsilon_i = \hbar \omega_{\vec{k}}^\ast\).

\[
n(\vec{k}) = \frac{1}{e^{\hbar \omega_{\vec{k}}/k_B T} - 1}.
\]

The signs in the factor \(n(\vec{k}) + \frac{1}{2} \pm \frac{1}{2}\) in Eq. 6.52 are such that it is \(n(\vec{k})\) for phonon annihilation, and \(n(\vec{k}) + 1\) for phonon creation – it is always possible to create a phonon, even at \(T = 0\) when no phonon excitations are present. The \(\delta(Q + \vec{k} \pm \vec{g})\) and \(\delta(\omega - \omega(\vec{k}))\) enforce the conservation of momentum and energy on the neutron scattering process. The phonons on different branches must be considered separately in Eq. 6.52, and it is traditional to add a "branch index," sometimes denoted \(j\), to \(\omega(\vec{k})\), since more than one \(\omega_j\) may correspond to a specific \(\vec{k}\).

Why should the expansion of an evidently classical exponential in Eq. 6.51 produce the series of neutron-phonon interactions? In the classical limit, the scattering of every neutron is inelastic, generating heat composed of a number of phonons with total energy, \(E\), the energy transfer from the neutron. For classical scattering, an atom of mass \(m\) will recoil with velocity \(v = \hbar Q/m\), with

---

\(^8\)Perhaps they exist as quantum beats of the wavetrains, but we ignore them because we do not have sufficient time resolution at the detector to sense them.

\(^9\)A new factor is the ratio \(k_i/k_f\), which expresses the effect on flux caused by the rate at which neutrons leave the sample. Compared to an elastically-scattered neutron, fewer neutrons per second will pass into an increment of solid angle \(d\Omega\) if they are slowed to smaller values of \(k_f\). Also, the use of solid angle eliminates the need for the factor \(r^{-2}\).
momentum transfer $\hbar Q$, and energy transfer

$$E_R = \frac{\hbar^2 Q^2}{2m}.$$ (6.54)

In the quantum limit where $Q$ is small, the energy recoil may or may not occur, depending on whether phonons are created or not. The total scattering, $S(Q, E)$, is the sum of components, $\sum_{n=0}^{\infty} S_n(Q, E)$, from neutrons scattered after creating different numbers, $n$, of phonons in the sample.\(^\text{10}\) The thermal energy is

$$k_B T = 2ma^2 \langle u^2 \rangle$$ (6.55)

for one harmonic mode of frequency $\omega$. From Eqs. 6.54 and 6.55

$$Q^2 \langle u^2 \rangle = \frac{E_R}{\hbar \omega} \frac{k_B T}{\hbar \omega}$$ (6.56)

This is the quantity in which which the multiphonon expansion is performed (Eq. 6.51). From Eq. 6.56 we see that the relative intensities of the sequence of phonon scattering terms scale with 1) the ratio of the recoil energy to the oscillator energy, and 2) the ratio of temperature to the oscillator energy. When the first ratio is small, quantum mechanics requires that some recoils excite the oscillator, but most do not. The same is true for temperature (and at low temperatures we of course have low heat capacity because the oscillator is usually not excited).

It remains to get the spectral shape of each order of the multiphonon scattering. The spectrum for one-phonon scattering weights more heavily the low-energy modes because they have larger amplitudes of motion, providing a factor of $g(\varepsilon)/\varepsilon$. The number of phonons is the Planck distribution $1/[\exp(\varepsilon/k_B T) - 1]$, and the two factors provide the shape of the 1-phonon profile, $A_1(\varepsilon)$:

$$A_1(\varepsilon) = \frac{g(\varepsilon)}{\varepsilon} \frac{1}{e^{\varepsilon/k_B T} - 1}. \quad (6.57)$$

When two phonons are created simultaneously, the total spectrum of energies is the convolution of the 1-phonon profile with the 1-phonon profile.\(^\text{11}\) The 2-phonon spectrum is the convolution of two 1-phonon profiles,

$$A_2(E) = A_1 \ast A_1 = \int_{-\infty}^{\infty} A_1(E - E') A_1(E') \, dE', \quad (6.58)$$

\(^{10}\) Phonon annihilation is handled by extending the range of $E$ to negative values for each $S_n(Q, E)$.

\(^{11}\) Consider each phonon excitation to be a random variable with a probability distribution of $A_1$. The sum of two random variables has a distribution that is the convolution of the probability distributions $A_1 \ast A_1$. 
The \( n \)-phonon profile is the convolution of another 1-phonon profile with the \((n-1)\)-phonon profile

\[
A_n(\epsilon) = A_1 * A_{n-1} = \int_{-\infty}^{\infty} A_1(\epsilon - \epsilon') A_{n-1}(\epsilon') \, d\epsilon'.
\]  

(6.59)

Starting with \( A_1 \), we can generate the spectral shapes of all orders of multi-phonon scattering by the systematic application of Eq. 6.59.

The total scattering is the sum of these spectral profiles, weighted by the corresponding terms of Eq. 6.51

\[
S(Q, E) = \sum_{n=0}^{\infty} \exp(-2W) \frac{(2W)^n}{n!} A_n(E),
\]  

(6.60)

where we have defined \( S(Q, E) \) as the double-differential cross-section \( d^2\sigma/d\Omega dE \) without prefactors.\(^{12}\)

In much experimental work, we seek to isolate the single scattering profile, \( A_1(E) \), from the measured \( S(Q, E) \), and then determine the phonon DOS \( g(E) \) by Eq. 6.57. One way to correct for multiphonon scattering is by iteration on an initial guess of the phonon DOS. With an initial guess at the DOS or \( A_1(E) \) (often the \( S(Q, E) \) with a constant background removed), the various \( A_2(E), A_3(E), A_4(E) \)... are calculated and subtracted from \( S(Q, E) \) to give a better \( A_1(E). \)

Iteration is complete when there is a good match between the measured \( S(Q, E) \) and the \( S(Q, E) \) recalculated with Eq. 6.60. When the multiphonon scattering is weak, perhaps only two iterations are necessary. When the multiphonon scattering is strong, the procedure becomes difficult and slow. If there is strong multiphonon scattering, however, the experiment is moving into the limit of classical scattering, so isolating the single phonon spectrum becomes more challenging. Section 7.3.2 develops further these concepts of multiphonon scattering.

6.3.3 Neutron Weighting

Strictly speaking, the phonon DOS from the procedure outlined above is not the true phonon DOS, but rather the neutron-weighted DOS. For an elemental scatterer, the neutron-weighted DOS is identical to the phonon DOS. This is not the usual case for a sample that contains more than one type of atom. The neutron-weighted phonon DOS is

\[
g_{NW}(\epsilon) \propto \sum_d g_d(\epsilon) \exp(-2W_d) \exp(2W) \frac{\sigma_d}{m_d}
\]  

(6.61)

\(^{12}\)This \( S(Q, E) \), called the “scattering law,” is normalized so that it is the double Fourier transform of the Van Hove correlation function \( G(\vec{r}, t) \).
where \( \exp(-2W_d) \), \( \sigma_d \) and \( M_d \) are the Debye-Waller factor, total scattering cross-section and mass of atom \( d \). The \( g_d(\varepsilon) \) are the phonon partial DOS functions of Eq. 8.62. The Debye-Waller factor is an explicit function of \( g_d(\varepsilon) \). The term \( \exp(2W) \) is the average Debye-Waller correction; it is calculated from the self-consistent neutron-weighted DOS. At low temperatures, the factor \( \exp[2(W - W_d)] \) is approximately unity.

For the case where \( \sigma_d/m_d \) is the same for all species \( d \), \( g_{NW}(\varepsilon) \approx g(\varepsilon) \), but in general obtaining the true phonon DOS from the neutron-weighted phonon DOS requires a full analysis of the lattice dynamics. This can be performed by simulational procedures or by some types of analytical methods. The neutron-weight correction as well as other approximations involved in the data analysis can be overcome by fitting a dynamics model to the neutron-scattering data directly. Although this approach is both scientifically and computationally demanding, we foresee no better method for extracting the vibrational dynamics from inelastic neutron scattering measurements.

**Further Reading**

The contents of the following are described in the Bibliography.


**Problems**

1. Suppose the scattering factor distribution is a rectangular function in time:

   \[
   f(\tau) = \begin{cases} 
   \frac{1}{t} f_0 & \text{when } 0 < \tau < t, \text{ and} \\
   0 & \text{otherwise.}
   \end{cases}
   \]

   \[ (6.62) \]

   \[ (6.63) \]

   Calculate the self-correlation function \( G_s(\tau) \), and the scattered intensity, \( I(\omega) \).

   *(Hint: See section 9.4.1 in main text. You can sample time in intervals \( \delta t \) if you like.)*
2.) Is it possible to have a self-correlation function $G_s(\tau)$ of the form:

$$G_s(\tau) = \begin{cases} 
\frac{1}{t} g_0 & \text{when } 0 < \tau < t, \\
0 & \text{otherwise}
\end{cases} \quad \text{(6.64)}$$

(a) If yes, give an example. If not, why not?

(b) For very long $t$ and short $\tau$ the practical situation may be different. If so, how?

3.) Make graphs of the self-correlation functions $G_s(t)$ for the three cases of Figs. 6.2, 6.3, 6.4.

4.) Perhaps with the help of Fig. 6.6,

(a) Propose two paths to get from the Van Hove function $G(R, \tau)$ to the diffraction pattern $I(Q)$.

(b) Suppose you have $f(r)$ and $f(t)$. Why can you not obtain a rigorous $f(r,t)$ using some combination of $f(r)$ and $f(t)$?

(c) Suppose you have $I(Q)$ and $I(\omega)$. Why can you not obtain a rigorous $S(Q, \omega)$ using some combination of $I(Q)$ and $I(\omega)$? List a couple of things you can learn about $S(Q, \omega)$ if you have both $I(Q)$ and $I(\omega)$.

(d) Suppose you have $I(Q)$ and $I(\omega)$, and suppose that the scattering is incoherent. How could you construct an estimate of the measured intensity $I(Q, \omega)$?
Chapter 7

Fundamental Theory of Neutron Scattering

7.1 Basic Quantum Mechanics of Neutron Scattering

This chapter develops a more rigorous treatment of nuclear scattering than was presented in the previous Sect. 6.3. It starts with Fermi’s golden rule, and avoids the explicit use of wavefunctions for the scatterer (i.e., it does not require phonon solutions like $u_\alpha(q, t)$ of (6.49)).

7.1.1 Fermi’s Golden Rule

Fermi’s golden rule gives the transition rate from an initial state to a final state, $W$, at time $t'$:

$$W(t') = \frac{2\pi}{\hbar} \left| \langle \Psi_f(\vec{r}, t') | V(\vec{r}, t') | \Psi_i(\vec{r}, t') \rangle \right|^2.$$  

(7.1)

For nuclear scattering of a neutron, the states $|\Psi\rangle$ include coordinates for the neutron and coordinates for the crystal. The interactions between them are through nuclear forces of very short range, compared to the thermal displacements of the atoms in the crystal. The potential in (7.1), $V_{\text{nuc}}(\vec{r}, t')$, moves during thermal vibrations over distances that are large compared to the dimensions of the potential itself. We will use a Fermi pseudopotential as in (6.40), with $\delta$-functions located at the instantaneous positions of the nuclei.\(^1\)

Although the Fermi pseudopotential moves with the vibrating nuclei, its strength (the scattering length $b$) remains unchanged. This conveniently allows

\(^1\)The electrons of the atom, and the chemical bondings between atoms, are not affected by the neutron. There are magnetic scatterings of neutrons from unpaired electrons, but these are considered later in this chapter.
us to first use basic conservations of momentum and energy to understand general features of inelastic neutron scattering, but later we will employ phonon excitations to present a more complete picture of thermal neutron scattering from crystals. We can therefore separate the state $|\Psi(t', t)\rangle$ into a lattice part $|\lambda_i\rangle$ and a neutron part $|\vec{k}_i\rangle$:

$$|\Psi_i(t', t)\rangle = |\lambda_i(\vec{r}_{nu}, t)\rangle |\vec{k}_i(\vec{r}_{ne}, t)\rangle,$$

(7.2)

where the independent coordinates $\vec{r}_{nu}$ and $\vec{r}_{ne}$ refer to the positions of the nucleus and neutron. We have assumed the neutron states are plane-wave states characterized by the wavevector $\vec{k}_i$ (as in (1.49) and (1.68), but here $k_i \neq k_j$). In practice, the $|\rangle |\rangle$ in (7.1) denotes an integration over all positional coordinates at the instant $t'$ when $W$ is evaluated. To get the total probability of the transition, $P_{i\rightarrow f}$, we integrate this over all times when the two states interact in the presence of the perturbing potential $V(\vec{r}', t)$:

$$P_{i\rightarrow f} = \frac{2\pi}{\hbar} \int \langle \lambda_i(\vec{r}', t) | \vec{k}_i(\vec{r}', t) | V(\vec{r}', t) | \lambda_i(\vec{r}', t) \rangle \langle \vec{k}_i(\vec{r}', t) | \rangle \langle \vec{k}_i(\vec{r}', t) | V(\vec{r}', t) | \lambda_i(\vec{r}', t) \rangle \langle \vec{k}_i(\vec{r}', t) | d\vec{r}' ,$$

(7.3)

where we have substituted (7.2) into (7.1), written out the $|\vec{r}'$, and integrated over all times.

Recall that the time evolution of the state of the scatterer is:

$$|\psi(t)\rangle = e^{-i\frac{\Delta E}{\hbar}}|\psi(t=0)\rangle ,$$

(7.4)

$$\langle \psi(t)| = \langle \psi(t=0)|e^{i\frac{\Delta E}{\hbar}} .$$

(7.5)

We assume that the states of the crystal $|\langle i\rangle\rangle$ are eigenstates with specific energies $\{\epsilon_i\}$, so we can simplify (7.4) and (7.5) using $\epsilon_i = \hbar\omega_i$ as:

$$|\lambda_i(0)\rangle = e^{-i\omega_i t}|\lambda_i(0)\rangle ,$$

(7.6)

$$\langle \lambda_i(0)| = \langle \lambda_i(0)|e^{i\omega_i t} .$$

(7.7)

The general formulation makes use of these expressions because they allow us to work with states of the crystal at $t = 0$ such as $|\lambda_1(0)\rangle$ and $|\lambda_2(0)\rangle$, which are constant and can be pulled out of any integration over time. Similarly for the position evolution of plane-wave states of the neutron $|\langle k\rangle\rangle$, which have constant momentum:\footnote{Plane waves prove their convenience here. For other states we would have to integrate over $k$-space.}

$$|\vec{k}_i(\vec{r})\rangle = e^{-i\vec{k}_i \cdot \vec{r}}|\vec{k}_i(\vec{r}=0)\rangle ,$$

(7.8)

$$\langle \vec{k}_i(\vec{r}| = \langle \vec{k}_i(\vec{r}=0)|e^{i\vec{k}_i \cdot \vec{r}} .$$

(7.9)
Returning to the evaluation of (7.3), into it we substitute (7.6), (7.7) and (7.8), (7.9):

\[
P_{1\to t} = \frac{2\pi}{\hbar} \int \langle \lambda_i(0)| e^{i\omega t} \langle \vec{k}_i(0)| e^{i\vec{P}_i \cdot \vec{r}_i} V(\vec{r}, t) e^{-i\vec{k}_f \cdot \vec{r}_f(0)} \rangle e^{-i\omega t|\lambda_i(0)} \rangle \langle \lambda_i(0)| e^{i\omega t} \langle \vec{k}_i(0)| e^{i\vec{P}_i \cdot \vec{r}_i} V(\vec{r}, t) e^{-i\vec{k}_f \cdot \vec{r}_f(0)} \rangle e^{-i\omega t|\lambda_i(0)} \rangle \ dt ,
\]

where the operators \( \vec{r} \) refer to the neutron coordinates, and the (0) refer to both \( t = 0 \) and \( \vec{r} = 0 \) (although we are not concerned about the time-dependence of the neutron wavefunction or the position-dependence of the crystal excitation).

The notation (0) refers to both \( t = 0 \) and \( \vec{r} = 0 \), since we made use of the relations (7.6)–(7.9). At \( t = 0 \), \( \vec{r} = 0 \), the phase factors, \( e^{i|\vec{r} - \vec{r}'|} \), of the plane-wave states of (1.49) and (1.68) are equal to 1. Therefore:

\[
\langle \vec{k}_i(0)| \vec{k}_i(0) \rangle = 1 .
\]

We define the difference in frequency as \( \omega \):

\[
\omega_f - \omega_i \equiv \omega .
\]

Substituting (7.12) and (7.13) into (7.11):

\[
P_{1\to t} = \frac{2\pi}{\hbar} \int \sum_j e^{i\omega t} b_j \langle \lambda_i(0)| e^{i\vec{P}_i \cdot \vec{r}_i} e^{-i\vec{k}_f \cdot \vec{r}_f(0)} \rangle \langle \lambda_i(0)| e^{i\omega t} \langle \vec{k}_i(0)| e^{i\vec{P}_i \cdot \vec{r}_i} e^{-i\vec{k}_f \cdot \vec{r}_f(0)} \rangle \ dt ,
\]

The integration is over all times, so we can redefine \( t \) as a difference between scattering times:

\[
t \equiv t_j - t_k ,
\]
and likewise we define the scattering vector $\vec{Q}$:

$$Q \equiv \vec{k}_i - \vec{k}_f . \quad (7.16)$$

Substituting these differences into (7.14):

$$P_{i \rightarrow f} = \frac{2\pi}{\hbar} \int e^{\text{int}} \sum_j b^*_j \langle \lambda_i(0) | e^{i\vec{Q}\cdot\vec{R}_j(t)} | \lambda_f(0) \rangle^* \langle \lambda_f(0) | e^{-i\vec{Q}\cdot\vec{R}_k(0)} | \lambda_i(0) \rangle \, dt , \quad (7.17)$$

Now we consider how scattering changes the state of the crystal, the $\{ | \lambda(0) \rangle \}$. First consider the final states, $\{ | \lambda_f(0) \rangle \}$, and work with them in a slightly-rearranged form of (7.17)

$$P_{i \rightarrow f} = \frac{2\pi}{\hbar} \int e^{\text{int}} \sum_f \sum_k b^*_f b_k \langle \lambda_i(0) | e^{i\vec{Q}\cdot\vec{R}_j(t)} | \lambda_f(0) \rangle \langle \lambda_f(0) | e^{-i\vec{Q}\cdot\vec{R}_k(0)} | \lambda_i(0) \rangle \, dt , \quad (7.18)$$

We do not have control over which final state is obtained, and in principle all acceptable final states will occur over the duration of a long experiment. In effect, an experiment sums over final states, but the final states are assumed to form a complete set:

$$\sum_f | \lambda_f(0) \rangle \langle \lambda_f(0) | = 1 , \quad (7.19)$$

for each term in the double sum (7.17). Equation (7.19) simplifies (7.18)

$$P_{i \rightarrow f} = \frac{2\pi}{\hbar} \int e^{\text{int}} \sum_f \sum_k b^*_f b_k \langle \lambda_i(0) | e^{i\vec{Q}\cdot\vec{R}_j(t)} | \lambda_f(0) \rangle \langle \lambda_f(0) | e^{-i\vec{Q}\cdot\vec{R}_k(0)} | \lambda_i(0) \rangle \, dt , \quad (7.20)$$

where going from $P_{i \rightarrow f}$ to $P_{i \rightarrow f,i}$ means that we are calculating the total scattering out of the initial state (not just from one $f$ as in (7.18)).

Next, consider the initial states $\{ | \lambda_i(0) \rangle \}$. The initial states are set by thermodynamics. Temperature will alter the distribution of initial states by the appropriate thermodynamic distribution, i.e., a Bose–Einstein distribution for phonons. Instead of writing this distribution $n(\varepsilon)$ directly, we define the brackets $\langle \rangle$ to mean the thermodynamic average. This is arguably just a notational change to (7.20) that is now

$$P_{i \rightarrow f} = \frac{2\pi}{\hbar} \int e^{\text{int}} \sum_f \sum_k b^*_f b_k \langle \lambda_i(0) | e^{i\vec{Q}\cdot\vec{R}_j(t)} | \rangle \langle \lambda_i(0) | e^{-i\vec{Q}\cdot\vec{R}_k(0)} | \rangle \, dt . \quad (7.21)$$

We make use of the prefactors described in Sect. 6.3 to convert the transition probability into a cross-section:

$$\frac{d^2\sigma}{d\Omega\,dE} = \frac{k_i}{\hbar} \frac{(2\pi)^3}{2V_0} \sum_f \sum_k b^*_f b_k \int e^{\text{int}} \langle \lambda_i | e^{i\vec{Q}\cdot\vec{R}_j(t)} | \rangle \langle \lambda_i | e^{-i\vec{Q}\cdot\vec{R}_k(0)} | \rangle \, dt . \quad (7.22)$$
Equation (7.22) is the most general result that can be obtained from Fermi’s Golden Rule. At its heart is a thermodynamic average of phase factors from scattering by atoms $j$ and $k$ at their different positions in space for different snapshots in time. Note that the space coordinates for the two atoms are generally not evaluated at the same time. If there were no time dependence to the atom positions, the Fourier transform would yield a delta function $\delta(\omega - 0)$, indicating pure elastic scattering. If there were no spatial periodicities, the thermodynamic average over all phase factors would not produce any constructive or destructive interferences at different angles, so there would be no structure of the cross section in solid angle, $\Omega$.

The topic of this book is inelastic scattering, so we assume motion of the nuclei (the scattering centers), and we will be alert to relations between the two phase factors (the exponentials in (7.22)) in space and in time. We need to address the thermal spread of atom positions over time. Even the analysis of small displacements requires confronting some subtleties of quantum mechanics, but later our treatment in Sect. 7.2.3 will assume the distributions of atom displacements are Gaussian functions, or at least the displacements are small. We will also assume that the scattering centers have the translational periodicity of a crystal. This is a more restrictive assumption about the sample, and can be misleading in cases of disordered solids. Therefore it is sometimes important to return to (7.22) for guidance on the scattering problem, since its only assumption is that the neutron is scattered once.

There is, however, another very general and useful feature of inelastic neutron scattering that depends only on thermodynamic equilibrium for the populations of the initial states of the scatterer. This principle of detailed balance is described next in Sect. 7.1.2.

### 7.1.2 Detailed Balance

The intensities of inelastic spectra depend on the ratio of energy transfer to temperature, at least in the usual case where the sample is in thermodynamic equilibrium before scattering. There is a detailed balance between the rates of two scattering processes, one with the creation of an excitation, and the other with the annihilation of the same excitation. Consider the temperature dependence of the positive and negative energy transfers between the neutron and the specimen. One extreme is when the sample is at a very low temperature (in practice, where $k_B T$ is much smaller than the energy resolution of the instrument). In this case there are no excitations present in the sample, so no scattering can occur with the annihilation of excitations. Excitations can be created by transfer of energy from the incident neutron, of course. At low temperatures the inelastic spectrum will have intensity on one side of the elastic peak, but no intensity on the other side. This is seen in Fig. 7.1, at 30 K, where there is no intensity for $S(-E)$, but a weak but reasonable spectrum is seen at $S(+E)$.

The other extreme is for very high temperatures where $k_B T$ is much larger than the energy transfer $|E|$. Because the probability of creating or annihilating
Figure 7.1: Incoherent inelastic scattering of phonons in fcc $^{57}$Fe$_{64}$Ni$_{36}$. Data were for equal incident flux, but different temperatures as labeled. Elastic peak at $\epsilon = 0$ rises to 15,000 counts in both spectra. Compare the relative intensites at $\pm \epsilon$, which are related by detailed balance. From (7.35), the ratio of the sharp features at $\pm 16$ meV should be 1.9 at 300 K and 600 at 30 K. (Incidentally, some multiphonon scattering is evident in the 300 K data above 37 meV.) These spectra were from nuclear resonant inelastic x-ray scattering of $^{57}$Fe with a high resolution monochromator at Sector 3 of the Advanced Photon Source (courtesy S.H. Lohaus and P. Guzman). The principles are similar to incoherent inelastic neutron scattering, but these are inelastic x-ray measurements.
one additional excitation such as a phonon makes little difference to the energy of the sample, we expect the measured intensities to be more similar at \( \pm E \). Scattering processes involving the creation or annihilation of excitations occur with similar probabilities when the sample is at high temperatures.

Each creation process has an inverse annihilation process, and the intensities of these two processes in an inelastic spectrum are related by a thermal factor that we show to be a simple Boltzmann factor. We will compare intensities at pairs of energies \( \pm E \), i.e., at \( S(-E) \) and \( S(+E) \). The systematics to be explained can be seen with Fig. 7.1.

The energy equivalence to 300 K is \( k_B 300 = 25 \text{ meV} \), and at 30 K it is 2.5 meV. The heights of each spectrum differ by a factor of \( e \) at these energies, i.e., at 300 K, with \( E \) in meV, \( S(+25) = 2.71828 S(-25) \), and at 30 K, \( S(+2.5) = 2.71828 S(-2.5) \). This ratio is visible at \( \pm 25 \text{ meV} \) for the 300 K spectrum but the 30 K spectrum is covered by the tails of the elastic line at 2.5 meV. Notice also the differences in the shapes of the two spectra on the phonon creation side of the elastic peak \( +E \). The 30 K spectrum is actually close to the shape of the phonon DOS itself, whereas the 300 K spectrum is enhanced at energies below approximately 25 meV, owing to the large phonon occupancy factor.

A picture inspired by quantum mechanics can account for the detailed balance of the intensities of spectral features at \( \pm E \) in Fig. 7.1. The density of states for excitations is symmetric, i.e., \( S(-E) = S(+E) \), because an excitation can be created or annihilated in each mode at \( E \). Suppose that the observed intensities are proportional to the thermal occupancies before scattering, which are the Planck factors \( n_{E,T} \)

\[
\begin{align*}
n_{E,T} &= \frac{1}{e^{\frac{E}{k_B T}} - 1}. \\
\end{align*}
\]  

(7.23)

The idea follows stimulated emission in optics, where both phonon absorption and phonon emission to the neutron increase with \( n_{E,T} \), but emission (phonon creation) has an extra spontaneous contribution of \( +1 \). This relates the intensities at \( \pm E \) as

\[
\begin{align*}
\frac{I(+E)}{I(-E)} &= \frac{n_{E,T} + 1}{n_{E,T}}. \\
\end{align*}
\]  

(7.24)

This Eq. 7.24 is easy to remember, and readily shows that as \( n_{E,T} \to 0 \) at low \( T \), only the right side of the spectrum will be observed (see 30 K spectrum in Fig. 7.1).

However, Eq. 7.24 is not quite so quantum mechanical as it may seem. Rearranging

\[
\begin{align*}
\frac{I(+E)}{I(-E)} &= \left( \frac{1}{e^{\frac{E}{k_B T}} - 1} + \frac{e^{\frac{E}{k_B T}} - 1}{e^{\frac{E}{k_B T}} - 1} \right) \left( e^{\frac{E}{k_B T}} - 1 \right), \\
\frac{I(+E)}{I(-E)} &= e^{\frac{E}{k_B T}}, \\
\end{align*}
\]  

(7.25)
and the intensities for phonon creation and annihilation are in the ratio of the simple Boltzmann factor $\exp(\frac{E}{k_B T})$. The Planck occupancy does express the quantum statistics of bosons, of course. Furthermore, a simple classical analysis might give the incorrect ratio of $I(+E)/I(-E) = \exp(\frac{2E}{k_B T})$ because spectral components at $\pm E$ are separated in energy by $2E$.

The condition of detailed balance follows from two assumptions:

- The probability of the initial state of the sample, the $|\lambda_i\rangle$ in (7.2), is as expected for thermodynamic equilibrium.

- The interaction operator for the transition probability, the $V(r', t')$ of (7.1), is Hermitian. (This is certainly true for the delta function (6.40) for the Fermi pseudopotential.)

To show the essence of the derivation of the detailed balance condition, assume the initial state $|\lambda_1\rangle$ exists, and consider the probability, $W'_{1\rightarrow 2}$, for a transition to a final state $|\lambda_2\rangle$, and the probability for the reverse transition, $W'_{2\rightarrow 1}$, from a pre-existing state $|\lambda_2\rangle$:

$$W'_{1\rightarrow 2} = |\langle \lambda_2 | V | \lambda_1 \rangle|^2,$$

$$W'_{2\rightarrow 1} = |\langle \lambda_1 | V | \lambda_2 \rangle|^2,$$

which are both products of a number with its complex conjugate:

$$W'_{1\rightarrow 2} = \langle \lambda_2 | V | \lambda_1 \rangle \left( \langle \lambda_2 | V | \lambda_1 \rangle \right)^*,$$

$$W'_{2\rightarrow 1} = \left( \langle \lambda_1 | V | \lambda_2 \rangle \right)^* \langle \lambda_1 | V | \lambda_2 \rangle.$$

For a Hermitian operator, recall that $V = (V^T)^* \equiv V^\dagger$. We use the transpose to operate on the other side of $V$, for which we use the complex conjugates of the bras and kets:

$$W'_{1\rightarrow 2} = \langle \lambda_2 | V | \lambda_1 \rangle \langle \lambda_1 | V | \lambda_2 \rangle,$$

$$W'_{2\rightarrow 1} = \langle \lambda_2 | V | \lambda_1 \rangle \langle \lambda_1 | V | \lambda_2 \rangle,$$

so:

$$W'_{1\rightarrow 2} = W'_{2\rightarrow 1} \equiv W'. $$

This result (7.32) is true so long as $V$ is Hermitian. Starting in the known states $|\lambda_1\rangle$ and $|\lambda_2\rangle$, the transition probabilities between these states are equal.

Now assume that the states $|\lambda_i\rangle$ are populated in thermodynamic equilibrium, differing by a Boltzmann factor. The measured cross sections are proportional to $W_{1\rightarrow 2}$ and $W_{2\rightarrow 1}$:

$$W_{1\rightarrow 2} = e^{-E_1/k_B T} Z^{-1} W_{1\rightarrow 2}' = \frac{e^{-E_1/k_B T}}{Z} W',$$

$$W_{2\rightarrow 1} = e^{-E_2/k_B T} Z^{-1} W_{2\rightarrow 1}' = \frac{e^{-E_2/k_B T}}{Z} W'.$$
where $Z$ is the partition function. Now that we have taken into consideration the fact that the initial states have probabilities as expected from thermodynamic equilibrium, we can relate the observed intensity for the transition $1 \rightarrow 2$ to the observed intensity for its reverse transition $2 \rightarrow 1$:

$$W_{1 \rightarrow 2} = e^{(E_2 - E_1)/k_BT} W_{2 \rightarrow 1}. \quad (7.35)$$

Suppose the state $|\lambda_2\rangle$ has an energy higher (more positive) than $|\lambda_1\rangle$, because the state $|\lambda_2\rangle$ has an extra excitation in the sample. The transition $1 \rightarrow 2$ is uphill energetically, and requires the neutron to transfer energy to create an excitation sample. Nevertheless, this transition is more intense experimentally because the initial state $|\lambda_1\rangle$ is more probable thermodynamically. Equation (7.35) shows that the intensity $W_{1 \rightarrow 2} > W_{2 \rightarrow 1}$ because $E_2 > E_1$ and the exponential is greater than 1. To clarify (7.35), we recognize that the difference in energy, $E = E_2 - E_1$, is the energy of the excitation in the solid. It is more traditional to write the condition of detailed balance as:

$$S(E) = e^{E/k_BT} S(-E). \quad (7.36)$$

where $E$ is the energy of the excitation in the solid, and the argument $-E$ corresponding to $W_{2 \rightarrow 1}$ signifies that $S(-E)$ is on the phonon annihilation side of the elastic peak in the spectrum. Detailed balance remains valid when a single scattering creates or annihilates multiple excitations – a detailed balance between forward and reverse processes still exists because the thermodynamic probabilities of the required initial states are set by $E$.

A practical use of detailed balance is to check the quality of experimental data. For example, in a spectrum measured at 300 K, equivalent to 25 meV, the intensities at $\pm 25$ meV on the two sides of the elastic peak must be in the ratio of $e^{-1}$. If this were not true, we might suspect instrument artifacts, such as differences in sensitivity or resolution. A noise background could also cause measured data to violate the condition of detailed balance, and perhaps detailed balance could be exploited to help subtract some sources of background from experimental data. We warn the reader, however, about such simple interpretations with data from a time-of-flight chopper spectrometer. The value of $Q$ varies across the energy scale of data from a particular detector, and the relationship is not symmetric, i.e., $Q(E) \neq Q(-E)$. When multiphonon scattering is strong, and the Debye–Waller factor is significant, detailed balance will not be observed in the experimental data unless they are rebinned for constant $Q$.

### 7.1.3 Crystalline Periodicity

The translational periodicity of crystals allows the reduction of the double sum in (7.22) to a single sum. We separate the atom position operators, $\vec{R}_j(t)$, into static, $\vec{x}_j$, and time-varying, $\vec{u}_j(t)$, components:

$$\vec{R}_j(t) = \vec{x}_j + \vec{u}_j(t), \quad (7.37)$$
where the static positions were broken into lattice vectors (index $l$) and basis vectors (index $\kappa$) in (7.37). The exponentials in (7.22) refer to pairs of atoms separated by a distance $\vec{x}_j - \vec{x}_k$. In an infinite crystal, where all unit cells are equivalent, these exponentials cannot depend on the absolute position of the unit cell, but only on the lattice translation vector $\vec{\delta}$. Any one of the $N$ unit cells can be taken as the origin, and the terms from the $l$th neighbor must be the same. This distance, $\vec{\delta}$, has no time dependence, and is a constant. It therefore commutes through the other distance operators, and we can write:

$$\frac{d^2\sigma}{d\Omega\,dE} = \frac{k_i}{k_f} \frac{(2\pi)^3 N|b|^2}{2V_0} \sum_l e^{i\vec{Q}\cdot\vec{x}_l} \int e^{i\omega t} \left( e^{i\vec{Q}\cdot\vec{u}_l(t)} e^{-i\vec{Q}\cdot\vec{u}_l(0)} \right) dt \quad (7.38)$$

Here, for simplicity we have also assumed all nuclei are the same so $b_j = b_k \equiv b$, and we have assumed a simple lattice without a basis. The result in (7.38) accounts for the interactions of waves scattered from pairs of nuclei, recognizing that their instantaneous displacements may differ by a phase factor.

This difference in phase factor gives a constant prefactor $e^{i\vec{Q}\cdot\vec{x}_l}$. The other exponentials are not constants, however, and need to be handled with care. The temptation would be to combine the exponentials into a single factor such as $e^{i\vec{Q}\cdot(\vec{R}_j(t) - \vec{R}_j(0))}$. Unfortunately, this tempting step would be incorrect, except for classical systems. The quantum argument, described next, is subtle.

### 7.2 More Subtle Quantum Behavior

#### 7.2.1 Commutation Relations

Calculating the expectation value of an operator $\mathcal{A}$ using the left- and right-hand sides of (7.4) and (7.5) gives:

$$\langle \psi(t)|\mathcal{A}|\psi(t)\rangle = \langle \psi(t=0)|e^{+i\mathcal{H}t/\hbar}\mathcal{A}e^{-i\mathcal{H}t/\hbar}|\psi(t=0)\rangle \quad (7.39)$$

Evidently the time-dependence of the matrix element can be transferred from the state functions $\psi$ to the operator $\mathcal{A}$ by replacing the operator by $e^{+i\mathcal{H}t/\hbar}\mathcal{A}e^{-i\mathcal{H}t/\hbar}$. This moves us into the “Heisenberg picture” of quantum mechanics where state functions are fixed, but the time-dependence is in the operator. The motivation for putting the dynamics into the operators is as follows. In passing from classical mechanics to quantum mechanics, we replace the position vector $\vec{R}$ with an operator, denoted $\hat{\vec{R}}(t)$. In particular, we will alter soon work with phase factors that are a time-dependent operators:

$$e^{i\vec{Q}\cdot\hat{\vec{R}}(t)} \rightarrow e^{i\vec{Q}\cdot\hat{\vec{R}}} \quad (7.40)$$

Changing $\vec{R}$ to the operator $\hat{\vec{R}}$ leads to a subtlety in calculating the intensity from the wave amplitude. It turns out that the operator $\hat{\vec{R}}(0)$ does not commute
with the operator $\vec{R}(t)$ for the same atom at a different time. The operators are related as:

$$\vec{R}(t) = e^{+i\mathcal{H}t/\hbar} \vec{R}(0) e^{-i\mathcal{H}t/\hbar},$$  \hspace{1cm} (7.41)$$

$$e^{-i\mathcal{H}t/\hbar} \vec{R}(t) = \vec{R}(0) e^{-i\mathcal{H}t/\hbar},$$  \hspace{1cm} (7.42)$$

and likewise for the exponentiated operators:

$$e^{i\mathcal{Q} \cdot \vec{R}(t)} = e^{i\mathcal{Q} \cdot \vec{R}(0)} e^{-i\mathcal{H}t/\hbar},$$  \hspace{1cm} (7.43)$$

$$e^{-i\mathcal{H}t/\hbar} e^{i\mathcal{Q} \cdot \vec{R}(t)} = e^{i\mathcal{Q} \cdot \vec{R}(0)} e^{-i\mathcal{H}t/\hbar}.$$  \hspace{1cm} (7.44)$$

We cannot interchange the order of $\vec{R}$ and $\mathcal{H}$ because $\mathcal{H}$ includes the momentum operator.

To get better expressions for calculating with these operators, we need a result about exponentiations of non-commuting operators. Operators in exponential functions are defined in terms of the power series expansion of the exponential. Consider the exponentiation of two operators $A$ and $B$:

$$\exp(A) = 1 + A + \frac{1}{2} AA + \ldots,$$  \hspace{1cm} (7.45)$$

$$\exp(-B) = 1 - B + \frac{1}{2} BB + \ldots,$$  \hspace{1cm} (7.46)$$

Now take the product, including all terms to the second order:

$$\exp(A) \exp(-B) = \left[1 + A + \frac{1}{2} AA + \ldots\right] \left[1 - B + \frac{1}{2} BB + \ldots\right].$$  \hspace{1cm} (7.47)$$

Group the terms in order of the $n$th power of operators $A$ and $B$ as $O^n$, and stop at the second order. (This truncations may seem like an approximation now, but it proves exact after a physical fact is brought out later.)

$$\exp(A) \exp(-B) = O^0 + O^1 + O^2,$$  \hspace{1cm} (7.48)$$

$$O^0 = 1,$$  \hspace{1cm} (7.49)$$

$$O^1 = A - B,$$  \hspace{1cm} (7.50)$$

$$O^2 = \frac{1}{2} AA - AB + \frac{1}{2} BB,$$  \hspace{1cm} (7.51)$$

where we have been fastidious about keeping the operator $A$ to the left of the operator $B$, because they may not necessarily commute.

Now evaluate the exponentiation of the sum $A + B$, again to second order in the operators:

$$\exp(A - B) = 1 + (A - B) + \frac{1}{2} (A - B)(A - B),$$  \hspace{1cm} (7.53)$$

$$\exp(A - B) = 1 + A - B + \frac{1}{2} [AA - AB - BA + BB],$$  \hspace{1cm} (7.54)$$
Note that (7.52) and (7.54) are unequal when $A$ does not commute with $B$. We need one more exponentiation expression to make an equality. It is, to second order in the operators

$$\exp\left(\frac{1}{2}[B, A]\right) = 1 + \frac{1}{2}[B, A], \quad (7.55)$$

$$\exp\left(\frac{1}{2}[B, A]\right) = 1 + \frac{1}{2}(BA - AB). \quad (7.56)$$

Multiplying the expressions for (7.54) and (7.56)

$$\exp(A - B) \exp\left(\frac{1}{2}[B, A]\right) = \left[1 + A - B + \frac{1}{2}(AA - AB - BA + BA)\right] \times \left[1 + \frac{1}{2}(BA - AB)\right] \quad (7.57)$$

and considering terms only to second order in the operators

$$\exp(A - B) \exp\left(\frac{1}{2}[B, A]\right) = 1 + A - B + \frac{1}{2}AA - AB + \frac{1}{2}BB, \quad (7.58)$$

This is exactly (7.52), so we conclude

$$\exp(A - B) \exp\left(\frac{1}{2}[B, A]\right) = \exp(A) \exp(-B). \quad (7.59)$$

This is the “Weyl identity.”

Not so fast, you may say. What about the terms $O^3$ and higher? The next subsection shows how these higher terms can be identically zero when the commutator $[B, A]$ is a constant, and gives a practical case where this happens. If you are satisfied with the approach in this section, you can skip the next section because it is not crucial for the development.

### 7.2.2 An Operator Identity (Optional Section)

Another useful identity can be obtained from an expansion for projection operations. We take the expression $e^{A}Be^{-A}$ and write out expansions for the exponentiated operators

$$e^{A} = 1 + A + \frac{1}{2!}A^2 + \frac{1}{3!}A^3 + ..., \quad (7.60)$$

$$e^{-A} = 1 - A + \frac{1}{2!}A^2 - \frac{1}{3!}A^3 + ..., \quad (7.61)$$

and develop a scheme for working with all terms to higher order beyond the cubic. Expanding the typical projection expression in the Heisenberg representation

$$e^{A}Be^{-A} = (1 + A + \frac{1}{2!}A^2 + \frac{1}{3!}A^3 + ...)B(1 - A + \frac{1}{2!}A^2 - \frac{1}{3!}A^3 + ...), \quad (7.62)$$
we seek to group all terms in the same order of operators, denoted $O^0$, $O^1$, $O^2$, ... such as the lowest ones that are easy to see in Eq. 7.62

\[
O^0 = B, \quad \tag{7.63}
\]

\[
O^1 = BA - AB = [B, A]. \quad \tag{7.64}
\]

It turns out that the big trick is already evident in the expression (7.64) for $O^1$. To go from $O^0$ to $O^1$, we commute $O^0$ with $A$. This works to all higher orders. For example, go from $O^1$ to $O^2$, first laboriously by writing out all the terms to second order

\[
O^2 = (1 + A + \frac{1}{2!}A^2)B(1 - A + \frac{1}{2!}A^2), \quad \tag{7.65}
\]

\[
O^2 = \frac{1}{2!}BA^2 - ABA + \frac{1}{2!}A^2B, \quad \tag{7.66}
\]

where the ends of the underlines in (7.65) denote the product factors that contribute terms to second order in $A$ in (7.66). Equation (7.66) can be written alternatively as

\[
O^2 = \frac{1}{2!}(BA^2 - ABA - A^2B), \quad \tag{7.67}
\]

\[
O^2 = \frac{1}{2!}((BA - AB)A - A(BA - AB)), \quad \tag{7.68}
\]

\[
O^2 = \frac{1}{2!}[B, A], A \tag{7.69}
\]

The same trick works for going from $O^2$ to $O^3$

\[
O^3 = \frac{1}{3!}[[B, A], A, A], \quad \tag{7.70}
\]

What we do by commuting with $A$, perhaps clear in (7.68) is making a term with a power of $A$ to the right, and a term with the opposite sign with $A$ to the left. This is just what we need when working the expansion to higher orders, using the method of underlines in (7.65). It is a good exercise to obtain $O^3$ by the method of (7.65), but to the next order there will be four underlines that denote the terms for $O^3$.

This elegance can be hard to use, but it is valuable in our case of position and momentum operators, $X$ and $P$. The important point is that the commutation of these operators is a constant,

\[
[X, P] = i\hbar. \quad \tag{7.71}
\]

Any later commutation with a constant is zero, because the two terms in the commutator are equal. Therefore, all terms beyond $O^1$ vanish identically. For
example, consider
\[ e^{-iPX/\hbar}Xe^{iPX'/\hbar} = X + [X, iPX'/\hbar], \]
\[ e^{-iPX/\hbar}Xe^{iPX'/\hbar} = X + \frac{i}{\hbar}(XPX' - PX'X), \]
\[ e^{-iPX/\hbar}Xe^{iPX'/\hbar} = X + \frac{i}{\hbar}(i\hbar + PX)X' - PX'X', \]
\[ e^{-iPX/\hbar}Xe^{iPX'/\hbar} = X - X', \]
where we substituted the commutation result (7.71) in the small ( ) in (7.74). This is an exact result when the commutator \([B, A]\) is a constant – the higher terms obtained by commuting \([B, A]\) with \(A\) give zero identically.

### 7.2.3 Gaussian Thermal Averages

Now that we have the expression (7.59), we can use it to rearrange (7.38) into a form that shows intensity contributions from different numbers of phonon excitations. To clarify the next steps, it is traditional to make the definitions:
\[ \vec{U} \equiv -i\vec{Q} \cdot \vec{u}_0(0), \]
\[ \vec{V} \equiv i\vec{Q} \cdot \vec{u}_l(t). \]
Now we substitute (7.76) and (7.77) into (7.59):
\[ \exp(\vec{U})\exp(\vec{V}) = \exp(\vec{U} + \vec{V})\exp\left(\frac{1}{2}[\vec{U} \cdot \vec{V} - \vec{V} \cdot \vec{U}]\right). \]

At this point we need to take a thermal average (the \(\langle\rangle\) in (7.38)). In doing so, we make the assumption that the vibrational atom displacements are distributed with a Gaussian spread. The Gaussian distribution function can be used to average a squared displacement, \(X^2\):
\[ \langle X^2 \rangle = \int_{-\infty}^{\infty} X^2 \frac{1}{\sqrt{\pi}\sigma^2} e^{-X^2/\sigma^2} dX, \]
\[ \langle X^2 \rangle = \frac{1}{2}\sigma^2, \]
a standard result.

For comparison, we next average an exponential \(e^X\):
\[ \langle e^X \rangle = \int_{-\infty}^{\infty} e^X \frac{1}{\sqrt{\pi}\sigma^2} e^{-X^2/\sigma^2} dX, \]
by completing the square:
\[ -\left(\frac{X}{\sigma} - \sigma/2\right)^2 = -\left(X^2/\sigma^2 - X + \sigma^2/4\right), \]
so by adding and subtracting the last term of (7.82) in the exponential:

\[ \langle e^X \rangle = \int_{-\infty}^{\infty} \frac{1}{\sqrt{\pi\sigma^2}} e^{-(X/\sigma - \sigma/2)^2} e^{\sigma^2/4} \, dX , \]  
(7.83)

\[ \langle e^X \rangle = e^{\sigma^2/4} \int_{-\infty}^{\infty} \frac{1}{\sqrt{\pi\sigma^2}} e^{-(X/\sigma - \sigma/2)^2} \, dX , \]  
(7.84)

\[ \langle e^X \rangle = e^{\sigma^2/4} , \]  
(7.85)

where the last result was obtained by integrating the normalized Gaussian function. By comparing (7.80) and (7.85), we obtain:

\[ \langle e^X \rangle = e^{X^2/2} . \]  
(7.86)

Using (7.86), we obtain a factor with our operators \( \vec{U} \) and \( \vec{V} \):

\[ \langle \exp(\vec{U} + \vec{V}) \rangle = \exp\left(\frac{1}{2}(\vec{U} + \vec{V})^2\right) . \]  
(7.87)

This is the “Bloch identity” for Gaussian thermal averages, usually written more simply with \( C = \vec{U} + \vec{V} \). We use this result to obtain the thermal average of (7.78):

\[ \langle \exp(\vec{U} \exp(\vec{V})) \rangle = \exp\left(\frac{1}{2}(\vec{U} + \vec{V})^2\right) \exp\left(\frac{1}{2}(\vec{U} \vec{V} - \vec{V} \vec{U})\right) , \]  
(7.88)

\[ \langle \exp(\vec{U} \exp(\vec{V})) \rangle = \exp\left(\frac{1}{2}(\vec{U}^2 + \vec{V}^2 + \vec{U}\vec{V} + \vec{V}\vec{U} + \vec{U}\vec{V} - \vec{V}\vec{U})\right) , \]  
(7.89)

\[ \langle \exp(\vec{U} \exp(\vec{V})) \rangle = \exp\left(\frac{1}{2}(\vec{U}^2 + \vec{V}^2)\right) \exp(\vec{U}\vec{V}) . \]  
(7.90)

By examining (7.76) and (7.77), we can see that:

\[ \langle \vec{U}^2 \rangle = \langle \vec{V}^2 \rangle , \]  
(7.91)

because the average vibrational amplitudes do not change over time, and all unit cells are equivalent in the crystal. This allows a final simplification of (7.90):

\[ \langle \exp(\vec{U} \exp(\vec{V})) \rangle = \exp(\vec{U}^2) \exp(\vec{U}\vec{V}) . \]  
(7.92)

Equation (7.92) was obtained with the one assumption of a Gaussian thermal spread of atom positions. Even if this in not quite the case, (7.92) is expected to be valid when the atom displacements are small. By expanding both sides of (7.86), and recognizing that the thermal average of odd powers of \( X \) are zero, (7.86) seems a reasonable approximation.
Finally, we use (7.92) to rewrite (7.38) in a way that will later let us identify individual phonon scatterings. Making use of the definitions (7.76) and (7.77):

\[
\frac{d^2\sigma}{d\Omega dE} = \frac{k_i}{k_f} \frac{(2\pi)^3 N|b|^2}{2V_0} \sum_l e^{iQ_l \cdot x_l} \int e^{i\omega t} \exp(\langle U^2 \rangle) \exp(\langle \bar{U} V \rangle) \, dt.
\] (7.93)

Taking the thermal averages has removed much of the time dependence in our factors. The only remaining time dependence is within the \langle \bar{U} V \rangle factor, so we rewrite:

\[
\frac{d^2\sigma}{d\Omega dE} = \frac{k_i}{k_f} \frac{(2\pi)^3 N|b|^2}{2V_0} \sum_l e^{iQ_l \cdot x_l} \exp(\langle U^2 \rangle) \int e^{i\omega t} \exp(\langle \bar{U} V \rangle) \, dt.
\] (7.94)

Using (7.76), the factor \exp(\langle U^2 \rangle) in (7.94) becomes \exp(-\langle Q \cdot \bar{u}(0) \rangle^2). We recognize this as a Debye–Waller factor. To its right in (7.94) is a sum over pairs of atoms separated by \(x_l\). There is a phase associated with the atom separation in the pair. Suppose for the moment that there were no displacements of the atoms from their lattice sites. In this case the final exponential \(\exp(\langle \bar{U} V \rangle)\) would equal \(e^0 = 1\). This is the case of elastic scattering, and the sum of phase factors of (7.94) would reduce to that of the elastic scattering result of (6.48). It is the final factor, the Fourier transform of \(\exp(\langle \bar{U} V \rangle)\), that makes (7.94) interesting, and it is in a convenient form for further development.

### 7.3 Practical Expressions for Phonon Scattering

#### 7.3.1 Impulse Approximation

It is relatively easy to adapt the general result (7.22) to the case where the energy of the incident neutron is much higher than the characteristic excitations in the solid. We therefore ignore the interatomic interactions, and consider the collision of the neutron with a single nucleus at \(\vec{R}\). This approaches the problem of hitting a ball with a classical projectile, so we lose some features of wave mechanics.\(^3\) We do have to account for momentum and energy transfer, of course. The impulse approximation begins with the replacement of the operator:

\[
\begin{align*}
\vec{R}_j(t) & \rightarrow \vec{R}_j(0) + t \vec{v}_j, \\
\vec{R}_j(t) & \rightarrow \vec{R}_j(0) + \frac{t}{M_j} \vec{p}_j.
\end{align*}
\] (7.95, 7.96)

\(^3\)Another viewpoint is that we expect multiple excitations to occur in the solid. The effects of coherence are generally washed out when multiple excitations occur. For incoherent inelastic scattering as in Sect. 6.2, we expect to analyze the scattering by considering only a single nucleus at a time. We expect the excitations to be incoherent.
where we expect \( t \) to be small since the neutron is moving fast. Because the incoherent character of the multiple excitations suppresses the phase relationships between different scatterers, and we consider terms \( j = k \) in (7.22). It is tempting to substitute (7.96) directly into (7.22) to obtain:

\[
\frac{d^2 \sigma}{d\Omega \, dE} \, ? = ? \frac{k_i}{k_i} \frac{(2\pi)^3}{2V_0} \sum_j \sum_k b_j^* b_k \delta_{jk} \\
\times \int e^{i\omega t} \left( e^{iQ_0^j(0) + i/M_j \hat{p}_j} - e^{-iQ_0^j(0)} \right) dt ,
\]

(7.97)

and with \( \vec{R}(0) = \vec{R}(0) \):

\[
\frac{d^2 \sigma}{d\Omega \, dE} \, ? = \frac{k_i}{k_i} \frac{(2\pi)^3}{2V_0} \sum_j |b_j|^2 \int e^{i\omega t} \left( e^{i/M_j \hat{p}_j} - e^{-i/M_j \hat{p}_j} \right) dt .
\]

(7.98)

The missing piece in (7.98) is a phase factor associated with the energy gain of the scatterer. The energy gain is kinetic, \( E_{\text{kin}} = p^2/2M_j = \hbar Q_j^2/(2M_j) \). Fermi’s Golden Rule, which connects the wavefunctions before and after the scattering, is sensitive to the phases of the initial and final states of the scatterer. The transfer of energy causes a relative change in the phase of these two states by the factor:

\[
\exp(iE_{\text{kin}} \hbar^{-1} t).
\]

This phase relationship for the total energy transfer leaves a minimal amount of quantum mechanics in the scattering problem (which we expect to go away at very high incident energies):

\[
\frac{d^2 \sigma}{d\Omega \, dE} = \frac{k_i}{k_i} \frac{(2\pi)^3}{2V_0} \sum_j |b_j|^2 \int e^{i\omega t} \left( e^{i/M_j \hat{p}_j} - e^{-i/M_j \hat{p}_j} \right) dt .
\]

(7.99)

In Sect. 7.2.3 we calculated the thermal average, \( \langle \rangle \), when the displacements of the scatterer had a Gaussian thermal spread. To apply this result to (7.99), we use the result: \( \langle e^{iX} \rangle = e^{-X^2/2} \).

\[
\frac{d^2 \sigma}{d\Omega \, dE} = \frac{k_i}{k_i} \frac{(2\pi)^3}{2V_0} \sum_j |b_j|^2 \int e^{i\omega t} \left( e^{i(M_j \hat{p}_j)^2/2} - e^{-((M_j \hat{p}_j)^2/2)} \right) dt .
\]

(7.100)

We complete the square in the exponential, defining:

\[
x^2 \equiv \left( \frac{\sqrt{a}t - b}{2\sqrt{a}} \right)^2 = a t^2 - bt + \frac{b^2}{4a} ,
\]

(7.101)

and we obtain a result:

\[
\int_{-\infty}^{\infty} e^{-at^2 + bt} \, dt = \int_{-\infty}^{\infty} e^{-x^2} \, dx = \sqrt{\frac{\pi}{a}} \, e^{b^2/4a} .
\]

(7.102)
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With the associations for \(a\) and \(b\):

\[
\begin{align*}
    a &= \frac{1}{2} \left( \frac{\langle \vec{Q} \cdot \vec{p}_j \rangle}{M_j} \right)^2, \\
    b &= i \left( \omega + \frac{\hbar Q^2}{2M_j} \right),
\end{align*}
\]

(7.103)

(7.104)

(7.100) becomes:

\[
\frac{d^2 \sigma}{d\Omega dE} = \frac{k_i (2\pi)^{7/2}}{k_i} \sum_j |b_j|^2 \left( \frac{M_j}{\langle \vec{Q} \cdot \vec{p}_j \rangle} \right)^2 \exp \left( -\frac{\left( \frac{\hbar Q^2}{2M_j} + \omega \right)^2}{2 \left( \frac{\langle \vec{Q} \cdot \vec{p}_j \rangle}{M_j} \right)^2} \right). \tag{7.105}
\]

The differential scattering cross section in (7.105) has the shape of a Gaussian function, centered at an energy \(\hbar \omega\):

\[
\hbar \omega = -\frac{\hbar^2 Q^2}{2M_j}. \tag{7.106}
\]

The center of the Gaussian is simply the energy transfer from a scattering with a single-particle recoil. This result could be obtained by classical mechanics. The spread of this Gaussian is obtained from the denominator in the Gaussian of (7.105). This width increases with \(Q\) and with the mean-squared velocity of the scatterers. The ratio of shift to width grows larger with \(Q\), however. In the classical limit of very large \(Q\), the width is negligible, so the energy and momentum transfers are well-defined.

7.3.2 Multiphonon Expansion

We return to develop (7.94) with the “multiphonon expansion,” which is obtained from the expansion of the exponential \(\exp(\vec{U} \vec{V})\). This is seen most easily in the incoherent approximation, where we replace:

\[
\exp(\vec{U} \vec{V}) \rightarrow \exp(\vec{U}) \exp(\vec{V}) = \exp(\vec{U}) \exp(\vec{V}). \tag{7.107}
\]

so that, treating \(\vec{U}\) as a displacement and not an operator:

\[
\frac{d^2 \sigma}{d\Omega dE} = \frac{k_i (2\pi)^3 N |b|^2}{2V_0} \sum_l e^{i\delta_l} \exp(\vec{U}_l^2) \times \int e^{i\omega t} \exp \left( -i\vec{Q} \cdot \vec{u} i\vec{Q} \cdot \vec{u} \right) dt. \tag{7.108}
\]
7.3. PRACTICAL EXPRESSIONS FOR PHONON SCATTERING

The harmonic oscillator energy $M\omega^2u^2_{\text{max}}$, is assumed quantized in units of $\hbar\omega$. From $\hbar\omega \propto M\omega^2u^2_{\text{max}}$, we replace the displacement $u = \sqrt{\hbar/(2M\omega)}$, for which we expect a time-dependence:

\[
\frac{d^2\sigma}{d\Omega dE} = \frac{k_f (2\pi)^3 N|b|^2}{2V_0} \sum_l e^{i\vec{Q}\cdot\vec{x}_l} \exp\left(\frac{1}{2}\right) \\
\times \int e^{i\omega t} \exp\left(\frac{\hbar Q^2}{2M\omega} Y(t)\right) dt , \tag{7.109}
\]

The function $Y(t)$ serves to convert between phonon energy and nuclear displacement, and varies with the phonon DOS, $g(E)$, and temperature through the Planck factor, $n(E)$. It can be shown to be [Lovesey Vol 1. Eq. 4.238]

\[
Y(t) = \int_{-\infty}^{\infty} \frac{g(E)}{E} n(E) e^{-i(E/\hbar)t} dE . \tag{7.110}
\]

Assuming the argument of the exponential in (7.109) is small, we expand this exponential

\[
\frac{d^2\sigma}{d\Omega dE} = \frac{k_f (2\pi)^3 N|b|^2}{2V_0} \sum_l e^{i\vec{Q}\cdot\vec{x}_l} \exp\left(\frac{1}{2}\right) \\
\times \int e^{i\omega t} \left(1 + \frac{\hbar Q^2}{2M\omega} Y(t) + \frac{1}{2} \left(\frac{\hbar Q^2}{2M\omega} Y(t)\right)^2 + \ldots\right) dt . \tag{7.111}
\]

The terms in the expansion of (7.111) are recognized as a series in powers of $E_{\text{Recoil}}/(\hbar\omega)$ — the recoil energy, $p^2/(2M)$, divided by the energy of the phonon, $\hbar\omega$. It is instructive to compare (7.111) to (6.51), which is re-written here. Note that it did not include a Debye–Waller factor:

\[
\Psi_{\text{sc}}(\vec{Q}, r) = -\frac{e^{i\vec{Q}\cdot\vec{r} - \omega t}}{|r|} \\
\times \sum_{l,x} b_\lambda e^{i\vec{Q}\cdot\vec{x}_l} \left(1 + i\vec{Q} \cdot \vec{u}_{\lambda x}(t) - \frac{1}{2} (\vec{Q} \cdot \vec{u}_{\lambda x}(t))^2 + \ldots\right) . \tag{7.112}
\]

Analyzing the two series in (7.111) and (7.112), term-by-term, we find:

- The first term, the 1, provides a Fourier transform of a constant, which is a delta function, $\delta(\omega)$. Since the excitation energy is therefore zero, this is an elastic scattering process. Note that the sum of phase factors $e^{i\vec{Q}\cdot\vec{x}_l}$ over lattice sites and the Debye–Waller factor $e^{i\vec{Q}\cdot\vec{x}_l}$ are as expected for diffracted neutron wavefunctions. Elastic nuclear scattering has no $Q$ dependence, except through the Debye–Waller factor.
• The second term, involving $\frac{\hbar^2 Q^2}{2M} / (\hbar \omega)$, is the inelastic scattering that occurs by exciting one phonon. We found in (6.52) that its Fourier transform led to a delta function $\delta(\omega - \omega(\vec{q}))$ from the conservation of energy. A conservation of momentum led to a similar delta function, $\delta(\vec{q} - \vec{Q} - \vec{q})$. Single phonon scattering increases with $Q$ as $Q^2$.

• The third term, involving $\left(\frac{\hbar^2 Q^2}{2M} \right)^2 / (\hbar \omega)$, is the scattering that involves the excitation of two phonons. This occurs in one scattering event, not through the creation of two phonons by two different deflections of the neutron. (The latter is “multiple scattering.”) When the time function is squared, such as $\cos^2(\omega t) = 1/2 (1 + \cos(2\omega t))$, the frequency is doubled, and energy conservation provides the delta function $\delta(\omega - 2\omega(\vec{q}))$. Two-phonon scattering involves twice the energy transfer as a one-phonon process. Note how it increases rapidly with $Q$, going as $Q^4$.

• Higher order terms, involving $+ \ldots$, involve the excitation of many phonons in one scattering of the neutron. These higher-order terms approach classical behavior. Typically the scattering of a particle with a large a large momentum transfer causes the excitation of many phonons, sometimes better described as the creation of heat.

We note that the analysis here has assumed ignored phase relationships in multiphonon scattering. This is typical of even more sophisticated treatments of the problem. Multiphonon scattering is usually analyzed in the incoherent approximation.

### 7.4 Magnetic Scattering

#### 7.4.1 Magnetic Form Factor and Scattering Amplitude

Magnetic scattering originates with the interaction between the spin of the neutron and the spins of electrons and/or the motions of electrons. Magnetic scattering is inherently more complicated than nuclear scattering because the potentials have vector character. The magnetic forces are also long range.

The scattering potential can be written in the general form $V = -\vec{\mu}_n \cdot B$, where $B$, which originates with the electrons, has a spin component $B_S$ and an orbital component $B_L$:

$$B = B_S + B_L.$$  \hspace{1cm} (7.113)
These components have different mathematical forms:

\[ B_L \propto \frac{\vec{R} \times \vec{p}}{R^2}, \quad (7.114) \]

\[ B_S \propto \vec{\nabla} \times \left( \frac{\vec{s} \times \vec{R}}{R^2} \right). \quad (7.115) \]

From (1.75) we obtain the scattering amplitude in the first Born approximation as the Fourier transform of the scattering potential

\[ V = -\vec{\mu}_n \cdot B \]

\[ f_{\text{mag}}(Q, E) = \sqrt{\frac{k_i}{k_i}} \frac{2m}{4\pi} \int_{-\infty}^{\infty} e^{iQ \vec{r}} (-\vec{\mu}_n \cdot B) \, d\vec{r}. \quad (7.116) \]

\[ f_{\text{mag}}(Q, E) = \sqrt{\frac{k_i}{k_i}} (-\gamma r_e) \int_{-\infty}^{\infty} e^{iQ \vec{r}} \frac{1}{\hbar} \vec{R} \times \vec{p} \, d\vec{r} + \vec{\nabla} \times \left( \frac{\vec{s} \times \vec{R}}{R^2} \right) \right) \, d\vec{r}, \quad (7.117) \]

where \( \vec{s} \) is the neutron spin, later to be the spin operator. Many constants were combined into the “classical electron radius,” \( r_e = e^2 / (m_e c^2) \), and \( \gamma \) is the gyromagnetic ratio of the neutron, \( \gamma = 1.913 \).

The evaluation of (7.117) is most expedient with two mathematical tricks:

\[ \int_{-\infty}^{\infty} \frac{\vec{R}}{R^2} e^{-iQ \vec{R}} \, d\vec{R} = -4\pi i \frac{Q}{Q}, \quad (7.118) \]

\[ \vec{\nabla} \times \left( \frac{\vec{s} \times \vec{R}}{R^2} \right) = \frac{1}{2\pi^2} \int_{-\infty}^{\infty} \vec{q} \times (\vec{s} \times \vec{q}) e^{-iQ \vec{q}} \, d\vec{q}. \quad (7.119) \]

The derivation of (7.118) is not difficult – it involves transformation to spherical coordinates where the \( R^2 \) in the denominator is cancelled by an \( R^2 \) in the differential volume element.\(^5\) Unfortunately, (7.119) is much more work to obtain unless one is rather clever with, or accepting of, vector identities. A sketch of its derivation is given in Appendix B.2 of Squires’ book.

In using the tricks (7.118) and (7.119), the necessary steps are:

- With the neutron at \( \vec{r} \) and the \( i \)th electron at \( \vec{r}_i \), the position for the magnetic field, \( \vec{R} \) in (7.114) and (7.115), is the distance separating \( \vec{r} \) and \( \vec{r}_i \): \( \vec{R} = \vec{r} - \vec{r}_i \).

\(^4\) Note that \( B_L \) has the form of the Biot-Savart law for the electron current (with electron momentum \( \vec{p} \)) about the atom. The \( B_S \) can be written as a curl of a vector potential, \( B_S = \vec{\nabla} \times A \), if \( A = \mu_e \times \vec{R}/R^2 \), and \( \mu_e = -e\hbar / m_e = -2\mu_B \).

\(^5\) Note that (7.118) is the inverse transformation of (A.27) in the Appendix, with the interchange of \( \vec{R} \) and \( \vec{Q} \).
The Fourier transform of $\vec{R} \times \vec{p} / R^2$ first involves replacing the exponential $e^{-i\vec{Q} \cdot \vec{R}} = e^{-i\vec{Q} \cdot \vec{r} + i\vec{Q} \cdot \vec{r}}$, in order to use coordinates of the neutron and the $i^{th}$ electron. The second phase factor $e^{-i\vec{Q} \cdot \vec{r}_i}$, is a constant, and is removed from the integration over all space. The remaining spatial integration becomes, from (7.118), $-i4\pi \vec{Q} \times \vec{p}/Q$.

Notice that the right-hand side of (7.119) has its only dependence on $\vec{R}$ in the exponential (which we again write as $e^{-i\vec{q} \cdot \vec{r}} = e^{-i\vec{q} \cdot \vec{r} + i\vec{q} \cdot \vec{r}}$). When (7.119) is substituted into (7.117), a double integral (over $\vec{r}$ and $\vec{q}$) is obtained. After the phase factor, $e^{i\vec{q} \cdot \vec{r}_i}$, is isolated, the $\vec{r}$-integral is of the form $\int e^{i(Q-q) \cdot \vec{r}} d\vec{r}$. The $\vec{r}$-integral therefore gives a three-dimensional $\delta$-function, $(2\pi)^3 \delta(\vec{Q} - \vec{q})$.

The result from these manipulations with (7.118) and (7.119) is:

$$f_{mag}(\vec{Q}, E) = \sqrt{\frac{k_f}{k_i}} (\gamma r_0) \times \vec{\sigma} \cdot \left( \int_{-\infty}^{\infty} e^{i\vec{q} \cdot \vec{r}} \frac{1}{2\pi} i \vec{Q} \times (\vec{s} \times \vec{q}) (2\pi)^3 \delta(\vec{Q} - \vec{q}) dq - \frac{i4\pi}{\hbar} e^{i\vec{q} \cdot \vec{r}_i} \frac{\vec{Q} \times \vec{p}}{Q} \right).$$  (7.120)

Integrating over the $\delta$-function forces $\vec{q} \rightarrow \vec{Q}$:

$$f_{mag}(\vec{Q}, E) = \sqrt{\frac{k_f}{k_i}} (\gamma r_0) e^{i\vec{Q} \cdot \vec{r}_i} \times \vec{\sigma} \cdot \left( \vec{Q} \times (\vec{s} \times \vec{Q}) - \frac{i}{\hbar} \frac{\vec{Q} \times \vec{p}}{Q} \right).$$  (7.121)

With the definition of $\vec{M}_\perp(\vec{Q})$, which involves a sum over all electrons in the sample:

$$\vec{M}_\perp(\vec{Q}) = \sum_i e^{i\vec{q} \cdot \vec{r}_i} \left( \vec{Q} \times (\vec{s} \times \vec{Q}) - \frac{i}{\hbar} \frac{\vec{Q} \times \vec{p}}{Q} \right),$$  (7.122)

$$f_{mag}(\vec{Q}, E) = 4\pi \sqrt{\frac{k_f}{k_i}} (\gamma r_0) \vec{\sigma} \cdot \vec{M}_\perp(\vec{Q}).$$  (7.123)

Equation (7.123) is a general expression for the magnetic scattering from the spin and orbital moment of the electrons. It includes a sum over the phase
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factors for electrons at all \( \{ \mathbf{r}_i \} \). Unfortunately, the spin and orbital terms in the large parentheses in (7.122) have different forms, and to see more clearly the vectorial character of magnetic scattering it is convenient to make them equivalent using the expression (proved with some effort in Squires Appendix H.1)

\[
\tilde{M}_{LL} = \frac{i}{\hbar Q} \sum_{\mathbf{r}_i} e^{i\mathbf{Q} \cdot \mathbf{r}_i} \mathbf{r}_i \mathbf{p} = \frac{1}{2\mu_B} \mathbf{Q} \times \left( \tilde{M}_L(\mathbf{Q}) \times \mathbf{Q} \right),
\]

(7.124)

where the Fourier transform of the magnetic form factor from the spatial distribution of electron currents is:

\[
\tilde{M}_L(\mathbf{Q}) \equiv \int \tilde{M}_L(\mathbf{r}) e^{i\mathbf{Q} \cdot \mathbf{r}} d\mathbf{r}.
\]

(7.125)

It is natural to write (7.122) as:

\[
\tilde{M}_\perp = \tilde{M}_L + \tilde{M}_{LL},
\]

(7.126)

\[
\tilde{M}_\perp = \frac{1}{2\mu_B} \mathbf{Q} \times \left( \tilde{M}(\mathbf{Q}) \times \mathbf{Q} \right),
\]

(7.127)

where \( \tilde{M}(\mathbf{Q}) \) is the Fourier transform of the spatial distribution of all magnetization (as in (7.125)):

\[
\tilde{M}(\mathbf{Q}) \equiv \int \tilde{M}(\mathbf{r}) e^{i\mathbf{Q} \cdot \mathbf{r}} d\mathbf{r},
\]

(7.128)

We arrive at the cleaner expression for the magnetic scattering factor

\[
f_{\text{mag}}(\mathbf{Q}, E) = 4\pi \sqrt{\frac{k_i}{k_f}} (-\gamma r_0) \sum_{\mathbf{r}_i} e^{i\mathbf{Q} \cdot \mathbf{r}_i} \mathbf{p} \cdot \left( \mathbf{Q} \times \left( \tilde{M}(\mathbf{Q}) \times \mathbf{Q} \right) \right),
\]

(7.129)

7.4.2 Vector Orientations in Magnetic Scattering

Equation (7.129) shows that the magnetic scattering is proportional to the vector \( \mathbf{Q} \times \tilde{M} \times \mathbf{Q} \). The maximum magnetic scattering therefore occurs when the direction of the spin, \( \mathbf{S} \), or magnetization, \( \tilde{M}(\mathbf{r}) \), is perpendicular to the scattering vector, \( \mathbf{Q} \). This is illustrated in Fig. 7.4.2, which shows intensity contours measured about the forward beam in a small-angle scattering experiment. A magnetic field of 8 kG was applied to the specimen, perpendicular to the direction of the incident beam. This field should be sufficient to saturate the magnetic moment of the sample (a soft magnetic material), aligning all its spins. Notice that the contours are oriented perpendicularly to the direction of the applied magnetic field. The scattering along the direction of the magnetic field is non-zero, however, because Ni-Fe has strong nuclear scattering. By
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Figure 7.2: Experimental intensity contours from small-angle neutron scattering (SANS) from fcc Ni-Fe in the presence of an 8 kG applied magnetic field. The forward beam was perpendicular to the plane of the paper. The intensity decreases with angle from the forward beam, but more rapidly in the direction of the applied magnetic field.

Figure 7.3: Important vectors for magnetic scattering. (left) The gray plane, parallel to the paper, is defined by the vectors \( \vec{M} \) and \( \vec{Q} \). The vector product \( \vec{M} \times \vec{Q} \) is perpendicular to the plane of the paper and \( \vec{M}_\perp \equiv \vec{Q} \times \vec{M} \times \vec{Q} \) again lies in the plane. (right) The vector of length \( \vec{M} \cdot \vec{Q} \) along the direction \( \vec{Q} \).

Comparing intensities parallel and perpendicular to the applied magnetic field, it is possible to extract individual profiles for magnetic and nuclear scattering.

The relationship between the generalized magnetization, \( \vec{M} \), its projection, \( \vec{M}_\perp \), and the scattering vector, \( \vec{Q} \), is illustrated with the help of Fig. 7.4.2 and its caption. By comparing the two parts of this figure, we find

\[
\vec{Q} \times \vec{M} \times \vec{Q} = \vec{M} - \vec{Q} (\vec{M} \cdot \vec{Q}) .
\]  

(7.130)

The cross-section for magnetic scattering is proportional to \( |f_{mag}|^2 \). We need to take the product of \( \vec{M}_\perp \) with its Hermitian adjoint to obtain the intensity

\[
\vec{M}_\perp \vec{M}^\dagger = \left( \vec{M} - \vec{Q} (\vec{M} \cdot \vec{Q}) \right) \left( \vec{M} - \vec{Q} (\vec{M} \cdot \vec{Q}) \right) .
\]  

(7.131)
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When distributing the product in (7.131), the two middle terms have the same form but opposite sign to the fourth term, so

\[ \tilde{M}_+^\dagger \tilde{M}_- = \tilde{M}_+ \tilde{M}_- - (\tilde{M}_+ \cdot \tilde{Q})(\tilde{M}_- \cdot \tilde{Q}) \cdot (7.132) \]

We resolve \( \tilde{M} \) and \( \tilde{Q} \) into Cartesian components

\[ \tilde{M}_+^\dagger \tilde{M}_- = \sum_{\alpha, \beta} S_{\alpha}^\dagger S_{\beta} - S_{\alpha}^\dagger Q_{\alpha} Q_{\beta} \cdot (7.133) \]

\[ \tilde{M}_+^\dagger \tilde{M}_- = \sum_{\alpha, \beta} (\delta_{\alpha\beta} - Q_{\alpha} Q_{\beta}) S_{\alpha}^\dagger S_{\beta} \cdot (7.134) \]

7.4.3 Averaging over Neutron Polarizations

To obtain an experimental cross-section from (7.129), we need to average over the spin orientations of the incident neutrons (orientations of \( \tilde{\sigma} \)). We expect to write the cross-section as

\[ \left| f_{\text{mag}} \right|^2 \frac{d^2 \sigma}{d\Omega dE} = (\gamma r e)^2 k_f \left( \lambda_f, \sigma_f \right| \tilde{\sigma} \cdot \tilde{M}_+ | \lambda_i, \sigma_i \left) \right|^2 \delta(E_f - E_i + \hbar \omega) \cdot (7.135) \]

Compared to nuclear scattering, (7.135) includes additional coordinates in the matrix element, \( \sigma_i \) and \( \sigma_f \), to account for the change in spin of the neutron after scattering. Again, the \( \lambda_i \) and \( \lambda_f \) refer to states of the scatterer. For magnetic scattering, a change in \( \lambda \) may originate with the creation or annihilation of an excitation of the electron spins. Further progress with these coordinates will require a magnetic dynamics model. The total cross-section requires that we sum over all initial and final states of the neutron:

\[ \left| f_{\text{mag}} \right|^2 \frac{d^2 \sigma}{d\Omega dE} = (\gamma r e)^2 k_f \sum_{i} \sum_{f} \left( \lambda_f, \sigma_f \right| \tilde{\sigma} \cdot \tilde{M}_+ | \lambda_i, \sigma_i \left) \right|^2 \delta(E_f - E_i + \hbar \omega) \cdot (7.136) \]

The coordinates \( \sigma_i \) and \( \sigma_f \) describe to the neutron spin, which we expect to be up or down (sometimes \( \uparrow \), \( \downarrow \)). Equation (7.135) is the cross-section for one scattering process, but in an experiment we expect to average over the spins of many neutrons. The total cross-section should include weighting functions such as \( p_\alpha \) to account for different probabilities of initial spin up and spin down neutrons, as for example with polarized beam experiments. The final spin states are assumed unbiased, however, and so have no associated weight function.

The operator in (7.136) can be resolved into its vector components:

\[ \tilde{\sigma} \cdot \tilde{M}_+ = \sigma_x \tilde{M}_+ x + \sigma_y \tilde{M}_+ y + \sigma_z \tilde{M}_+ z \cdot (7.137) \]
The coordinates of $\sigma_a$ pertain to the neutrons only, and the coordinates of $\tilde{M}_{\lambda z}$ pertain only to the electrons. The products in (7.137) serve to group the factors involving electrons and neutrons, but they separate as for example:

$$\langle \lambda_f, \sigma_f | \sigma_z \tilde{M}_{\lambda z} | \lambda_i, \sigma_i \rangle = \langle \lambda_f | \tilde{M}_{\lambda z} | \lambda_i \rangle \langle \sigma_f | \sigma_z | \sigma_i \rangle .$$  \hspace{1cm} (7.138)

The cross-section of (7.136) can be separated into nine different terms by use of (7.137) and (7.138) (note the sequencing of subscripts $x$ and $y$):

$$\frac{d^2\sigma}{d\Omega \, dE} = (\gamma r_e)^2 \frac{k_f}{k_i} \sum_i \sum_f \left( \langle \lambda_i | \tilde{M}_{\lambda x} | \lambda_f \rangle \langle \sigma_i | \sigma_x | \sigma_f \rangle \langle \sigma_f | \sigma_z | \sigma_i \rangle \langle \lambda_f | \tilde{M}_{\lambda z} | \lambda_i \rangle \\
+ \langle \lambda_i | \tilde{M}_{\lambda y} | \lambda_f \rangle \langle \sigma_i | \sigma_y | \sigma_f \rangle \langle \sigma_f | \sigma_z | \sigma_i \rangle \langle \lambda_f | \tilde{M}_{\lambda z} | \lambda_i \rangle + \ldots \right) \delta(E_f - E_i + \hbar \omega) .$$  \hspace{1cm} (7.139)

By closure, $\sum_f |\sigma_f\rangle \langle \sigma_f| = 1$, there is simplification of the spin factors

$$\frac{d^2\sigma}{d\Omega \, dE} = (\gamma r_e)^2 \frac{k_f}{k_i} \sum_i \sum_f \left( \langle \lambda_i | \tilde{M}_{\lambda x} | \lambda_f \rangle \langle \sigma_i | \sigma_x | \sigma_f \rangle \langle \sigma_f | \sigma_z | \sigma_i \rangle \langle \lambda_f | \tilde{M}_{\lambda z} | \lambda_i \rangle \\
+ \langle \lambda_i | \tilde{M}_{\lambda y} | \lambda_f \rangle \langle \sigma_i | \sigma_y | \sigma_f \rangle \langle \sigma_f | \sigma_z | \sigma_i \rangle \langle \lambda_f | \tilde{M}_{\lambda z} | \lambda_i \rangle + \ldots \right) \delta(E_f - E_i + \hbar \omega) .$$  \hspace{1cm} (7.140)

The spin operators have the properties\footnote{These relations are obtained, for example, from explicit forms of the Pauli spin matrices.}

$$\sigma_x | \uparrow \rangle = | \downarrow \rangle , \quad \sigma_y | \uparrow \rangle = +i | \downarrow \rangle , \quad \sigma_z | \uparrow \rangle = + | \uparrow \rangle ,$$

$$\sigma_x | \downarrow \rangle = | \uparrow \rangle , \quad \sigma_y | \downarrow \rangle = -i | \downarrow \rangle , \quad \sigma_z | \downarrow \rangle = - | \downarrow \rangle .$$  \hspace{1cm} (7.141)

Equation (7.140) includes terms with the factors $\langle \uparrow | \sigma_x \sigma_y | \uparrow \rangle$ and $\langle \uparrow | \sigma_y \sigma_z | \uparrow \rangle$. Evaluating them with (7.141) gives:

$$\langle \uparrow | \sigma_x \sigma_y | \uparrow \rangle = \langle \uparrow | \sigma_x (+i) | \downarrow \rangle = \langle \uparrow | + i \uparrow \rangle = +i (\uparrow | \uparrow \rangle = +i ,$$

$$\langle \uparrow | \sigma_y \sigma_z | \uparrow \rangle = \langle \uparrow | \sigma_y | \downarrow \rangle = \langle \uparrow | - i \uparrow \rangle = -i (\uparrow | \uparrow \rangle = -i ,$$  \hspace{1cm} (7.142)

(7.143)
where the last result used the normalization $\langle \uparrow | \uparrow \rangle = 1$. It is not surprising that (7.142) and (7.143) give opposite results because the spin operators $\sigma_x$ and $\sigma_y$ do not commute. The consequence is that there is a pairwise cancellation of the six terms in (7.140) that involve the subscripts $xy$, $yx$, $xz$, $zx$, $yz$, $zy$.

For unpolarized neutrons with $|\uparrow\rangle$ and $|\downarrow\rangle$ of equal probabilities, we expect no bias for the three remaining terms of (7.140), which becomes

$$\frac{d^2\sigma}{d\Omega dE} = (yr_e)^2 \frac{k_f}{k_i} \sum_i \sum_f$$
$$\left( \langle \lambda_i|\vec{M}_{\perp\alpha}|\lambda_f \rangle \langle \sigma_{\alpha}\sigma_i|\lambda_f|\vec{M}_{\perp\beta}|\lambda_i \rangle \right.$$
$$+\langle \lambda_i|\vec{M}_{\perp\alpha}|\lambda_f \rangle \langle \sigma_{\alpha}\sigma_i|\lambda_f|\vec{M}_{\perp\beta}|\lambda_i \rangle$$
$$+\langle \lambda_i|\vec{M}_{\perp\alpha}|\lambda_f \rangle \langle \sigma_{\alpha}\sigma_i|\lambda_f|\vec{M}_{\perp\beta}|\lambda_i \rangle \right) \delta(E_f-E_i+\hbar\omega) . \quad (7.144)$$

The three remaining terms have factors $\langle \sigma_i|\sigma_j^2|\sigma_i \rangle$, $\langle \sigma_i|\sigma_j^2|\sigma_i \rangle$, and $\langle \sigma_i|\sigma_j^2|\sigma_i \rangle$. These terms evaluate to 1 as for example

$$\langle \uparrow | \sigma_i^2 | \uparrow \rangle = \langle \uparrow | \sigma_i | \downarrow \rangle = \langle \uparrow | \uparrow \rangle = 1 . \quad (7.145)$$

simplifying (7.144)

$$\frac{d^2\sigma}{d\Omega dE} = (yr_e)^2 \frac{k_f}{k_i} \sum_i \sum_f$$
$$\left( \langle \lambda_i|\vec{M}_{\perp\alpha}|\lambda_f \rangle \langle \lambda_f|\vec{M}_{\perp\beta}|\lambda_i \rangle \right.$$}
$$+\langle \lambda_i|\vec{M}_{\perp\alpha}|\lambda_f \rangle \langle \lambda_f|\vec{M}_{\perp\beta}|\lambda_i \rangle$$
$$+\langle \lambda_i|\vec{M}_{\perp\alpha}|\lambda_f \rangle \langle \lambda_f|\vec{M}_{\perp\beta}|\lambda_i \rangle \right) \delta(E_f-E_i+\hbar\omega) . \quad (7.146)$$

In (7.146) we recognize that that the three terms in parentheses are the projections onto a set of final states of the three components of the vector $\vec{M}_{\perp}$. This can be obtained as $\langle \lambda_f|\vec{M}_{\perp}|\lambda_f \rangle \langle \lambda_f|\vec{M}_{\perp}|\lambda_i \rangle$. We therefore make use of (7.134) to rewrite (7.146) as

$$\frac{d^2\sigma}{d\Omega dE} = (yr_e)^2 \frac{k_f}{k_i} \sum_{\alpha\beta} \delta_{\alpha\beta} - \vec{M}_{\perp} \vec{M}_{\perp}$$
$$\sum_i \sum_f \langle \lambda_i|\vec{M}_{\perp\alpha}|\lambda_f \rangle \langle \lambda_f|\vec{M}_{\perp\beta}|\lambda_i \rangle \delta(E_f-E_i+\hbar\omega) . \quad (7.147)$$
Further Reading

The contents of the following are described in the Bibliography.


Problems

1.) (a) Develop the operator identity (7.70), working the algebra given before it in Sect. 7.2.2.

(b) Show that the Weyl identity (7.59) is obtained when (7.71) is achieved.
Chapter 8

Phonon Dynamics

8.1 Phonon Thermodynamics

8.1.1 Phonon Statistical Mechanics

Phonons are useful for understanding the collective motions of atoms in crystals because as normal modes of a solid, phonon modes can be treated independently. Independent contributions to thermodynamic functions are then possible, simplifying analyses of the vibrational partition function and vibrational entropy, for example.

Phonons are bosons, so there is no limit to how many of them can be present in each mode, or oscillator. Consider a set of oscillators \( N \), all of the same energy \( \epsilon \), and assume these oscillators can exchange energy. Suppose there are a number \( m \) of phonons that are free to distribute between these different oscillators. We calculate the phonon entropy from the equation carved in stone in Fig. 8.1

\[
S = k_B \ln \Omega ,
\]  

(8.1)

where there are a number of ways \( \Omega \), to arrange the \( m \) phonons in the \( N \) oscillators.

Figure 8.2 illustrates five possibilities for distributing 4 phonons among 4 oscillators. The top configuration in Fig. 8.2 has all four phonons in the left oscillator. The bottom configuration places one phonon in each oscillator. The middle three are other possibilities, such as three phonons in the first oscillator, one in the second, and none in the other two.

What is important about Fig. 8.2 is that it inspires the trick for calculating \( \Omega \) combinatorially. The trick is to consider the three vertical bars and the four circles as distributed at random over the seven positions in each row. The number \( \Omega \) becomes the number of ways to distribute \( m \) indistinguishable phonons over \((N - 1) + m\) possible sites (equal to the probability of distributing \( N - 1 \) indistinguishable bars over \((N - 1) + m\) possible sites). The result is a
Figure 8.1: The Boltzmann monument in Vienna. The constant $k$ is related to $k_B$ by the factor 2.3026 if log denotes $\log_{10}$. Our notation uses $\Omega$ instead of $W$ to avoid conflict with Debye–Waller factors.

Figure 8.2: Five configurations of placing 4 ($m$) phonons into 4 ($N$) oscillators. Vertical lines demark oscillators; circles denote phonons.
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For systems of many oscillators we can replace \( N - 1 \) with \( N \), of course:

\[
\Omega = \frac{(N + m)!}{N! m!}.
\]  

(8.2)

Substituting (8.2) into (8.1), and using the Stirling approximation of \( \ln(x!) \approx x \ln x - x \)

\[
S_{\text{osc}} = k_B \left[ (N + m) \ln(N + m) - N \ln N - m \ln m \right].
\]  

(8.3)

We now consider the entropy per oscillator, \( S_{\text{osc}}/N \), and the occupancy of each oscillator, \( n \equiv m/N \).

\[
S_{\text{osc}} \frac{N}{N} = k_B \left[ (1 + n) \ln(1 + n) - \ln n \ln n \right],
\]  

(8.4)

\[
S_{\text{osc}} \frac{N}{N} = k_B \left[ (1 + n) \ln(1 + n) + (1 + n) \ln N - N \ln n - n \ln N \right],
\]  

(8.5)

\[
S_{\text{osc}} \frac{N}{N} = k_B \left[ (1 + n) \ln(1 + n) - (1 + n) \ln n - n \ln n \right],
\]  

(8.6)

\[
S_{\text{osc}} \frac{N}{N} = +k_B \left[ (1 + n) \ln(1 + n) - n \ln n \right].
\]  

(8.7)

It is interesting to compare (8.7) to the analogous equation for fermions, where \( m \) electrons are distributed over \( N \) states of energy \( \epsilon \), for example. For fermions, the combinatorics give \( \Omega = N! / [(N - m)! m!] \). Fermion entropy, incidentally, is the same as the entropy of mixing of atoms on a crystal lattice, were each lattice site can hold only one atom:

\[
S_{\text{mix}} \frac{N}{N} = -k_B \left[ (1 - c) \ln(1 - c) + c \ln c \right].
\]  

(8.8)

Equation (8.7) and (8.8) have two differences – all signs are reversed, and \( n \) can exceed 1, whereas \( 0 \leq c \leq 1 \).

8.1.2 Phonon Free Energy

With (8.7) for the entropy per oscillator, we construct a Helmholtz free energy for a system of oscillators, where the phonon energy is simply the number of phonons per oscillator times the phonon energy, \( n \epsilon \)

\[
F_{\text{osc}} = E_{\text{osc}} - TS_{\text{osc}},
\]  

(8.9)

\[
F_{\text{osc}} = n \epsilon - k_B \left[ (1 + n) \ln(1 + n) - n \ln n \right].
\]  

(8.10)

---

1The first phonon can be placed in \( N-1 \) possible sites, the second has one less, i.e. \( N-2 \) possibilities, and the \( m \)th phonon can be placed in \( N \) ways. The product of these independent probabilities is \( (N-1)!/(N-1)! \). There is, however, an overcounting because we put the phonons down individually, labeling the first, second, etc., but the phonons are indistinguishable. The overcounting is the number of ways of distributing \( m \) phonons over \( m \) sites, which is \( m! \), so we divide our result by \( m! \) in (8.2).
We seek the equilibrium number of phonons per oscillator at temperature $T$ by calculating the minimum of $F_{\text{osc}}$ with respect to $n$:

$$\frac{\partial F_{\text{osc}}}{\partial n} = 0,$$  
(8.11)

$$0 = \varepsilon - T k_B \left[ \ln(1 + n) + 1 - \ln n - 1 \right],$$  
(8.12)

$$-\frac{\varepsilon}{k_B T} = \ln \frac{n}{1 + n},$$  
(8.13)

$$e^{-\beta \varepsilon} = \frac{n}{1 + n},$$  
(8.14)

$$e^{-\beta \varepsilon} = n(1 - e^{-\beta \varepsilon}),$$  
(8.15)

$$n(T) = \frac{1}{e^{\beta \varepsilon} - 1}.$$  
(8.16)

where $\beta \equiv (k_B T)^{-1}$. Equation (8.16) is the Planck distribution for phonons, or the Bose-Einstein distribution (for zero chemical potential). It is a consequence of the statistics of configurations illustrated in Fig. 8.2.

### 8.1.3 Phonon Entropy and Some Chemical Trends

Here we extend the thermodynamics to a collection of phonon modes with different energies, and we illustrate a handy calculational approach with the partition function. A quantum harmonic oscillator has energies that increase by $\varepsilon$ for each phonon, and the partition function for a single harmonic oscillator is

$$Z_i = \sum_{n} e^{-\beta(n + 1/2)\varepsilon_i},$$  
(8.17)

$$Z_i = \frac{e^{-\beta \varepsilon_i / 2}}{1 - e^{-\beta \varepsilon_i}},$$  
(8.18)

where (8.18) was obtained by identifying (8.17) as a geometric series times the constant factor $\exp(-\beta \varepsilon_i / 2)$. The partition function for a harmonic solid with $N$ atoms and $3N$ independent oscillators is the product of these single oscillator partition functions,

$$Z_N = \prod_{i=1}^{3N} \frac{e^{-\beta \varepsilon_i / 2}}{1 - e^{-\beta \varepsilon_i}},$$  
(8.19)

from which we can calculate the phonon free energy as $F = -k_B T \ln Z$,

$$F_{\text{ph}} = \frac{1}{2} \sum_{i} \varepsilon_i + k_B T \sum_{i} \ln \left(1 - e^{-\beta \varepsilon_i}\right),$$  
(8.20)
and the phonon entropy by differentiating \( F \) with respect to \( T \):

\[
S_{\text{ph}} = k_B \sum_{i}^{3N} \left[-\ln \left( 1 - e^{-\beta \epsilon_i} \right) + \frac{\beta \epsilon_i}{e^{\beta \epsilon_i} - 1} \right].
\] (8.21)

It is often useful to work with a phonon density of states (DOS), \( g(\epsilon) \), where \( 3Ng(\epsilon) \, d\epsilon \) phonon modes are in an energy interval \( d\epsilon \). For a DOS acquired as digital data in \( m \) intervals of width \( \Delta \epsilon \) (so \( \epsilon_j = j\Delta \epsilon \)), the partition function can be computed numerically as:

\[
Z_N = \prod_{j=1}^{m} \left( \frac{e^{-\beta \epsilon_j/2}}{1 - e^{-\beta \epsilon_j}} \right)^{3N(g(\epsilon_j))\Delta \epsilon}.
\] (8.22)

Given the phonon DOS, \( g(\epsilon) \), using (8.7) it is possible to obtain this useful expression for the phonon entropy of a harmonic material at any temperature

\[
S_{\text{ph}} = 3k_B \int_{0}^{\infty} g(\epsilon) \left[ (n(\epsilon) + 1) \ln(n(\epsilon) + 1) - n(\epsilon) \ln(n(\epsilon)) \right] \, d\epsilon,
\] (8.23)

where \( g(\epsilon) \) is normalized to 1 and \( n(\epsilon) \) is the Planck distribution for phonon occupancy at the temperature of interest. A handy expression for the high-temperature limit of the difference in phonon entropy between two harmonic phases, \( \alpha \) and \( \beta \) can be obtained readily from (8.21):

\[
S_{\beta - \alpha}^{\text{ph}} = 3k_B \int_{0}^{\infty} (g^\alpha(\epsilon) - g^\beta(\epsilon)) \ln(n(\epsilon)) \, d\epsilon.
\] (8.24)

The DOS is central to understanding the phonon contributions to thermodynamics, but \( g(\epsilon) \) must be known to high accuracy because small differences in DOS curves are often important. For example, if we apply (8.24) to a case where the phonon DOS curves of the \( \alpha \) and \( \beta \) phases have the same shape, but differ in energy scaling by 10%, we obtain a change in phonon entropy of \( \Delta S_{\text{ph}} = 3k_B \ln(1.1) \approx 0.3k_B/\text{atom} \). This change in phonon entropy is almost half of the maximum possible change in configurational entropy of the order-disorder transformation of a binary alloy, which is \( \ln 2 \times 0.69 \times 0.69 \approx 0.69 \times 0.69 \times 0.3k_B/\text{atom} \). When comparing the phonon entropies of different alloy phases, a 1% accuracy in the difference in logarithmic-averaged phonon energy is often required. Although the role of phonon entropy on phase transformations has been discussed for many years [1,2], only in recent times have measurements and calculations become adequate for assessing its thermodynamic importance to phase stability [3-5]. Today it is clear that phonon entropy usually differs enough between alloy phases and compositions to have thermodynamic importance.

At low temperatures it is possible to use electronic structure calculations with the local density approximation to calculate phonon frequencies with reasonable accuracy, but a 1% accuracy remains a challenge. Inelastic neutron
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scattering faces its challenges too. A triple axis spectrometer at a reactor neutron source is highly accurate in its energy and momentum measurements, and when a full set of phonon dispersions are measured on single crystals of ordered compounds, the phonon DOS derived from a lattice dynamics model is highly reliable. This technique is not so reliable for disordered solid solutions, unfortunately, because the dispersions are inherently broadened, and this broadening need not be symmetrical in energy. Direct measurements of the full phonon DOS of disordered alloys are possible with chopper spectrometers at pulsed neutron sources, but other difficulties arise. In general, the neutron scattering from phonon excitations does not receive an equal contribution from the different elements in an alloy, and the displacement amplitudes of the different elements may be different in different phonons. This causes a “neutron-weight” that is difficult to correct, as discussed in Sect. 6.3.3, although differential measurements are often possible.

Our knowledge about the phonon entropy of alloy phases is being organized today, but some trends are clear, and some correlations between phonon entropy and alloy properties are known [5,6]. The Hume–Rothery rules of alloy thermodynamics are based on the atomic properties of: 1) electronegativity, 2) metallic radius, and 3) electron-to-atom ratio, with the first factors being the most important. For phonon entropy, we should perhaps add 4) atom mass, to this list. It is found that across the periodic table, where atomic weight varies from 1 to 238 and beyond, for matrix having atoms of mass $M_0$, and solutes having mass $M_S$, the phonon entropy upon alloying tends to scale with $\ln\left(\frac{M_0}{M_S}\right)$ [7]. This is a general consequence of the relationship for an oscillator of $\omega = \sqrt{\frac{k}{M}}$, where the frequency, $\omega$, scales as the inverse root of the mass, $M$, assuming that the spring constant, $k$, is in fact a constant. There is a large scatter in this correlation, however, and it is not a reliable one for most alloys. The spring constants, or more specifically the interatomic force constants $\Phi_{\alpha\alpha'}$ (Eq. 8.47), are in fact not constant in alloys. In particular, a larger solute atom will cause a local compression. Since interatomic forces generally become stiffer under compression, alloying with an atom having both a large size and a large mass will produce an uncertain result for the vibrational entropy.

Van de Walle and Ceder [8] proposed a model based upon a bond-stiffness versus bond-length argument. In their model, an atom pair in different local atomic configurations will have different bond stiffnesses, with greater stiffness for shorter interatomic distances. These characteristics seem transferable when atom pairs are in different crystal structures. The model is useful for semi-quantitative arguments, and when calibrated for specific elements by ab-initio calculations, for example, this model can be used for comparing phonon entropies of different alloy phases.

In alloying there are chemical trends of phonon entropy. In a systematic study of transition metal solutes in vanadium, it was found that the phonon entropy had a robust correlation with the difference in electronegativity between the solute atom and the vanadium atom. Results, shown in Fig. 8.3, indicate that this correlation works well for solutes from the 3$d$, 4$d$, and 5$d$
rows of the periodic table, in spite of their large differences in mass and atomic size. Electronic structure calculations on transition metals have shown that the large differences in electronegativity cause large charge transfers between the solute and its first-nearest-neighbor atoms. These alter the charge screening sufficiently to vary the interatomic forces, as implied by a substantial decrease in electron states at the Fermi level [5, 6, 9, 10]. A simple explanation for transition metals is that with large charge transfers, the bonding tends to become more ionic, leading to fewer metallic-like states near the Fermi level that can become excited in response to atom displacements. The crystal becomes more difficult to distort, and the interatomic forces become stiffer. For alloying of free electron metals, the solute may donate electrons to the band, stiffening it. There is also an effect from hybridizing the occupied local states at the host atoms and the unoccupied local states at the solute, which can cause effects at least as large [11].

8.2 Heat Capacity

8.2.1 Harmonic Heat Capacity

First consider a harmonic model where $C_V$ is obtained with a fixed set of oscillator frequencies, or a fixed phonon DOS. The temperature derivative of the Planck occupancy distribution (Eq. 8.16) with respect to $T$, weighted by the energy of the mode, $\varepsilon_i$, shows that each mode of energy $\varepsilon_i$ contributes to the heat capacity at $T$ the amount

$$C_{V,i}(T) = k_B \left( \frac{\varepsilon_i}{k_B T} \right)^2 \frac{\exp(\varepsilon_i/k_B T)}{(\exp(\varepsilon_i/k_B T) - 1)^2}.$$

(8.25)
In the high temperature (classical) limit where \( k_B T \gg \varepsilon_i \), the heat capacity becomes \( k_B \) per mode. \(^2\) Equation 8.25 is consistent with Eq. 8.21 if we integrate over \( T \) for all modes

\[
S_{V,\text{vib}}(T) = \int_0^T \sum_i^{3N} \frac{C_V(T')}{T'} \, dT'.
\] (8.26)

(Usually we integrate against the phonon DOS, \( g(\varepsilon) \), to obtain \( S_{V,\text{vib}}(T) \).)

A Debye model predicts some general features of the heat capacity. Figure 8.4a shows heat capacity curves for two materials with similar Debye temperatures. The curves look similar at low \( T \), and are essentially identical at very high \( T \). Their most pronounced difference is found at approximately one-quarter of the Debye temperature. The integrand of Eq. 8.26 has a maximum below about a fifth of the Debye temperature, so experimental studies of vibrational entropy by heat capacity should include this range of temperatures. The difference between these two curves, shown in Fig. 8.4b, is typical of the shape of a differential heat capacity curve obtained from real phonon DOSs, although there are differences in detail. The Debye model assumes a phonon DOS that is a quadratic function of \( \varepsilon \), and increases monotonically up to a cutoff energy \( \varepsilon_D \).

The actual DOS often has a more abrupt rise at low energies that comes from the lower sound velocity of the low transverse modes, which are also weighted at their cutoff energy by the effects of dispersion (e.g., from a high DOS at the Brillouin zone boundary). Plots of \( \varepsilon_D \) versus \( T \) typically show a dip and a rise from a minimum at roughly a tenth of the Debye temperature \( \theta_D = \varepsilon_D / k_B \), but this varies from material to material.

### 8.2.2 Quasiharmonic Thermodynamics

More typically measured by calorimetry is the amount of heat going into the solid at constant pressure, \( C_p \), defined as

\[
C_p(T) = T \frac{dS}{dT}_p.
\] (8.27)

The difference between \( C_p \) and \( C_V \) is a classical thermodynamic relationship

\[
C_p - C_V = 9Bv\alpha^2T,
\] (8.28)

where \( B \) is bulk modulus, \( \alpha \) is the linear coefficient of thermal expansion, \( v \) is specific volume, and 9 is the square of three dimensions. Equation 8.28 can

\(^2\)Even with phonon softening this could remain true. In the high-temperature limit, with phonon occupancies of \( k_B T / \varepsilon_i \) per mode, the softening of the modes gives an increase in occupancy proportional to \( 1/\varepsilon_i \), but a decrease in energy proportional to \( \varepsilon_i \). The product of these two factors produces a heat capacity \( C_V \) at high temperatures that is unaffected by mode softening, so \( C_V \) can remain as \( 3k_B / \text{atom} \) at high temperatures. This is not so if there are other phenomena at work, such as a buildup of elastic energy as a crystal expands.
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Figure 8.4: (a) Heat capacity versus temperature for two Debye solids with Debye temperatures of 450 and 500 K. (b) Difference of the two curves in part a.

be derived from formal thermodynamic manipulations, aided for example by techniques of Jacobian transformations.

A physical derivation of Eq. 8.28 can be performed by considering the free energy of a crystal having only 1) phonons, and 2) thermal expansion. There are energy terms from the phonons and from the elastic expansion, and an entropy from the phonons (an average thermal expansion does not generate entropy itself). The free energy is

\[ F(T) = E_{\text{elas}}(T) + E_{\text{ph}}(T) - TS_{\text{ph}}. \]  

The elastic energy of thermal expansion, using \( \delta V/V_0 = 3\alpha T \) is

\[ E_{\text{elas}} = \frac{1}{2} B \frac{(\delta V)^2}{V_0}, \]
\[ E_{\text{elas}} = \frac{9}{2} B V_0 \alpha^2 T^2, \]  

(8.30)

The phonon energy in the classical limit is

\[ E_{\text{ph}}(T) = 3Nk_B T. \]  

(8.31)

In the high-temperature limit, the phonon entropy is (cf., Eq. 8.21 for large \( \beta \))

\[ S_{\text{ph}}(T) = k_B \sum_j 3N \ln \left( \frac{k_B T}{\hbar \omega_j} \right). \]  

(8.32)
For small $\delta V/V^0$ (where the superscript 0 denotes a reference value at zero temperature and pressure), we expect a typical phonon frequency $\omega_j$ to shift as

$$\omega_j = \omega_j^0 (1 - \gamma \delta V/V^0),$$  

(8.33)

$$\ln(\omega_j) \approx \ln(\omega_j^0) - \gamma \frac{\delta V}{V^0},$$  

(8.34)

where $\gamma$ is the Grüneisen parameter. Using the approach of Eq. 8.34 with and $\delta V/V^0 = 3 \alpha T$ for the thermal expansion

$$S_{ph}(T) = k_B \sum_j^{3N} \ln \left( \frac{k_B T}{\hbar \omega_j^0 (1 - 3 \gamma \alpha T)} \right),$$  

(8.35)

$$S_{ph}(T) = \left[ k_B \sum_j^{3N} \ln \left( \frac{k_B T}{\hbar \omega_j^0} \right) \right] + 9N \gamma \alpha k_B T,$$  

(8.36)

$$S_{ph}(T) = S_{ph}^0(T) + 9N \gamma \alpha k_B T = S_{ph}^0 + S_q,$$  

(8.37)

where the first term, $S_{ph}^0(T)$, is the harmonic phonon entropy with phonon frequencies characteristic of $T = 0$, and unchanged with temperature. Using Eqs. 8.30, 8.31 and 8.37 in the free energy expression for the expanded crystal with phonons, Eq. 8.29

$$F(T) = \frac{9}{2} BV_0 \alpha^2 T^2 + 3Nk_B T - TS_{ph}^0(T) - 9N \gamma \alpha k_B T^2.$$  

(8.38)

First we obtain the equilibrium thermal expansion coefficient, $\alpha$, by calculating

$$\frac{\partial F(T)}{\partial \alpha} = 0 = 9BV_0 \alpha T^2 - 9N \gamma k_B T^2,$$  

$$\alpha = \frac{C_V \gamma}{3BV_0},$$  

(8.39)

where we have used the classical result $C_V = 3Nk_B$.\(^3\) If the electronic entropy depends on volume, an additional term, $C_{elV} \gamma_{el}/(3BV_0)$ is added to the right-hand side of Eq. 8.39.

Substituting Eq. 8.39 for the thermal expansion into 8.38 for the free energy

$$F(T) = F^0(T) - \frac{9}{2} BV_0 \alpha^2 T^2,$$  

(8.40)

where $F^0(T)$ is the harmonic free energy at finite temperature with zero thermal expansion ($\alpha = 0$). We use Eq. 8.39 to express the quasiharmonic entropy $S_q$ of Eq. 8.36 or 8.37 as

$$S_q = 9BV_0 \alpha^2 T$$  

(8.41)

---

\(^3\)The effect of phonon energy is small as discussed in Footnote 2, although there is also a small effect from a reduction in zero-point phonon energy as the solid expands.
8.2. HEAT CAPACITY

Figure 8.5: Heat capacity versus temperature for trial models of cerium metal. Inset is the phonon DOS at 0 K, and the harmonic heat capacity curve was obtained from this DOS ($3R$ is the classical limit). The elastic energy from thermal expansion was added to the harmonic curve to obtain the quasiharmonic one. Using experimental $\alpha$ and $B$ in Eq. 8.39, a $\gamma$ was obtained from Eq. 8.39 that predicted the phonon DOS would soften by 6.5% from 0 to 300 K. (The real heat capacity also includes an electronic contribution.)

(consistent with differentiating Eq. 8.40 with respect to $T$). Finally, if we substitute Eq. 8.41 into Eq. 8.27, we can recover Eq. 8.28. Typical effects on heat capacity are shown in Fig. 8.5. This $S_q$ of Eq. 8.41 should always be considered when comparing the vibrational thermodynamics of different materials at elevated temperatures.

8.2.3 Anharmonic Heat Capacity

In mechanics, the word “anharmonic” describes any oscillator with generalized forces that deviate from linearity with generalized coordinates. In vibrational thermodynamics, “anharmonic” is used more restrictively. The word “quasiharmonic” accounts for the effects of thermal expansion on phonons just considered (e.g., Eqs. 8.28, 8.41), effects that are not strictly harmonic. An implicit assumption of quasiharmonic theory is that an oscillation has a well-defined, long-lived, harmonic-like frequency that changes modestly and predictably with changes in $T$ and $P$.

Equation 8.39 is useful for separating anharmonic from quasiharmonic behavior. Suppose we know (by inelastic scattering measurements for example) the thermal softening of the phonon modes, from which we calculate an average Grüneisen parameter $\gamma$. With the phonon DOS we can also calculate $C_V$ (or its molar quantity), and conventional measurements can provide $B$, $V_0$ (or
molar volume \( v \), and \( \alpha \), accounting for all unknowns in Eq. 8.39. If Eq. 8.39 proves to be true with the \( \gamma \) measured from the phonon softening, it is possible that the solid is "quasiharmonic."

Anharmonic behavior is identified as phonon softening or stiffening that is inconsistent with the Grüneisen parameter \( \gamma \) needed for equality in Eq. 8.39. Anharmonicity involves phenomena beyond those of independent phonons. Higher-order phonon-phonon interactions are the source of anharmonic behavior as discussed in Sect. 9.3.1. Section 9.3.2 discusses electron-phonon interactions, which also vitiate Eq. 8.39. Substantial deviations from quasiharmonic behavior are known for Cr (6), V (7), Ce (8), Mo, W, Si, Ge (9), and anharmonic or non-harmonic effects could be important in many other systems.

### 8.2.4 Thermodynamic Entropy

The temperature dependence of the entropy

\[
\frac{dS(T, V)}{dT} = \left( \frac{\partial S}{\partial T} \right)_V + \left( \frac{\partial S}{\partial V} \right)_T \frac{dV}{dT}.
\]

contains harmonic, quasiharmonic, and anharmonic contributions, along with non-harmonic effects from electron-phonon interactions. For the harmonic part of the vibrational entropy we use Eq. 8.23, with a phonon DOS, \( g_0(\varepsilon) \), from low temperature measurement or calculation (ideally \( T = 0 \) K). The harmonic part contributes to the first term on the right-hand-side of Eq. 8.42, i.e., \( \partial S/\partial T \)_V.

Harmonic phonons undergo no change in frequency with \( T \) or \( V \). The temperature dependence of \( S_h(T) \) originates with the Planck occupancy factors \( n(\varepsilon, T) \).

Quasiharmonic phonons have frequencies that depend on volume only, and make a contribution through the second term on the right-hand-side of Eq. 8.42. At a fixed volume, however, they behave as harmonic oscillators. Their frequencies can change with temperature, but only because thermal expansion alters the volume of the solid.

Anharmonic entropy, \( S_a \), may contribute to either of the two terms on the right-hand side of Eq. 8.42, but especially the first, a pure temperature-dependence of the vibrational entropy at fixed volume caused by changes in the interatomic force constants with temperature. Consider for example a quartic term in the interatomic potential, which alters the force constants with larger thermal displacements. The quartic term does not affect the thermal expansion,

---

4. However, some phonon modes may be anharmonic and others quasiharmonic, or the quasiharmonic modes may have different Grüneisen parameters.

5. It remains a challenge to differentiate between these sources of non-harmonic behavior (5), so sometimes the effects of electron-phonon interactions are called "anharmonic." More often "anharmonic" is reserved for phonon-phonon interactions only.

6. For harmonic solids at high temperatures, differences in the phonon DOS will cause a difference in entropy for different phases, but this difference in entropy does not change with temperature. (At high \( T \), \( S \) increases with \( T \) by the same amount for each phase.)
but it does change the phonon frequencies. It causes a type of “phonon-phonon interaction,” so called because the existence of phonons affects the energy to create new phonons. Other temperature-dependences of force constants at constant volume may arise if the nature of the bonding changes as atoms are displaced further from their equilibrium positions, or perhaps thermal excitations of electrons alter the sensitivity of electronic energy to atom displacements (Section 9.3.2 addresses such effects from adiabatic electron-phonon interactions). The second term in Eq. 8.42 is associated with changes of the interatomic force constants with volume. The quasiharmonic contribution is expected to account for most of it, but the practice of using harmonic entropy expressions, while correct to first order (5), is not fully verified. The cubic term associated with thermal expansion also shortens phonon lifetimes, causing energy spreads of phonon linewidths. This may not alter the vibrational entropy to first order, but again the robustness of the harmonic approximation is not fully tested.

References

8.3 Lattice Dynamics

8.3.1 Atomic Force-Constants

Since most of the atom mass is in the nucleus, for lattice dynamics we consider the Hamiltonian for nuclear motions in a crystal

\[ H_n = \sum_{l,\kappa} \frac{p_{lk}^2}{2m_\kappa} + \Phi, \]

and the characteristics of the atom vibrations require attention to the potential energy, \( \Phi \). Following the development of Maradudin, et al. (10) for a crystal lattice with a basis, the basis vectors are \( \{ \vec{r}_\kappa, \kappa = 1, 2, ..., R \} \). Each atom \( \kappa \), in the unit cell \( l \), is allowed to vibrate about equilibrium with the displacement \( \vec{u}_l(\kappa) \).

The instantaneous position \( \vec{R}_{lk}(t) \) of atom \( lk \) at time \( t \) is

\[ \vec{R}_{lk}(t) = \vec{r}_l + \vec{r}_\kappa + \vec{u}_{lk}(t). \]

We use Cartesian components \( u_{alc} \) for the displacement vector, where \( a = [x, y, z] \). The total potential energy of the crystal, \( \Phi \), is a function of the instantaneous positions of all the atoms in the crystal, but it is expanded in a Taylor series of the atomic displacements about \( \vec{u}_l(\kappa) = 0 \) (i.e., the equilibrium positions of the atoms \( \vec{R}_{lk} = \vec{r}_l + \vec{r}_\kappa \))

\[ \Phi = \Phi_0 + \sum_{alc} \Phi_{alc} u_{alc} \]

\[ + \frac{1}{2} \sum_{alc} \sum_{a'l'c'} \Phi_{aa'lc'c'} u_{alc} u_{a'l'c'} + ... , \]

where the coefficients of the Taylor series are the derivatives of the potential with respect to the displacements:

\[ \Phi_{alc} = \left. \frac{\partial \Phi}{\partial u_{alc}} \right|_0, \]

\[ \Phi_{aa'lc'c'} = \left. \frac{\partial^2 \Phi}{\partial u_{alc} \partial u_{a'l'c'}} \right|_0, \]

where the subscript zero means that derivatives are evaluated in the equilibrium configuration (all displacements equal to zero) and \( \Phi_0 \) is the static potential energy of the crystal. Because the force on any atom must vanish in the equilibrium configuration, we have (10)

\[ \Phi_{alc} = 0 \quad \forall \ a, l, \kappa . \]

In the harmonic approximation of lattice dynamics we keep only the terms of the series written explicitly in Eq. 8.45 – we neglect terms of order three and
higher in the displacements

\[ H_n = \sum_{\ell \kappa} \frac{p_{\ell \kappa}^2}{2m_\kappa} + \Phi_0 + \frac{1}{2} \sum_{\alpha \ell \kappa} \sum_{\beta \ell' \kappa'} \Phi_{\alpha \beta; \ell \kappa; \ell' \kappa'} u_{\alpha \ell \kappa} u_{\beta \ell' \kappa'}, \]  

where \( m_\kappa \) is the mass of the atom at basis index \( \kappa \) of the unit cell. We rewrite the Hamiltonian in matrix form

\[ H_n = \sum_{\ell \kappa} \frac{p_{\ell \kappa}^2}{2m_\kappa} + \Phi_0 + \frac{1}{2} \sum_{\ell \kappa} \sum_{\ell' \kappa'} \Phi_{\ell \kappa; \ell' \kappa'} \Phi^{T}_{\ell' \kappa' \ell \kappa} u_{\ell \kappa} \Phi_{\ell \kappa; \ell' \kappa'}, \]  

where a 3×3 force-constant sub-matrix is defined for each atom pair \((l_\kappa; l'_\kappa')\)

\[ \Phi_{l \kappa; l' \kappa'} = \begin{bmatrix} \Phi_{\alpha \alpha; l \kappa; l' \kappa'} \end{bmatrix}. \]  

If \((l, \kappa) \neq (l', \kappa')\), Eq. 8.47 applies. If \((l, \kappa) = (l', \kappa')\), \( \Phi_{\alpha \alpha; l \kappa; l' \kappa'} \) is a “self-force constant,” derived from the requirement of no overall translation of the crystal

\[ \Phi_{l \kappa; l' \kappa'} = -\sum_{(l', \kappa') \neq (l, \kappa)} \Phi_{l \kappa; l' \kappa'}. \]  

Because \(-\Phi_{l \kappa; l' \kappa'} u_{l' \kappa'}\) is the force acting upon atom \((l \kappa)\) when atom \((l' \kappa')\) is displaced by \( u_{l' \kappa'}\), it follows that \( \Phi_{l \kappa; l' \kappa'} \) must be a real symmetric matrix:

\[ \Phi_{l \kappa; l' \kappa'} = \begin{bmatrix} a & b & c \\ b & d & e \\ c & e & f \end{bmatrix}. \]  

Any crystal is invariant when translated by a lattice vector, so the force constant matrices must also have the following property:

\[ \Phi_{l \kappa; l' \kappa'} = \Phi_{0 \kappa; (l' - l) \kappa} = \Phi_{(l - l') \kappa; 0 \kappa}. \]  

### 8.3.2 Equations of Motion

In the harmonic approximation, the equations of motion for all nuclei are

\[ m_\kappa \ddot{u}_{l \kappa}(t) = -\sum_{l', \kappa'} \Phi_{l \kappa; l' \kappa'} \ddot{u}_{l' \kappa'}(t) \quad \forall \ l, \kappa. \]  

In Eq. 8.55 there are \( 3 \times R \times N_{\text{cell}} \) equations of motion to solve for a finite crystal containing \( N_{\text{cell}} \) unit cells. We seek solutions having the form of plane waves

\[ \ddot{u}_{l \kappa}(t) = \sum_{l', \kappa'} u_{l' \kappa'}(t). \]
of wavevector \( \vec{k} \), angular frequency \( \omega_{k'} \) and “polarization” \( \vec{c}_{kj}(\vec{k}) \) (where \( j \) is a “branch index” discussed below)

\[
\vec{u}_{lki}(t) = \frac{2\hbar}{N m \omega_{kj}} \vec{c}_{kj}(\vec{k}) e^{i(\vec{k} \cdot \vec{r}_l - \omega_{kj} t)},
\]

(8.56)

\[
\vec{u}_{lki}(t) = \hbar \sqrt{\frac{2N(e_{kj}, T) + 1}{N m \epsilon_{kj}}} \vec{c}_{kj}(\vec{k}) e^{i(\vec{k} \cdot \vec{r}_l - \omega_{kj} t)},
\]

(8.57)

where we take the real part to obtain physical displacements. The phase factor, \( e^{i\vec{k} \cdot \vec{r}_l} \), provides all the long-range spatial modulation of \( \vec{u}_{lki}(t) \). The dependence on \( \kappa \), a short-range basis vector index, is taken out of the phase factor and placed in the complex constant \( \vec{c}_{kj}(\vec{k}) \). It is convenient for the \( \vec{c}_{kj}(\vec{k}) \) of Eq. 8.56 to have modulus unity, as does the exponential. The prefactors are consistent with the quantization of energy for one phonon, \( \hbar \omega = m \omega^{2}(u^2) = \frac{1}{2} m \omega^{2} \xi_{\text{max}}, \) or for the thermal population of phonons in mode \( \epsilon_{kj} \), having energy \( E = \epsilon_{kj}[n(\epsilon_{kj}, T) + \frac{1}{2}] \).

(For root-mean-squared displacements, delete the \( \sqrt{2} \) in Eqs. 8.56 and 8.57.)

We impose periodic boundary conditions on the finite crystal. These require the set of possible wavevectors \( \{\vec{k}\} \) to have \( N_{\text{cell}} \) values, a large number that gives a very fine mesh of \( \vec{k} \)-points in reciprocal space. With \( R \) atoms in the basis of the unit cell, the crystal has \( 3 \times R \times N_{\text{cell}} \) vibrational modes, in agreement with its total number of mechanical degrees of freedom. Each wavevector \( \vec{k} \) is associated \( a\text{-priori} \) with \( 3 \times R \) types of vibrational modes, identified by a branch index, \( j \). Each of the \( 3R \) different modes corresponds to a different polarization vector \( \vec{c}_{kj}(\vec{k}) \) and angular frequency \( \omega_{kj} \) (\( 1 \leq j \leq 3R \)), although degeneracies can be induced by symmetry.

### 8.3.3 The Eigenvalue Problem for the Polarization Vector

The polarization vector, \( \vec{c}_{kj}(\vec{k}) \), is a characteristic of each vibrational mode \( \vec{k}, j \). The vector \( \vec{c}_{kj}(\vec{k}) \) contains information on the excursion of each atom \( \kappa \) in the unit cell for the phonon mode mode \( \vec{k}, j \). Specifically, it gives 1) the displacement direction of the atom, and 2) its phase lag in time with respect to the other atoms. The vectors \( \vec{c}_{kj}(\vec{k}) \) for all the atoms in the basis (\( 1 \leq \kappa \leq R \)) and their associated angular frequencies \( \omega_{kj} \) can be calculated by diagonalizing the “dynamical matrix” \( D(\vec{k}) \). The dynamical matrix is obtained by substituting Eq. 8.56 into 8.55. It has the dimensions \( (3N \times 3N) \) and is constructed from \( (3 \times 3) \) submatrices \( D_{\kappa \kappa}(\vec{k}) \)

\[
D(\vec{k}) = \begin{pmatrix}
D_{11}(\vec{k}) & \cdots & D_{1N}(\vec{k}) \\
\vdots & \ddots & \vdots \\
D_{N1}(\vec{k}) & \cdots & D_{NN}(\vec{k})
\end{pmatrix}.
\]

(8.58)
Each sub-matrix $D_{\kappa\kappa'}(\vec{k})$ is the Fourier transform of the force-constant matrix $\Phi_{\kappa l \kappa' l'}$, considered as a function of $(l' - l)$:

$$D_{\kappa\kappa'}(\vec{k}) = \frac{1}{\sqrt{m_\kappa m_{\kappa'}}} \sum_{l'} \Phi_{\kappa l \kappa' l'} e^{i \vec{k} \cdot (\vec{r}_{l'} - \vec{r}_0)}, \quad (8.59)$$

where we set $l = 0$ because the summation is over all $l'$ and the origin is arbitrary. By similarly collecting the polarization vectors into a vector of size $3 \times \mathcal{R}$, we rewrite the system of differential equations (Eq. 8.55) with the plane wave solutions (Eq. 8.56) as an eigenvalue problem:

$$\mathcal{D}(\vec{k}) \vec{e}_j(\vec{k}) = \omega_{kj}^2 \vec{e}_j(\vec{k}), \quad (8.60)$$

where

$$\vec{e}_j(\vec{k}) = \begin{pmatrix} e_{x1}(\vec{k}) \\ e_{y1}(\vec{k}) \\ e_{z1}(\vec{k}) \\ \vdots \\ e_{2N}(\vec{k}) \end{pmatrix}.$$  

(8.61)

It can be shown that the $(3\mathcal{R} \times 3\mathcal{R})$ dynamical matrix $\mathcal{D}(\vec{k})$ is hermitian (for any value of $\vec{k}$). It is fully diagonalizable, and the $\omega_{kj}^2$ are real. The $3\mathcal{R}$ eigenvectors and eigenvalues of the dynamical matrix evaluated at a particular wavevector $\vec{k}$ then correspond to the $3\mathcal{R}$ eigenmodes of vibration of the crystal for that wavevector.

### 8.3.4 Calculation of the Phonon Density of States

To calculate the phonon density of states (DOS) of the crystal, $g(\epsilon)$, the dynamical matrix is diagonalized at a large number of points in reciprocal space (typically covering the first Brillouin zone). The diagonalization of $\mathcal{D}(\vec{k})$ at each $\vec{k}$ point returns $3\mathcal{R}$ eigenvalues of angular frequency $\omega_{kj}^2 (1 \leq j \leq 3\mathcal{R})$, which are then binned into the DOS histogram.

A phonon partial DOS, $g_d(\epsilon)$, is a similar quantity, but it gives the spectral distribution of motion by one atom, the species $d$ in the unit cell. Unlike the total DOS, $g(\epsilon)$, the eigenvalues of the crystal are not weighted equally in the partial DOS

$$g_d(\epsilon) = \sum_g \sum_{a\alpha} \delta_{dc} |e_{a\alpha}(\vec{k})|^2 g(\epsilon), \quad (8.62)$$
where the Kroneker delta is zero unless the atom species \( d \) is at the site \( \kappa \) (when \( \delta_{dd} = 1 \)). Because the eigenvalues of the dynamical matrix are normalized for each \( \vec{k} \) as

\[
\sum_{\alpha j} |e_{\alpha j}(\vec{k})|^2 = 1 , \tag{8.63}
\]

the total DOS is the sum of the partial DOSs of all atoms in the unit cell,

\[
g(\varepsilon) = \sum_d g_d(\varepsilon) . \tag{8.64}
\]

To calculate a partial DOS, for each diagonalization of the dynamical matrix at a specific \( \vec{k} \) point, the partial DOS histogram for atom \( d \) is incremented at frequency \( \omega_{\vec{k}j} \) by the amount \( \sum_a e_{adj}(\vec{k})^* e_{adj}(\vec{k}) \). The partial DOS for atom \( d \) is large at energies where there are many modes with large displacements of atom \( d \). The histogram for the phonon partial DOSs are computed simultaneously with the total DOS.

### 8.3.5 Symmetry Constraints on the Force-Constant Matrices

#### References


### 8.4 Computer Simulations of Lattice Dynamics

#### 8.4.1 Density Functional Theory

A large branch of computational materials science owes its predictive power to developments in density functional theory (DFT), in conjunction with advances in computing power. There are excellent reviews and books on density functional theory and its applications to materials, some of which offer a broader context of condensed matter theory (11)(12)(13)(14). This rich field is far too extensive to cover here. The techniques for calculating phonons from first principles also deserve a much more extensive presentation (43)(15)(16)(17)(18)(19)(20)(21)(22)(23).
This brief overview of some key methods and concepts is presented with apologies that numerous intellectual achievements in this rapidly-moving field cannot be covered.

Density functional theory elevates the importance of the electron density with respect to the electron wavefunctions, relying on a theorem that all properties of an electronic system can be obtained from the ground state electron density (24)(25). A functional relationship must exist between the density and the physical quantity of interest, but finding this relationship is a challenge. The successful approach developed by Kohn and Sham (26) assumes that the ground state electron density of a system of interacting electrons can be equal to that of a non-interacting system of electrons, with an interacting electron density. Working with independent electrons makes calculations practical, freeing us from time-ordered-fermion field operators for electron-electron interactions. In particular, the electron exchange and Coulomb correlation energy, which are nonlocal multibody interactions, can be successfully approximated with functionals of the local density, or the density and its local gradients, for example. The density is obtained from the electron wavefunctions, so computational algorithms require iteration to self consistency of the potential and the wavefunctions. The local density functional theory has been surprisingly successful, although problems occur for strongly-correlated electron systems.

8.4.2 First-Principles Phonon Calculations

Principles for calculating phonon dispersions from the electronic energy were outlined in the classic work of Ziman (27) and others (28). Two approaches are used. The direct methods impose displacements of atoms, and calculate energies of distorted configurations. In the supercell method, pairs of atoms are displaced by \( \vec{u} \) and \( \vec{u}' \) along directions of symmetry, for example, and from the energies of the distorted structures, \( E(\vec{u}, \vec{u}') \), the force constants \( \Phi_{\alpha\beta,\kappa\kappa'} \) of Eq. 8.55 can be calculated as

\[
\Phi_{\alpha\beta,\kappa\kappa'} = \frac{\partial^2 E(\vec{u}, \vec{u}')}{{\partial u}_\alpha {\partial u'}_\beta} \bigg|_0,
\]

(cf., Eq. 8.47). With these ab-initio force constants, the lattice dynamics can be calculated (17). The Hellman-Feynman theorem shows that errors in the wavefunctions that cause only second-order errors in energy are responsible for first-order errors in forces, so energy convergence is stringent (43)(21). With the frozen phonon method, the potential energy of an individual phonon is calculated directly by generating periodic distortions of a cell or supercell (29)(30). Calculations with direct methods are versatile, and can account for unusual structures. Frozen phonons of different amplitudes are a natural way to account for anharmonic potential energies (31)(32)(33), although a good sampling of all phonons is challenging. When periodic boundary conditions are used, the cell should be comfortably larger than the longest-range interatomic force, and the phonon wavevector should be commensurate with the structure. With many
atoms and electrons in a large supercell, however, the calculations can become unwieldy, especially since the computation time often scales as the cube of the number of electrons.

The second approach calculates the sensitivities to small density modulation, usually with a linear response theory (15)(16)(18)(34). These are second derivatives of the energy, and the simplest approach to the problem is with second order perturbation theory. For periodic crystals with both electrons and phonons expressed as Bloch functions, the density susceptibility function for the Fourier coefficient ($\vec{k}$) of the electron density depends on the Fourier coefficient of the phonon modulation ($\vec{k}'$) as

$$\chi(\vec{k}, \vec{k}') = 2 \sum_i \sum_j \langle j | e^{-i\vec{k} \cdot \vec{r}} | i \rangle \langle j | e^{i\vec{k}' \cdot \vec{r}} | i \rangle \frac{\epsilon_i - \epsilon_j}{\epsilon_i - \epsilon_j},$$

where the intermediate electronic states $|j\rangle$ are unoccupied, and the states $|i\rangle$ are occupied. A considerable simplification is that $\vec{k}$ and $\vec{k}'$ must be related by a reciprocal lattice vector, reducing the number of terms in Eq. 8.66, although for more realistic interacting systems, the situation is not so simple. A variational approach for obtaining the energy to second order has been developed to obtain a linear response to periodic perturbations (18). In this method, the dynamical matrix can be calculated directly (19). It is also possible to recover real-space force constants by Fourier transformation of the dynamical matrix for appropriate selections of $\vec{k}$.

### 8.4.3 Molecular Dynamics

Molecular dynamics simulations integrate equations of motion to follow in time the positions and velocities of all atoms. Many molecular dynamics simulations are classical or semi-classical, using forces obtained as gradients of model potential energy functions, for example. The atoms are allowed to move under Newton’s law for a short time, and the forces are recalculated for the new configuration of atoms. These simulations can be extended beyond a microcanonical ensemble by allowing contact of a small simulation system with a reservoir, accounting for fluctuations of temperature or pressure, for example. Realistic thermostatic or barostatic control can be achieved by extending the dynamical system with degrees of freedom that represent the external system (35)(36). Molecular dynamics is not restricted to harmonic forces, and large-amplitude displacements allow investigations of anharmonic behavior. Accurate force fields that account for all detail of the local atomic environment can be a challenge. Such force fields can be obtained from quantum mechanics, but often they are empirical.

The velocity-velocity autocorrelation function $\langle \vec{v}(t) | \vec{v}(0) \rangle$ can be calculated from the trajectory of each atom in a molecular dynamics simulation.\(^8\) The

---

\(^8\)The notation indicates that the velocity history of each atom is multiplied by the velocity history
Fourier transform of the velocity-velocity autocorrelation function gives the phonon DOS (37)

\[ g(\omega) = \int e^{i\omega t} \frac{\langle v(t)|v(0)\rangle}{\langle v(0)|v(0)\rangle} \, dt. \tag{8.67} \]

In another approach, it is possible to obtain differences in thermodynamic quantities such as phonon free energy by parameterizing a Hamiltonian to allow the system to switch from one structure to another, and integrating along the path of this switching parameter.

It is possible to solve for the forces on the moving nuclei by performing a full DFT calculation at each timestep of the simulation, but this is cumbersome. The Car–Parrinello algorithm makes practical quantum molecular dynamics simulations of greater complexity (38)(39). It uses the electron degrees of freedom as dynamical variables in a set of coupled equations of motion for the nuclei and the electrons. The nuclei have a classical kinetic energy of \( M/2 \left[ \frac{d^2u}{dt^2} \right] \). The trick to the algorithm is that the electrons are given a fictitious kinetic energy of \( \mu/2 \left[ \frac{d\psi}{dt} \right]^2 \), with a mass \( \mu \) that is an adjustable parameter (that helps with the stability of the numerical calculations). With this coupling of nuclear and electronic motions, constrained by the orthogonality of the electron wavefunctions, the electrons stay near their ground states during the simulation. The electron wavefunctions do not need to be recalculated at each step of the simulation, offering an improvement in efficiency. The Car–Parrinello algorithm is an adiabatic approximation.

8.5 Group Theory and Lattice Dynamics

8.5.1 Real Space

The paper by A. A. Maradudin and S. H. Vosko “Symmetry Properties of the Normal Modes of a Crystal”, Reviews of Modern Physics 40, 1-37 (1968), shows how group theory can be used to understand the normal modes of crystal vibrations. The focus of their substantial manuscript is on using the symmetry of the reciprocal lattice to find the eigenvectors of the dynamical matrix and classify them. The degeneracies of the different normal modes are addressed rigorously, and their association with the symmetry elements of the crystal are useful in labeling them. Projection operator methods offer the possibility of finding eigenvectors and eigenvalues without diagonalizing the dynamical matrix.

The paper by J. L. Warren “Further Considerations on the Symmetry Properties of the Normal Modes of a Crystal”, Reviews of Modern Physics 40, 38-76 (1968), also describes the point group of the bond. The bond means the interactions between an atom and its neighbors. The group of the bond refers

\[ \text{shifted by } t, \text{ and integrated over all time.} \]
to the real space symmetries of the interatomic interactions, i.e., how the force constants transform under the point group operations at a central atom.

Consider the space group operator in Seitz notation:

\[ S \equiv [S | \vec{v}(S) + \vec{x}(m)] , \tag{8.68} \]

where \( S \) is a rotation operation, \( \vec{x}(m) \) is a lattice translation, and \( \vec{v}(S) \) is an additional displacement that is required for non-symmorphic crystals (i.e., crystals with screw axes or glide planes).\(^9\) Apply this operator to a displacement vector \( u \) (the displacement of an atom off its site during a vibration):

\[ u_\alpha'(L, K) = \sum_S S_{\alpha\beta} u_\beta(l, \kappa) , \tag{8.69} \]

where the operation serves to mix the Cartesian components (denoted subscript \( \alpha \)) of the displacement vector for the atom at \( l, \kappa \), and translates the vector to the atom at \( L, K \). The potential energy of the crystal depends quadratically on the \( u_\alpha \) times a force constant matrix involving the spatial derivatives of the potential along the Cartesian axes of the \( u_\alpha \). The potential energy of the crystal is invariant under symmetry operations of (8.68). This leads to the equality:

\[ \Phi_{\mu
u}(LK, L'K') = \sum_{\alpha\beta} S_{\mu\alpha} S_{\nu\beta} \Phi_{\alpha\beta}(l, l' \kappa') . \tag{8.70} \]

The matrix equation (8.70) must be true for all valid force constant matrices \( \Phi_{\alpha\beta}(l, l' \kappa') \). It is instructive to set \( S = I \), the identity, so there is no rotation, and only translations are imposed. This can be used to show that force constants can depend only on relative separations between atoms, not absolute positions.

A number of other tests of force constants can be performed with (8.70). Imposing rotations that mix the Cartesian axes can be used to demonstrate that some force constants are equal. Force constants that must equal their negative are found – this result means that the force constant is zero. Equation (8.70) could be handy for determining allowed force constants if a set of space group matrices, \( S \), are available.

### 8.5.2 \( k \)-space

**Quantum Mechanics**

In quantum mechanics, group theory addresses the symmetry of the hamiltonian, \( H \). If an operator, \( R \), commutes with \( H \),

\[ RH = HR , \tag{8.71} \]

\[ \sum_j R_{ij} H_{jk} = \sum_j H_{ij} R_{jk} , \tag{8.72} \]

---

\(^9\)Note that this vector \( \vec{v}(S) \) mixes the rotation and translation operations.
it is convenient to select basis functions for which \( R \) is diagonal, so the single remaining terms in the sums are:

\[
R_{ii} H_{ik} = H_{ik} R_{kk}, \quad (8.73)
\]

\[
(R_{ii} - R_{kk}) H_{ik} = 0. \quad (8.74)
\]

Equation (8.74) shows that when \( i \neq k \), then the off-diagonal elements \( H_{ik} = 0 \).

The basis functions for which \( R \) is diagonal are the same for which \( H \) is diagonal.

For example, consider the rotational symmetry of a hydrogen atom about the \( \hat{z} \)-axis. The Abelian group of rotations has 1-dimensional representations in terms of functions \( e^{i\phi} \). Done. Obtaining the \( \phi \)-dependence this way was much easier than solving the Schrödinger equation in spherical coordinates.

**Lattice Dynamics**

The analogous question for lattice dynamics is, “Can we use symmetry to get the eigenvectors of the dynamical matrix without solving the eigenvalue problem in detail?” The dynamical matrix is:

\[
D_{\alpha\beta}(\kappa\kappa'|\vec{k}) = \frac{1}{\sqrt{M_\kappa M_{\kappa'}}} \sum_l \Phi_{\alpha\beta}(l; \kappa') \exp[-i\vec{k} \cdot (\vec{x}(l) - \vec{x}(l'))], \quad (8.75)
\]

and the relevant eigenvalue equation is:

\[
\sum_{\beta\kappa'} D_{\alpha\beta}(\kappa\kappa'|\vec{k}) u_{\beta}(\kappa') = \omega^2 u_{\alpha}(\kappa). \quad (8.76)
\]

All functions in (8.76) live in reciprocal space. The entire paper of Maradudin and Vosko is in reciprocal space, or \( k \)-space.

Consider the matrix \( \Gamma_S(\vec{k}) \) that performs symmetry operations \( S \) on the dynamical matrix. Doing two such operations is tricky, since the \( \vec{k} \) used in constructing the matrix for the second operation depends on the result of the first operation. Maradudin and Vosko choose instead to work with the space group of a single wavevector \( \vec{k} \), and designate the symmetry operations by \( R_i = \{ R | \vec{v}(S) + \vec{x}(m) \} \). These symmetry operations on \( \vec{k} \) produce equivalent wavevectors. In particular, the rotational part of \( R_i \) therefore must generate a \( \vec{k} \) that differs only by a reciprocal lattice vector \( \vec{g} \):

\[
R \vec{k} = \vec{k} - \vec{g}. \quad (8.77)
\]

This is quite restrictive, and means that \( \vec{g} \) is zero for most vectors \( \vec{k} \). The situation is much like the allowed \( k \)-vectors for ordered structures in the Landau-Khachaturyan formalism of second-order phase transitions. The operations of (8.77) eliminate the equivalent vectors from the star of \( \vec{k} \). Inequivalent \( \vec{k} \) can exist only at special points such as the boundaries of Brillouin zones.
### Table 8.1: Elementary Variables Used by Maradudin and Vosko

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>( l, l' )</td>
<td>indices of unit cells</td>
</tr>
<tr>
<td>( \kappa, \kappa' )</td>
<td>indices of atom in basis</td>
</tr>
<tr>
<td>( r )</td>
<td>number of atoms in basis</td>
</tr>
<tr>
<td>( \vec{x}_{l,\kappa} )</td>
<td>atom site in a crystal</td>
</tr>
<tr>
<td>( \vec{x}_l )</td>
<td>lattice site in a crystal</td>
</tr>
<tr>
<td>( \vec{x}_{\kappa} )</td>
<td>basis vector</td>
</tr>
<tr>
<td>( M_{\kappa} )</td>
<td>mass of atom in basis</td>
</tr>
<tr>
<td>( \alpha, \beta )</td>
<td>Cartesian indices ( {x, y, z} )</td>
</tr>
<tr>
<td>( \vec{u}(\kappa) ) and ( u_\alpha(\kappa), u_\beta(\kappa) )</td>
<td>atom displacement in a phonon, and Cartesian components</td>
</tr>
<tr>
<td>( \Phi_{\alpha\beta}(l; l', \kappa') )</td>
<td>force constant between two atoms</td>
</tr>
<tr>
<td>( \vec{e}(\vec{k}, \sigma, \lambda) )</td>
<td>phonon polarization vector (with components for all ( \kappa ) in unit cell)</td>
</tr>
<tr>
<td>( \sigma ) (or ( s ))</td>
<td>denotes (or index for) a distinct value of ( \omega_j^2 )</td>
</tr>
<tr>
<td>( \lambda ) (or ( a ))</td>
<td>denotes (or index for) independent eigenvector for each ( \sigma ) or ( \omega_j^2 )</td>
</tr>
<tr>
<td>( f_\sigma )</td>
<td>number of eigenvectors for each degenerate energy ((1 \leq a \leq f_\sigma))</td>
</tr>
<tr>
<td>( S \equiv {S</td>
<td>\vec{v}(S) + \vec{x}(m)} )</td>
</tr>
<tr>
<td>( h )</td>
<td>order of the group (number of elements)</td>
</tr>
<tr>
<td>( S )</td>
<td>matrix of rotation or improper rotation</td>
</tr>
<tr>
<td>( \vec{v}(S) )</td>
<td>little displacement vector for a screw axis operation</td>
</tr>
<tr>
<td>( \vec{x}(m) )</td>
<td>lattice translation vector</td>
</tr>
<tr>
<td>( \Gamma_S(\vec{k}; {S</td>
<td>\vec{v}(S) + \vec{x}(m)}) )</td>
</tr>
<tr>
<td>( R_i \equiv {R</td>
<td>\vec{v}(S) + \vec{x}(m)} )</td>
</tr>
</tbody>
</table>
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So although:

\[ D(\mathbf{k}_S) = \Gamma_S^{-1}(\mathbf{k}) D(\mathbf{k}) \Gamma_S(\mathbf{k}) , \tag{8.78} \]

and therefore in general:

\[ D(\mathbf{k}) \neq \Gamma^{-1}_S(\mathbf{k}) D(\mathbf{k}) \Gamma_S(\mathbf{k}) , \tag{8.79} \]

with the restriction to the group of the wavevector \( \mathbf{k} \), the unitary matrices \( \Gamma_R \) commute with \( D \):

\[ D(\mathbf{k}) = \Gamma_R^{-1}(\mathbf{k}) D(\mathbf{k}) \Gamma_R(\mathbf{k}) . \tag{8.80} \]

These \( \{\Gamma_R(\mathbf{k})\} \) actually form a 3\( r \)-dimensional unitary representation of the space group of the wavevector, \( \mathbf{k} \). They could be used to develop the symmetry properties of the eigenvectors of the dynamical matrix, but this is not the approach of Maradudin and Vosko.

**Multiplier Representation**

Maradudin and Vosko define a new matrix \( T(\mathbf{k}; R_i) \), which differs from the \( \Gamma_R(\mathbf{k}) \) by an exponential phase factor of modulus unity:

\[ T(\mathbf{k}; R_i | \mathbf{v}(S) + \mathbf{x}(m)) = \exp[i \mathbf{g} \cdot (\mathbf{v}(S) + \mathbf{x}(m))] \times \Gamma_R(\mathbf{k}; R_i | \mathbf{v}(S) + \mathbf{x}(m)) , \tag{8.81} \]

This choice of \( T(\mathbf{k}; R_i) \) is somewhat unusual because these matrices do not form a group in the usual sense – the product of two of them is not an element of the group:

\[ T(\mathbf{k}; R_i) T(\mathbf{k}; R_j) = \exp[i \mathbf{g} \cdot (\mathbf{v}(R_i) + \mathbf{x}(m))] T(\mathbf{k}; R_i R_j) . \tag{8.82} \]

(where the product of exponentials of two cases of (8.81) were simplified by noting the restrictive condition (8.77) on the allowed \( R \), giving the condition \( \mathbf{g} \cdot \mathbf{x}(m) = 2\pi m \), and exponential phase factors of +1). The phase factor \( \exp[i \mathbf{g} \cdot (\mathbf{v}(S)) \) can differ from +1, although it differs from +1 only for crystals with screw axes AND special \( k \)-points.\(^{10}\)

Matrices \( \{T(\mathbf{k}; R_i)\} \) that obey a multiplication rule of (8.82) are said to form a “multiplier representation” of the group of \( \{R_i\} \). The phase factor is the “multiplier.” For most crystals this is +1, and we are back to ordinary group representations. Even if the multiplier is complex, however, if we can get the

\(^{10}\)Is it important to study phonons (or magnons) at special \( k \)-points in crystals with screw axes (non-symmorphic crystals)?
eigenvectors of these $T$, we can get the eigenvectors of the dynamical matrix, $D$. We rewrite (8.76):

$$D(\vec{k})\vec{e}(\vec{k}\sigma\lambda) = \omega_\sigma^2 \vec{e}(\vec{k}\sigma\lambda), \quad \lambda = 1, 2, 3 \ldots f_\sigma, \quad (8.83)$$

noting that:

$$D(\vec{k})[T(\vec{k}; R_i)\vec{e}(\vec{k}\sigma\lambda)] = \omega_\sigma^2 [T(\vec{k}; R_i)\vec{e}(\vec{k}\sigma\lambda)]. \quad (8.84)$$

The $\{T\}$ mix the degenerate $\vec{e}(\vec{k}\sigma\lambda)$ (i.e., they mix those $\lambda$ that go with the same $\sigma$). Showing this more clearly, we write:

$$T(\vec{k}; R_i)\vec{e}(\vec{k}\sigma\lambda) = \sum_{\lambda'} f_{\sigma} X_{\lambda'\lambda}(\sigma) \tau(\sigma)(\vec{k}; R_i) \vec{e}(\vec{k}\sigma\lambda'), \quad (8.85)$$

for every $R_i$ in the group of the wavevector $\vec{k}$. The $f_{\sigma}$ functions transform among themselves under $T(\vec{k}; R_i)$. Furthermore, from (8.80) we can show that $T(\vec{k}; R_i)$ commutes with $D$. These conditions are sufficient to show that the $\{\tau(\sigma)(\vec{k}; R_i)\}$ provide an $f_{\sigma}$-dimensional irreducible multiplier representation of the point group of the wavevector $\vec{k}$.

Now make a vector of all eigenvectors:

$$\vec{e}(\vec{k}) = \left[ \hat{\vec{e}}(\vec{k}\sigma_1\lambda_1), \hat{\vec{e}}(\vec{k}\sigma_1\lambda_2), \ldots \hat{\vec{e}}(\vec{k}\sigma_2\lambda_1), \hat{\vec{e}}(\vec{k}\sigma_2\lambda_2), \ldots \right], \quad (8.86)$$

which transforms as:

$$T(\vec{k}; R_i)\vec{e}(\vec{k}) = \vec{e}(\vec{k}) \Delta(\vec{k}; R_i), \quad (8.87)$$

and the $3r \times 3r$ dimensional matrix $\Delta(\vec{k}; R_i)$ has the block-diagonal form:

$$\Delta(\vec{k}; R_i) = \begin{bmatrix} \tau^{(1)}(\vec{k}; R_i) & 0 & 0 & \ldots \\ 0 & \tau^{(2)}(\vec{k}; R_i) & 0 & \ldots \\ 0 & 0 & \tau^{(3)}(\vec{k}; R_i) & \ldots \end{bmatrix}. \quad (8.88)$$

The unitary matrices $\tau^{(\sigma)}(\vec{k}; R_i)$ are known for all 230 space groups. In a formal sense, the problem is solved.

**Projection Operators**

In a practical sense, we can generate the eigenfunctions by projection operator machinery. Maradudin and Vosko show that:

$$P^{(\sigma)}_{\lambda A}(\vec{k}) = f_{\sigma}/\hbar \sum_{R_i} \tau^{(\sigma)}_{\lambda A}(\vec{k}; R_i)^* T(\vec{k}; R_i) \quad (8.89)$$
is a projection operator.

The projection operator method is considered standard practice, and is not
developed further by Maradudin and Vosko. Perhaps it is useful to review it
in the context of group theory in quantum mechanics.

Suppose a set of \( \{ \phi_{\lambda}^{(j)} \} \) are partner functions in the representation \((j)\), and
transform among themselves under operation of the symmetry operator \( P_{\mathbf{R}} \).
Explicitly:

\[
P_{\mathbf{R}} \phi_{\lambda}^{(j)} = \sum_{\lambda=1}^{l_j} \phi_{\lambda}^{(j)} \Gamma_{\lambda\kappa}^{(j)}(\mathbf{R}),
\]

Multiply by the complex conjugates of all representation matrices and sum over
all elements in the group:

\[
\sum_{\mathbf{R}} \Gamma_{\lambda\kappa}^{(j)}(\mathbf{R})^* P_{\mathbf{R}} \phi_{\lambda}^{(j)} = \sum_{\mathbf{R}} \sum_{\lambda=1}^{l_j} \Gamma_{\lambda\kappa}^{(j)}(\mathbf{R})^* \Gamma_{\lambda\kappa}^{(j)}(\mathbf{R}) \phi_{\lambda}^{(j)},
\]

where the last line was obtained through the Great Orthogonality Theorem of
group representation theory. Equation (8.92) shows how a projection operator,
\( P_{\lambda\kappa}^{(j)} \),

\[
P_{\lambda\kappa}^{(j)} = \frac{l_j}{h} \sum_{\mathbf{R}} \Gamma_{\lambda\kappa}^{(j)}(\mathbf{R})^* P_{\mathbf{R}},
\]

pulls out the function \( \phi_{\lambda}^{(j)} \).

For the eigenvectors of the dynamical matrix for a particular \( \mathbf{k} \), the projec-
tion operator of (8.89) does the same thing. Since the matricies \( \tau^{(\sigma)}(\mathbf{k}, \mathbf{R}) \) are
available, one could use projection operator techniques to operate on an arbi-
trary vector and generate all eigenfunctions of the dynamical matrix. Labels
can be assigned to various phonons based on group theory designations. An-
other step could be to substitute the \( \mathbf{e}^{(\sigma \lambda \alpha)}(\mathbf{k}) \) into (8.83) and obtain \( \omega_{\alpha}^2 \) by matrix
multiplication rather than by matrix inversion. Is this useful, or just elegant?
Projection operator methods will reveal all degeneracies of normal modes for
a particular \( \mathbf{k} \). In such cases where a set of partner eigenvectors is found, it is
not possible to obtain them directly. This requires diagonalizing the dynamical
matrix to solve for the eigenvectors, or more accurately a block diagonal form
of the dynamical matrix as in (8.88).

Compatibility Relations

The paper by J. L. Warren “Further Considerations on the Symmetry Properties
of the Normal Modes of a Crystal”, Reviews of Modern Physics 40, 38-76 (1968),
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adds to the discussion of Maradudin and Vosko. Compatibility relationships are described in more detail in Warren’s paper. Compatibility relationships refer to the new degeneracies that appear when special high symmetry directions in the Brillouin zone come together at special points, such as at the point \( \Gamma \) at the center of the zone, or at the points \( \{X, W, K, L\} \) on the surface of the zone.

For example, when moving along a line in \( k \)-space along the (100) direction, an abrupt change in symmetry occurs when the edge of the zone (X) or center of the zone (\( \Gamma \)) is reached. The source of this new symmetry is the fact that at these special points, translational vectors of the reciprocal lattice cause the points to become equivalent (identical) to other points in the reciprocal lattice.\(^{11}\)

For example, at an arbitrary position along the (100) direction, the symmetry elements are \( \{E, C_4^2, 2C_4, 2\sigma_v, 2\sigma_d\} \). This fourfold axis obviously does not have threefold symmetry elements, but the situation changes at the origin, point \( \Gamma \). The (111) direction, which includes a threefold axis, also converges at \( \Gamma \). Some representations along the (100) axis are expected to be compatible with some representations along the (111) axis when the two axes meet at the origin. The larger symmetry group at the origin, point \( \Gamma \), contains the elements of the threefold and fourfold axes as subgroups. Some representations of the groups from the threefold and fourfold axes are compatible at \( \Gamma \), while some are not. Those that are compatible at \( \Gamma \) must have eigenvalues that are degenerate in energy. Their associated phonon branches rigorously converge in energy at \( \Gamma \).

The way to test for compatibility is to inspect the characters, \( \chi(R) = \text{Tr}[\Gamma(R)] \), of the representations of the three groups. This can be done with the decomposition formula of group representation theory in exactly the same way as is done for analyzing the degeneracies in crystal field theory. (After all, we are considering only the point group of the wavevector.) The decomposition formula for the characters is:

\[
a_i = \frac{1}{N} \sum_{R} \chi(R)^* \chi(R),
\]

(8.94)

where \( a_i \) is the number of times that a particular representation of the subgroup (e.g., of the threefold axis) appears in the group representation of the special point (e.g., \( \Gamma \)).

Equation (8.94) is most easily used by inspection of character tables.\(^{12}\) For example, if all characters of a representation along the fourfold axis match the characters of a representation at \( \Gamma \), these representations are compatible. In a more complicated situation where there is only one \( a_i = 1 \), the sum of the characters from the representations along the fourfold axis must equal the character of a representation at \( \Gamma \). For the simplest example, the characters of all symmetry elements in the \( \Delta_1 \) representation of the group of the fourfold

---

\(^{11}\)These \( k \)-vectors are not identical when they are just slightly displaced off these special points, and related eigenvectors have to be considered independently.

\(^{12}\)These are found in the classic paper by L. P. Bouckaert, R. Smoluchowski, and E. P. Wigner, Phys. Rev. 50, 58 (1936), and an excellent description is provided in Chapter 8 of M. Tinkham’s book on Group Theory and Quantum Mechanics.
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axis are $+1$. The same symmetry elements have characters $+1$ for the $\Gamma$ point for the representation $\Gamma_1$. The threefold axis has a different set of symmetry elements than the fourfold axis, but these elements are also $+1$ for the $\Gamma_1$ representation, and are $+1$ only for the $\Lambda_1$ representation. We deduce that the $\Delta_1$ representation of the (100) fourfold axis and the $\Lambda_1$ representation of the (111) threefold are compatible at the point $\Gamma$. These two representations have the same basis functions as $\Gamma_1$, but the symmetry operators at $\Gamma$ interchange these basis functions because they are degenerate in energy. The significance that the basis functions of the $\Delta_1$ and $\Lambda_1$ representations must have exactly the same energy at the point $\Gamma$ in $k$-space. They are not degenerate just away from the $\Gamma$ point (unless accidentally, or in the case of no crystal potential), but group theory says that they are rigorously degenerate at $\Gamma$. This is a result of potential use in the analysis of phonon data. We can determine the required conditions for the merging of dispersion curves at special point in $k$-space. (Incidentally, $s$-type functions are suitable basis functions for the symmetric representations in this simple example.)

8.5.3 Time-reversal symmetry in the dynamical matrices

Time reversal symmetry must be considered in magnetic systems. Consider the spin as a rotation in a particular direction. The spin is flipped upon time reversal. Time reversal symmetry must also be considered in certain cases for non-magnetic systems, and results in additional degeneracies. In most cases time-reversal symmetry does not result in additional degeneracies, however. (Perhaps we could consider additional degeneracies as “accidental,” but get the software to flag them so we can inspect for this later.)

When the Hamiltonian is invariant by the time-reversal operation (the direction of momentum, or spin is inverted), the system is said to have the time-reversal symmetry. The system without the external magnetic field has generally the time-reversal symmetry. The time-reversal symmetry often gives the nontrivial physical significance like the additional degeneracy of the system. One of most well-known examples is the Kramers degeneracy, i.e. a system of an odd number of electrons has the extra degeneracy, but an even number does not. Of course, the present ionic lattice system has such symmetry, too. Therefore, it needs to be checked if there occur additional degeneracies or not due to the symmetry, which can be done by investigating the symmetry of the dynamical matrices.

Maradudin and Vosko begins the discussion by considering the additional rotational element $S\cdot\vec{k} = -\vec{k}$ in the point group of the crystal. In the same way as $R$ (in the the point group $G_0$), with each element $S\cdot R$ (in the coset $S\cdot G_0$) we associate a new matrix operator $T(\vec{k}; S\cdot R)$, which is defined anti-unitary as it should be. The analysis for $T(\vec{k}; R)$ in section 4.4.3 can be extended straightforwardly to include the anti-unitary matrix operator $T(\vec{k}; A)$ (the element $S\cdot R$ will be denoted by $A$ hereafter). It is highly desirable to derive the condition for the existence of extra degeneracies due to the time-reversal symmetry, by look-
ing into the linear dependence of the eigenvectors \( \vec{\tau}(k; A) \) and \( T(\vec{k}; A)\vec{\tau}(k; A) \) (it is clear both should be eigenvectors because the system has the time-reversal symmetry). We define

\[
\vec{\tau}(k; A) = T(\vec{k}; A)\vec{\tau}(k; A)
\]

and consider and compare the transformation properties of \( \vec{\tau}(k; A) \) and \( \vec{\tau}(k; A) \) under \( T(\vec{k}; R) \)

\[
T(\vec{k}; R)\vec{\tau}(k; A) = \sum_{\lambda'} f_\lambda \tau^{(0)}(\vec{k}, R)\vec{\tau}(k; A'),
\]

\[
T(\vec{k}; R)\vec{\tau}(k; A) = \sum_{\lambda'} \bar{\tau}^{(0)}(\vec{k}, R)\vec{\tau}(k; A'),
\]

where it is found that the irreducible multiplier representations \( \{ \vec{\tau}(\vec{k}; R) \} \) and \( \{ \tau^{(0)}(\vec{k}; R) \} \) belong to the same factor system. They can be either equivalent or inequivalent. In particular, they can be separated into three types depending on the relationship between two irreducible multiplier representations.

First, we think of a case the irreducible multiplier representations \( \vec{\tau}(\vec{k}; R) \) and \( \tau^{(0)}(\vec{k}; R) \) are inequivalent, which will be referred to as the third type. In this case, the eigenvectors \( \vec{\tau}(k; A) \) and \( \vec{\tau}(k; A) \) are orthogonal and the \( f_\lambda \)-fold degeneracy in the dynamical matrix is doubled by \( 2f_\lambda \) by the time-reversal symmetry.

On the other hand, it is a bit complicated for an alternative situation \( \vec{\tau}(\vec{k}; R) \), the uniqueness up to a phase factor is followed for the matrix \( \beta \). By Schur’s Lemmas, the matrix \( \beta\bar{\tau}(\vec{k}; A_{0}) \) commuting with all the matrices \( \{ \tau^{(0)}(\vec{k}; R) \} \) of an irreducible multiplier representation of the point group must be proportional to the unit matrix. This results in two cases;

\[
\beta\beta^* = \phi(\vec{k}; A_{0}; A_{0})\tau^{(0)}(\vec{k}; A_{0})
\]

\[
\beta\beta^* = -\phi(\vec{k}; A_{0}; A_{0})\tau^{(0)}(\vec{k}; A_{0})
\]

where \( \phi(\vec{k}; A_{0}; A_{0}) \) is a ”multiplier”. Further, as is often the case, \( \beta\beta^* = \pm 1 \) for a suitable element \( A_{0} \) in \( S_{G_{0}} \). Now we can turn our attention to the problem of liner independence of \( \vec{\tau}(k; A) \) and \( \vec{\tau}(k; A) \). For the purpose, it would be easy to investigate the scalar product of two eigenvectors. For the necessary algebra,
it is important to note the scalar products of a unitary and an anti-unitary
transformation on vectors are distinguished as
\[
\langle T(\vec{k}; R) \phi, T(\vec{k}; R) \psi \rangle = \langle \phi, \psi \rangle ,
\]
\[
\langle T(\vec{k}; A_0) \phi, T(\vec{k}; A_0) \psi \rangle = \langle \psi, \phi \rangle ,
\]
and also the properties for the successive transformations under \( T(\vec{k}; R) \) and \( T(\vec{k}; A_0) \) on the eigenvectors \( \vec{e}(ksa\lambda) \) and \( \vec{\bar{e}}(ksa\lambda) \). After a little algebra, it is
finally found that, adopting \( \beta_\beta^* = \pm 1 \),
\[
\langle \vec{\bar{e}}(ksa\lambda'), \vec{e}(ksa\lambda) \rangle = \pm \langle \vec{\bar{e}}(ksa\lambda'), \vec{e}(ksa\lambda) \rangle ,
\]
where the upper plus sign is corresponding to a case in Eq.(4) and the lower
minus sign to a case in Eq.(5), respectively. Clearly, if \( \beta_\beta^* \) satisfies Eq.(5), then
\[
\langle \vec{\bar{e}}(ksa\lambda'), \vec{e}(ksa\lambda) \rangle = 0 .
\]
This means that two eigenvectors are linearly independent and the minimum
dimension of the subspace that is invariant under \( \{ T(\vec{k}; R) \} \) is \( 2 f_s \). In the case,
the set of eigenvectors transforms according to an irreducible multiplier rep-
resentation of the second type under the symmetry operations. The other case
where \( \beta_\beta^* \) satisfies Eq.(4) cannot say anything about the linear independence of
two eigenvectors. If they are linearly dependent, they can differ by at most an
arbitrary phase factor and there occurs no additional degeneracy. On the other
hand, if they are linearly independent, this can be referred to as an acciden-
tal degeneracy. This is a case under the irreducible multiplier representations
called the first type.

The above discusses the criteria for establishing the type of representation by
dealing with the relation between \( \{ \vec{\bar{\tau}}(s) (\vec{k}; R) \} \) and \( \{ \tau(s) (\vec{k}; R) \} \). Before closing this
section, it may be heuristic to introduce the analogous criterion for irreducible
multiplier representations. Starting from the orthogonality theorem,
\[
\sum_{R} \chi^{(s)}_{\mu\nu}(\vec{k}; R)^* \chi^{(s)}_{\nu'\mu'}(\vec{k}; R) = (h/f_s) \delta_{\mu\nu} \delta_{\nu'\mu'} , \quad \text{first or second type,}
\]
\[
= 0 , \quad \text{third type,}
\]
where \( h \) is the order of the group \( G_0 \). Using the relation between \( \chi^{(s)}_{\mu\nu}(\vec{k}; R) \) and \( \chi^{(s)}_{\mu\nu}(\vec{k}; A_0^{-1} R A_0) \) and Eqs.(4) and (5), we arrive at another interesting criterion
\[
\sum_{R} \phi(\vec{k}; A_0 R, A_0 R)^* \chi^{(s)}_{\lambda\lambda'}(\vec{k}; A_0^{-1} R A_0 R) = +(h/f_s) \delta_{\lambda\lambda'} , \quad \text{first type,}
\]
\[
= -(h/f_s) \delta_{\lambda\lambda'} , \quad \text{second type,}
\]
\[
= 0 , \quad \text{third type.}
\]
It is often customary to express the criterion in terms of the characters $\chi^s$, i.e. $\chi^s(\vec{k}; R) = \text{Tr} \tau^s(\vec{k}; R)$,

$$\sum_R \phi(\vec{k}; A_0R, A_0R) \chi^s(\vec{k}; A_0RA_0R) = +h, \quad \text{first type},$$
$$= -h, \quad \text{second type},$$
$$= 0, \quad \text{third type}. $$

The criterion can be also expressed as a sum over the elements $A_0$ of the coset $S_\text{G}_0$ instead of $R$ of the point group $G_0$

$$\sum_{A_0} \phi(\vec{k}; A_0, A_0) \chi^s(\vec{k}; A_0) = +h, \quad \text{first type},$$
$$= -h, \quad \text{second type},$$
$$= 0, \quad \text{third type}. $$

8.5.4 Implementation in DANSE

Some results from group theory seem both useful and practical to implement in software. Others are not such good value:

- The point group of the bond can be used to fill out the force constant matrix, to check for inconsistencies of the force constants, and to help in generating them. This should, in principle, be facilitated by the matrix operations that are available as Pythonized routines in Computational Crystallographic Toolbox from the Lawrence Berkeley National Lab [?].

- The group of the wavevector can be assessed. If the Brillouin zone has been assessed previously by group theory, we can use the compatibility relationships to determine which dispersions are degenerate in energy at special points. This is important information when assessing the behavior of fuzzy data. Knowing that a convergence of curves is expected puts constraints on data, and allows a larger number of counts to be analyzed simultaneously, improving reliability.

- Time reversal symmetry might be useful to consider in magnetic systems. It seems plausible that one could assess the merging of magnon dispersions at special points, but we need to see how well this has been worked out by theorists. It would probably be too much to actually calculate character decompositions on the computer for individual cases.

- I think that actually reducing the dynamical matrix by symmetry is too much to do. The procedure may be elegant to implement, but the net gain does not seem large enough when compared to brute force diagonalization, which works rather well so far.
Chapter 9

Spin Dynamics

9.1 Spin as a Source of Magnetism

Electrons are the source of magnetism in materials. Although the atomic nuclei also have spins and magnetic moments, nuclear moments are smaller by a factor of 1,000. The electron contributes to magnetism in two separate ways. One is from orbital motion – the classical origin of magnetism. The Hamiltonian for electron motion in an electromagnetic field is

\[ H = \frac{1}{2m} \left[ \frac{\vec{p}^2}{c} + \frac{e}{c} \vec{A}(\vec{r}) \right]^2. \]  

(9.1)

Here we assume that the magnetic field \( \vec{H} \) is uniform and take the Coulomb gauge of \( \nabla \cdot \vec{A} = 0, \) consistent with \( \vec{A}(\vec{r}) = \frac{1}{2} (\vec{H} \times \vec{r}) \):

The first order term with respect to \( \vec{A}(\vec{r}) \) gives

\[ H^{(1)} = \frac{e}{2mc} \vec{H} \cdot (\vec{r} \times \vec{p}) \]  

(9.2)

\[ = \mu_B \vec{H} \cdot \vec{L}, \]  

(9.3)

where \( \vec{L} = \vec{r} \times \vec{p} \) (orbital angular momentum) and \( \mu_B = e \hbar / 2mc \) (Bohr magneton). Incidentally, the second order term in (9.1) is:

\[ H^{(2)} = \frac{e^2}{8mc^2} (\vec{r} \times \vec{H})^2, \]  

(9.4)

which is simply the magnetic induction, i.e. Lenz’s law. The first order term (9.3) gives the paramagnetism leading to the magnetization parallel to the magnetic field, while the second order term (9.4) gives the diamagnetism leading to the magnetization antiparallel to the magnetic field. For atoms with closed shells (i.e. \( \vec{L} = 0 \)) or superconducting metals, diamagnetism plays the dominant role. Except for such cases, however, paramagnetism is usually more important than diamagnetism.
There is another important contribution to magnetism from the electron besides its orbital motion. Electron spin angular momentum is the essential ingredient of magnetism in most systems having interesting magnetic properties. In metallic systems where the electron wavefunction is well approximated as a plane wave, it can be shown easily that \( \vec{L} = 0 \). The angular momentum also vanishes in insulators when the force the electron feels in a solid deviates much from the spherical symmetry the electron would feel in an isolated atom. In these cases, an orbital motion of the electron cannot contribute to the magnetism.

Electron spins can be localized or itinerant, and these two cases provide different types of magnetism. Usually, localized spins are found at ions in magnetic insulators and itinerant spins are on conduction electrons in metallic systems, although there are other important magnetic systems with both itinerant and localized spins that show unusual properties. The following sections describe the characteristic magnetic properties of localized spins, itinerant spins, localized spins immersed in itinerant spins, and strongly correlated electrons.

9.1.1 Localized Spins

Consider a system of localized spins, where the spins are arranged periodically on a crystal, and interact with each other through an exchange interaction. To understand effects of the spin-spin exchange interaction in the system, several kinds of spin models are available. These textbook models have a long history. Depending on the spin dimensionality, there are the Heisenberg model, the XY model, and the Ising model or \( n \)-state Potts model. The commonly-used Heisenberg model treats three-dimensional spins, the XY model treats two-dimensional spins, and the Ising model or \( n \)-state Potts model treats one-dimensional spins. This section discusses magnetic properties of localized spins within the ferromagnetic Heisenberg model.

For an external magnetic field \( h \hat{z} \), we have the spin interaction Hamiltonian \( (J > 0) \) on the square lattice\(^1\)

\[
\mathcal{H} = -J \sum_{\langle ij \rangle} \vec{S}_i \cdot \vec{S}_j - g\mu_B \sum_i h S^z_i ,
\]

where the first term gives the spin-spin interaction intrinsic to the Heisenberg model. Now introduce the “mean field” (or “molecular field”) approximation. Details of interactions between neighboring spins are ignored, and the Hamiltonian \( \mathcal{H} \) is replaced by:

\[
\mathcal{H} = -J \sum_{\langle ij \rangle} \vec{S}_i \cdot \langle \vec{S}_j \rangle - g\mu_B h \sum_i S^z_i .
\]

\(^1\)For geometrical reasons, on some lattices the spins cannot have long-range order.
The magnetization \( \langle M_z \rangle = -\mu_0 g \sum_i (S_i) = -\mu_0 B N \langle S_z \rangle \) can then be obtained through \( \langle S_z \rangle = \sum S_i e^{-\beta H} / Z \), where \( Z = \sum e^{-\beta H} \). The evaluation of \( \langle S_z \rangle \) is simple because \( H \) now includes only scalar quantities. Following the definition of the susceptibility \( \chi = \lim_{h \to 0} \langle M_z \rangle / h \),

\[
\chi = \frac{C}{T - T_C},
\]

(9.7)

where \( T_C \) and \( C \) are called the Curie temperature and the Curie constant, respectively. The temperature-dependence of \( \chi \) in (9.7) is characteristic of a localized spin system.

For understanding dynamical properties of a magnetic system, a starting point is its elementary excitations. For the case of three-dimensional spins in the Heisenberg model, the elementary excitation is known as a spin wave (or magnon). We consider the operator \( O_\mathbf{q} \) that creates a specific excitation in the system,

\[
[H, O_\mathbf{q}] = \omega_\mathbf{q} O_\mathbf{q}, \quad (9.8)
\]

where \( \omega_\mathbf{q} \) is the energy of the excitation. The operator \( O_\mathbf{q} \) raises the spin wave and provides the spin wave energy. Introducing the Fourier transformation of spin operators, we reexpress the Heisenberg Hamiltonian as:

\[
H = -\sum_\mathbf{q} j(\mathbf{q}) \tilde{S}(\mathbf{q}) \cdot \tilde{S}(-\mathbf{q}) \quad (9.9)
\]

\[
= -\sum_\mathbf{q} j(\mathbf{q}) \left[ S_z(\mathbf{q}) S_z(-\mathbf{q}) + \frac{1}{2} \left( S_+(\mathbf{q}) S_-(\mathbf{q}) + S_-(\mathbf{q}) S_+(\mathbf{q}) \right) \right], \quad (9.10)
\]

where we introduce \( \tilde{S}(\mathbf{q}) = \sum_i \tilde{S}_i e^{iq_i R_i} \), and \( S_\pm(\mathbf{q}) = \sum_i S_i^\pm e^{iq_i R_i} \) \((S_\pm = S_i^x \pm i S_i^y)\). Assuming the ground state is ferromagnetic, the low energy excited states should correspond to states with spins slightly deviated from perfect alignment. \( S_-(\mathbf{q}) \) is an operator that performs this role – it creates the spin wave excitation (magnon). The excitation energy of magnon is determined from (9.8):

\[
[H, S_-(-\mathbf{q})] = \omega_\mathbf{q} S_-(-\mathbf{q}), \quad (9.11)
\]

which is not, however, exactly solvable and requires an approximation. The most immediate approach is the mean field approximation for \( H \), which yields:

\[
\omega_\mathbf{q} = 2 \langle S_z \rangle [J(0) - J(\mathbf{q})] \quad (9.12)
\]

\[
\approx 2(J/N) \langle S_z \rangle a^2 \mathbf{q}^2, \quad (9.13)
\]

where for the three-dimensional cubic lattice:

\[
J(\mathbf{q}) = 2(J/N) [\cos(q_x a) + \cos(q_y a) + \cos(q_z a)]. \quad (9.14)
\]

At low temperatures, \( \langle S_z \rangle \approx NS \) and the magnon energy is

\[
\omega_\mathbf{q} = 2JSa^2 \mathbf{q}^2. \quad (9.15)
\]
The magnon energy spectrum is gapless, i.e., \( \omega \rightarrow 0 \) as \( \vec{q} \rightarrow 0 \). The zero temperature magnetization \( (M_0) \) then decreases with increasing \( T \) as magnons are created. At small \( T \):

\[
M(T) = M_0 - g\mu_B \sum_q n_q, \tag{9.16}
\]

where \( n_q \) is the Bose distribution function for magnons. A very famous result is the decrease of magnetization at low \( T \) is:

\[
\frac{M(T)}{M_0 - 1} \propto T^{3/2}, \tag{9.17}
\]

which is obtained by a simple integration of (9.16).

### 9.1.2 Itinerant Spins

Both itinerant electrons in metals and localized spins in insulators exhibit interesting magnetism. The magnetization of the system, \( \langle M_z \rangle \), in an external magnetic field \( h \) is:

\[
\langle M_z \rangle = -\mu_B \sum_{\vec{k}} \left[ \langle c^\dagger_{\vec{k} \uparrow} c_{\vec{k} \uparrow} \rangle - \langle c^\dagger_{\vec{k} \downarrow} c_{\vec{k} \downarrow} \rangle \right], \tag{9.18}
\]

where \( c^\dagger_{\vec{k} \uparrow} (c_{\vec{k} \uparrow}) \) is an electron creation (annihilation) operator with a momentum \( \vec{k} \) and spin up. In the noninteracting electron gas, the magnetic susceptibility (called the Pauli susceptibility \( \chi_P \)) is:

\[
\chi_P = 2\mu_B^2 N(\epsilon_F) \left[ 1 + O(T/\epsilon_F)^2 \right], \tag{9.19}
\]

where \( N(\epsilon_F) \) is the electron density of states at the Fermi level and \( \epsilon_F \) is the Fermi energy. \( \chi_P \) is almost constant at low temperatures and has a very weak temperature dependence.

Now consider a ferromagnetic metal. In ferromagnetic metals, the exchange interaction between electrons can raise the ferromagnetic order by overcoming the competition between kinetic energy and exchange energy. The susceptibility, \( \chi_S \), depends on the electron exchange interaction \( \mathcal{V} \), called the Stoner susceptibility. It can be obtained by extension of \( \chi_P \) as:

\[
\chi_S = \frac{\chi_P}{1 - \frac{1}{2\mu_B^2} \mathcal{V}\chi_P}. \tag{9.20}
\]

Noting that \( \chi_P \approx 2\mu_B^2 N(\epsilon_F) \) at low temperatures, \( \chi_S \) becomes:

\[
\chi_S = 2\mu_B^2 \frac{N(\epsilon_F)}{1 - \mathcal{V}N(\epsilon_F)}. \tag{9.21}
\]
9.1. SPIN AS A SOURCE OF MAGNETISM

From (9.21), \(1 - \bar{V}N(\epsilon_F) \geq 0\) is a ferromagnetic instability condition at \(T = 0\), called the “Stoner condition.” For transition metals, the \(d\)-band is narrow and \(N(\epsilon_F)\) can be large, so the Stoner condition is often satisfied (as for Fe, Co, Ni).

The temperature dependence of \(\chi_S\) is very weak at room temperature, especially compared to the Curie–Weiss susceptibility. This can be easily understood because only electrons near the Fermi level can participate in thermally-driven magnetic excitations. Nevertheless, the temperature dependence of the Stoner susceptibility provides context for a longstanding puzzle of the ferromagnetism observed in Fe \((T_C = 1044\ \text{K})\), Ni \((627\ \text{K})\), and Co \((1388\ \text{K})\). Putting \(\chi_P = 2\mu_B^2 N(\epsilon_F)[1 - aT^2]\), and \(\chi_S(T_C) = \infty\) at the transition temperature \(T_C\), we obtain for \(\chi_S(T)\):

\[
\chi_S = \frac{2\mu_B^2/\bar{V}a}{T^2 - T_C^2} = \frac{2\mu_B^2/\bar{V}a}{(T + T_C)(T - T_C)},
\]

and, in particular, near \(T_C\):

\[
\chi_S \approx \frac{\mu_B^2/\bar{V}aT_C}{T - T_C}.
\]

The Stoner susceptibility therefore shows a Curie–Weiss behavior near \(T_C\). However, it is well known that ferromagnetic metals like Fe, Ni, and Co show Curie–Weiss behavior over much wider temperature ranges. The observed Curie–Weiss behavior has been one of the most important sources of controversy over whether spins in ferromagnetic transition metals are itinerant or localized. Improving the Stoner spin susceptibility within a picture of itinerant spins remains an open problem.

Dynamical responses of spins in metals are also very different from those in localized spin systems. For conduction electrons, magnetic responses of their spins are directly related to details of the electron band structure. In the noninteracting electron gas, \(\chi_0(\vec{q}, \omega)\) governing the magnetic responses of the system is

\[
\chi_0(\vec{q}, \omega) = 2\mu_B^2 \sum_k \frac{n_{\vec{k}} - n_{\vec{k} + \vec{q}}}{\epsilon_{\vec{k} + \vec{q}} - \epsilon_{\vec{k}} - \omega},
\]

where \(n_{\vec{k}}\) is the Fermi distribution function and \(\epsilon_{\vec{k}}\) is the band energy at \(\vec{k}\). We note that the Pauli susceptibility \(\chi_P\) corresponds to \(\chi_0(\vec{q}, \omega)\) at \(\vec{q} = 0\) and \(\omega = 0\). In the same way as was found for non-interacting systems, for interacting electron systems in the paramagnetic state, \(\chi_0(\vec{q}, \omega)\) is:

\[
\chi(\vec{q}, \omega) = 2\mu_B^2 \frac{F(\vec{q}, \omega)}{1 - \bar{V}(\vec{q})F(\vec{q}, \omega)}.
\]

The susceptibility \(\chi(\vec{q}, \omega)\) is directly related to the neutron scattering cross section as:

\[
S(\vec{q}, \omega) = \frac{2}{1 - e^{-\beta\omega}} \text{Im} \chi(\vec{q}, \omega).
\]

It is curious that for temperatures well above \(T_C\), neutrons are still scattered magnetically by metallic spins. A further consideration makes this even more
curious. In thermal neutron scattering experiments, the neutron velocity $v_n$ is typically smaller than the electron velocity $v_F$ by a factor of $10^{-3}$. This means that a neutron interacts with approximately $10^3$ electrons as it moves across the magnetic moment of an atom. The average spin of $10^3$ electrons will be zero at $T > T_C$. The answer to this puzzle is provided by further analysis of (9.25). Neutrons are scattered by collective motions of electron spins (called spin-fluctuations), not by individual electron spins. More precisely, the paramagnetic scattering of neutrons is caused by fluctuations of the magnetic force on neutrons exerted by spins of itinerant electrons. On the other hand, in the ferromagnetic state ($T < T_C$), we need to consider effects from the spin polarization from the splitting of the electron band states into up and down spin states. In this case of itinerant band electrons, the elementary excitations are like the spin wave and Stoner excitation in the ferromagnetic states, obtained from the condition:

$$1 = \tilde{V}(q) \sum_k \frac{n_{\vec{k} \uparrow} - n_{\vec{k} + \vec{q} \downarrow}}{\varepsilon_{\vec{k} + \vec{q} \downarrow} - \varepsilon_{\vec{k} \uparrow} - \omega}.$$  \hspace{1cm} (9.27)

Even though the spin wave is usually discussed in the context of systems with localized spins, it is interesting that spin waves are also excited in the ferromagnetic ordered state of itinerant spins. Like the insulating case, we find $M(T)/M_0 - 1 \propto T^{3/2}$, as in (9.17) for ferromagnetic metals. This experimental finding strongly implies the existence of spin waves in metallic magnetism.

Returning to the problem of the ferromagnetic transition metals Fe, Co, and Ni, we ask again, “Are these spins itinerant?” or “Can Stoner theory properly explain their magnetism?” The answers are still extremely unclear. There have been attempts to incorporate the electron-phonon interaction within the itinerant picture. These have improved the Stoner susceptibility somewhat and have led to a Curie–Weiss-like behavior with a smaller $T_C$ than obtained from band theory, more consistent with experiment. Nevertheless, many features consistent with local moments are observed experimentally. The observed change in the specific heat of Fe at $T_C$, for example, is associated with an entropy of order $k_B \ln 2$, as expected for localized electrons. Spin-resolved photoemission experiments show that the exchange splitting does not vanish at $T > T_C$, strong experimental indication of local moments in ferromagnetic transition metals. These controversies demonstrate that a better theory is needed to properly account for electron correlations.


3A compromise model that includes both itinerant and localized features with “correlated delocalized electrons” is available, where the magnetic moments at different sites fluctuate in magnitude and direction at finite temperatures.
9.1.3 Localized Spins Embedded in Itinerant Spins

A mixed magnetic system with localized spins (or magnetic impurities) embedded in itinerant spins provides many exotic magnetic properties that differ from the properties of from systems having localized or itinerant spins, but showing features of both. Consider two cases:

- Mn atoms in Cu metal, where Mn (3d) electrons play the role of a magnetic impurity of \( S = 5/2 \), and Cu gives a sea of conduction electrons,
- An f-electron system, for example Ce which has a localized f-level and a 5d conduction band.

The governing Hamiltonian\(^4\) for both cases can be written as

\[
\mathcal{H} = t \sum_{ij} c_{i\sigma}^\dagger c_{j\sigma} + \sum_i \vec{S}_i \cdot \vec{\sigma}_i, \tag{9.28}
\]

where \( \sigma \) is the spin of conduction electrons and \( \vec{S}_i \) is the localized spin. Although both cases can be explained by \( \mathcal{H} \), they correspond to different limits of \( \mathcal{H} \).

First, for Mn impurities in Cu, we set \( t \gg J \), so the Hamiltonian expresses an effective indirect interaction of localized spins mediated by conduction electrons. This interaction is called the RKKY (Ruderman-Kittel-Kasuya-Yosida) interaction:

\[
\mathcal{H}_{\text{RKKY}} = J(k_F |\vec{r}_i - \vec{r}_j|) \vec{S}_i \cdot \vec{S}_j, \tag{9.29}
\]

and \( J(x) \) has the functional form:

\[
J(x) \propto - \frac{\cos x}{x^3} + \frac{\sin x}{x^4}. \tag{9.30}
\]

The RKKY interaction is also observed in magnetic multilayer systems composed of layers of Fe/Cu, for example, where Cu plays the role of a nonmagnetic metallic spacer. Here it is found that the exchange coupling constant oscillates and has both positive and negative values depending on the thickness of the Cu layer, consistent with the \( k_F \) for the conduction electron sea in Cu.

The second case is the opposite limit of \( t \ll J \), and approaches the Kondo model for heavy fermion systems. In the past decades, very unusual low temperature behaviors have been observed in rare earth metals (e.g., Ce) and actinides (e.g., U). The linear specific heat at low temperatures shows an unusually high coefficient, \( \gamma \), of order 1 J/mol K\(^2\), in contrast to a 1 mJ/mol K\(^2\) typical of ordinary metals. High values of \( \gamma \) are typical of “heavy fermion” systems.\(^5\) Furthermore, these heavy fermion systems have an electrical resistivity

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\(^4\)This is called the “Kondo lattice model” (KLM). Recently, the ferromagnetic KLM (\( J < 0 \)) has been frequently adopted to describe the colossal magnetoresistance in manganese oxides, where \( J \) is the Hund coupling.

\(^5\)Values of \( \gamma \) for UPt\(_3\), CeAl\(_3\), CeCu\(_2\)Si\(_2\), UBe\(_{13}\) are 0.45 J/mol K\(^2\), 1.6 J/mol K\(^2\), 1.1 J/mol K\(^2\), and 1.1 J/mol K\(^2\), respectively.
of $\rho_0 + AT^2$ at low $T$ with huge values of $A$ of order $10 \mu \Omega \text{cmK}^{-2}$, whereas $A$ is of order $10^{-5} \mu \Omega \text{cmK}^{-2}$ or less for ordinary metals. Heavy fermion systems exhibit another basic and universal magnetic property. Below a characteristic temperature $T^*$, heavy fermion systems show Fermi liquid behavior with a huge effective mass, and then constant (but very high) Pauli susceptibility. Above $T^*$, they show a Curie–Weiss susceptibility that originates from the localized $f$ electrons. At $T^*$, the quasi-particles are screened by conduction electrons and a singlet (nonmagnetic) state is formed, owing to strong electron correlations. Although heavy fermion systems show differences in their properties, the disappearance of a magnetic moment is a common feature.

Another unusual property originating with magnetic scattering is the Kondo effect. The Kondo effect is a logarithmic increase of the electrical resistivity of the Kondo system when the temperature is reduced. It is caused by spin flip scatterings of conduction electrons at magnetic impurities.

### 9.1.4 Strongly Correlated Electrons

Since the discovery of high $T_c$ superconductors, enormous theoretical effort has been made to understand the two-dimensional Hubbard model, which was originally introduced to understand the metal-insulator transition in transition metals. The Hubbard Hamiltonian is:

$$\mathcal{H} = t \sum_{ij} c_{i\sigma}^\dagger c_{j\sigma} + U \sum_i n_i^\uparrow n_i^\downarrow.$$  \hfill (9.31)

In high $T_c$ superconductors, $c_{i\sigma}^\dagger (c_{i\sigma})$ is the creation (annihilation) operator of the electron (or the hole) at the highest antibonding orbital (predominantly $d_{x^2-y^2}$) of Cu$^{2+}$ positioned at $i$ in CuO$_2$ plane. Approximately, $t \sim 0.1$ eV and $U \sim 1$ eV. It is immediately clear that the energy scale of the Coulomb correlation energy, $U$, is too high compared to the interesting energy scales, which are around 10 meV for critical temperature of 100 K. The low energy excitation relevant to superconductivity is therefore believed to originate with spins, not with charge, and effort has been made to derive an effective spin Hamiltonian that depends on the hole concentration (doping).

In the undoped case (half-filling), there is one electron at each site and the hopping into the nearest neighbor costs the energy $U$. In the limit of $t \ll U$, the electrons look localized at each site because of the high energy barrier $U$. The Hubbard model can be transformed into the Heisenberg spin Hamiltonian:

$$\mathcal{H} = J \sum_{ij} \vec{S}_i \cdot \vec{S}_j,$$  \hfill (9.32)

$$J = 4t^2/U.$$  \hfill (9.33)

$^6T^*$ is not necessarily same as $T_K$, which is the usual Kondo temperature for a single Kondo ion, and in most cases we see $T^* < T_K$ owing to the condensation of Kondo ions.

$^7$Examples of some high $T_c$ materials are La$_{2-x}$Sr$_x$CuO$_4$ ($T_c \sim 40$ K), YBa$_2$Cu$_3$O$_7$ ($T_c = 92$ K), Bi$_2$Sr$_2$Ca$_2$Cu$_3$O$_{10}$ ($T_c = 110$ K), and Tl$_2$Sr$_2$Ca$_2$Cu$_3$O$_{10}$ ($T_c \sim 125$ K).
In this limit of \( t \ll U \), the system acts as an antiferromagnetic insulator. By introducing a small fraction, \( x \), of holes, the superconducting phase appears. Typical fractions for \( \text{Bi}_2\text{Sr}_2\text{CaCu}_2\text{O}_{6+x} \) (Bi2212) are \( x \sim 0.1 \) (this may be underdoped, but overdoping occurs for \( x \sim 0.2 \)). The effective Hamiltonian for a small number of holes has also been reduced from the Hubbard model, and is called the \( t-J \) model:

\[
H_{t-J} = t \sum_{ij} a_{i\sigma}^\dagger a_{j\sigma} + J \sum_{ij} \vec{S}_i \cdot \vec{S}_j ,
\]

(9.34)

where \( a_{i\sigma}^\dagger = c_{i\sigma}^\dagger (1 - n_{i\sigma}) \) and \( a_{i\sigma} = c_{i\sigma} (1 - n_{i\sigma}) \). In the limit of half-filling, \( H_{t-J} \) is the Hamiltonian of a Heisenberg antiferromagnet. Research interests are in small deviations from half-filling, where holes move in the antiferromagnetic lattice and counteract antiferromagnetic long-range order. This leads to antiferromagnetic spin-fluctuations peaked at a momentum near \( \vec{Q} = (\pi, \pi) \). In the last decades, spin fluctuations have been found to play fundamental roles in high \( T_c \) superconductors. In the antiferromagnetic Fermi liquid model, many key properties of superconducting cuprates have been understood by a strong interaction between quasi-particles and spin-fluctuations. For example, the spin-fluctuation model has been successful in explaining transport properties such as electrical resistivity (\( \rho \propto T \)), the magnitude of \( T_c \), results from nuclear magnetic resonance (NMR) experiments, angle-resolved photoemission spectroscopy (ARPES) experiments, and other experimental results. The detailed phenomenological propagator for spin-fluctuations is:

\[
\chi_{sf}(\vec{q}, \omega) = \frac{\chi_{Q}}{1 + \xi^2 (\vec{q} - \vec{Q})^2 - i \omega / \omega_{sf}} ,
\]

(9.35)

which has proved applicable for interpreting NMR measurements and neutron scattering experiments. Here \( \omega_{sf} \) is the spin-fluctuation energy and \( \xi \) is the antiferromagnetic correlation length. Through the value of \( \xi \), the spin-fluctuation depends on the hole concentration. In the limit of \( \xi \rightarrow \infty \), the system in the pure magnon region, i.e. the undoped antiferromagnetic insulator with long-range order. An interaction between quasi-particles and spin-fluctuations is especially strong in underdoped materials. It has been shown recently that strong anisotropies in ARPES data for underdoped Bi2212 originate with a strong coupling between quasi-particles and spin-fluctuations. Incidentally, for other superconducting materials, i.e., \( \text{La}_{2-x}\text{Sr}_x\text{CuO}_4 \) and \( \text{YBa}_2\text{Cu}_3\text{O}_{7-x} \), data from inelastic neutron scattering and measurements of the spin-lattice relaxation rate by NMR indicate that the spin-fluctuations induce an opening of the gap in the spin excitation\(^8\) in the CuO\(_2\) plane, with an energy comparable to the BCS gap. The apparent gap develops well above \( T_c \).

\(^8\)The spin gap corresponds to the pseudogap of a small energy scale.
9.2 Simulations of Spin Dynamics

9.2.1 Monte Carlo Method

Section 9 introduced different spin systems and their excitations. Several approaches are available for calculating the states and state evolution of these spin systems. Perhaps the most conventional approach, although not a simulation per se, is the diagrammatic expansion of the Green’s function. This formal approach is not practical for many complex systems, however. Instead, one may use the local density approximation (LDA), or LDA+U to study band magnetism, or the small-size exact diagonalization for more localized spin systems. However, for more complete descriptions of the dynamics of spins, the most suitable and reliable method is often a Monte Carlo simulation. There are many variants of Monte Carlo simulations developed and optimized for specific problems. This section explains the Monte Carlo approach for calculating the states and dynamics of classical lattice spin systems. Monte Carlo simulations of classical systems are easier to understand than quantum Monte Carlo simulations, which are mentioned at the end of this section.

A Monte Carlo simulation is a Markovian process. Such processes will reach a steady state of a system that is independent of the initial configuration. Unfortunately, especially at low temperatures, this final state of equilibrium may require a very long time to achieve. If one knows the ground state a priori, at low temperatures it may be appropriate to start with the system in a ground state configuration, and allow temperature to produce disorder in this configuration. The ground state is often unknown, however, and this is typical of more complicated systems. It may be possible to start with several candidate ground state configurations, and identify the true ground state structure as the one that does not evolve with time.

An alternative approach is sometimes called “simulated annealing,” where the simulation begins with the system in a fully random state characteristic of infinite temperature. Equilibrium at lower temperatures is achieved by gradually reducing the temperature, and allowing the system to relax under the spin-spin interactions. Some delicacy is required for balancing the slowness of cooling with the need to minimize the time of the simulation.

Finally, systems that undergo spontaneous symmetry breaking may pose special problems. Starting from the random spin configuration at $T = \infty$ in the isotropic ferromagnet, one may find that the ordering direction would rotate without any preferred direction even below $T_C$. In such a case, therefore, one may include an infinitesimal anisotropy in the Hamiltonian to induce the symmetry breaking, or one may start with a symmetry-broken ground state configuration at $T = 0$ and raising the temperature. Without an applied bias, the formation of local domains, each with its own direction of spin alignment, is a common feature. Elimination of the domain boundaries is favorable ener-

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\[ \text{For example, by initially aligning all the spins along a certain direction.} \]
getically, but may take a long time in practice.

9.2.2 Spin Updates in Monte Carlo Simulations

Metropolis Algorithm

In the Metropolis algorithm, a new configuration is generated from an existing one by using a transition probability that depends on the energy difference between the two configurations. The state of thermodynamic equilibrium satisfies the detailed balance between two states $n$ and $m$,

$$P_n W_{n\rightarrow m} = P_m W_{m\rightarrow n}, \quad (9.36)$$

where $P_n$ is the probability of the system being in the state $n$ and $W_{n\rightarrow m}$ is the transition rate from $n \rightarrow m$. The Metropolis algorithm selects the simplest choice of the transition rate that is consistent with detailed balance

$$W_{n\rightarrow m} = \begin{cases} \exp(-\Delta E/k_B T) & \Delta E > 0 \\ 1 & \Delta E < 0 \end{cases}, \quad (9.37)$$

where $\Delta E = E_n - E_m$. The Metropolis algorithm updates one spin at a time in a given configuration at temperature $T$:

1. select the spin at site $i$,
2. evaluate $\Delta E$ by updating the spin at $i$,
3. generate a random number $\eta$, where $0 < \eta < 1$,
4. accept the updated configuration if $\eta < e^{-\Delta E/k_B T}$, or reject otherwise,
5. return to step 1 for a different spin at site $i + 1$.

For the model of continuous spins of $|S_i| = 1$, oriented in three dimensions, one may update the spin at site $i$ by generating two random numbers $\eta_1$ and $\eta_2$, such that $\zeta^2 = \zeta_1^2 + \zeta_2^2 < 1$ such that $\zeta_1 = 1 - \eta_1$ and $\zeta_2 = 1 - \eta_2$ (where $0 < \eta_1, \eta_2 < 1$)

$$S_x = 2\zeta_1 \sqrt{1 - \zeta^2}, \quad (9.38)$$
$$S_y = 2\zeta_2 \sqrt{1 - \zeta^2}, \quad (9.39)$$
$$S_z = 1 - 2\zeta^2. \quad (9.40)$$

The Metropolis algorithm ensures that the steady state of the system is the actual state of thermodynamic equilibrium. This is proved by assuming that the system is in thermodynamic equilibrium, and then showing that the Metropolis algorithm has the transition rates needed to keep it there. In thermodynamic
equilibrium, the probability \( P_n \) is the Boltzmann factor normalized by the partition function, \( e^{-E_n/k_B T}/Z \), a central result of statistical physics. Substituting into (9.36)

\[
e^{-E_n/k_B T} W_{n \rightarrow m} = e^{-E_m/k_B T} W_{m \rightarrow n},
\]

(9.41)

\[
e^{-\left(E_n - E_m\right)/k_B T} = \frac{W_{m \rightarrow n}}{W_{n \rightarrow m}}
\]

(9.42)

Note that \( W_{n \rightarrow m} = 1 \) (since the transition \( n \rightarrow m \) is downhill energetically, it will always occur according to (9.37)). Equation (9.42) becomes:

\[
W_{m \rightarrow n} = e^{-\left(\Delta E\right)/k_B T},
\]

(9.43)

which is the rate used by the Metropolis algorithm in step 4 or (9.37).

All Markovian processes converge to a steady state\(^{10}\), and the Metropolis algorithm assures us that thermodynamic equilibrium will be achieved between all pairs of spins in the system. It does not ensure that equilibration will occur in a reasonable time, however.

### 9.2.3 Low Temperatures

At low temperatures, a more sophisticated update algorithm is necessary. Most random updates cause a large energy exponent \( \Delta E/k_B T \), so the Metropolis algorithm will reject most changes to the spin configuration. The equilibration procedure can then become far too slow to be practical. To speed things up, the randomly-selected changes in spin can be made smaller, for example. This can be done by attenuating the random changes of spin by a factor \( \delta (0 \leq \delta \leq 1) \), i.e. \( \Delta S \) should be replaced by \( \delta \Delta S \). The factor \( \delta \) can be adjusted so that the acceptance rate is around 50% on the average.

An actual implementation could be based on a parameterized temperature-dependent solid angle, \( \Omega(T) \), of spin \( i \) with respect to its original orientation \( S_i \). This is begun by defining \( \alpha = \tan^{-1} \frac{S^y_i}{S^x_i} \) and \( \beta = \cos^{-1} \frac{S^z_i} \), from which we define the rotation matrices \( R_x(\alpha) \) and \( R_y(\beta) \):

\[
R_x(\alpha) = \begin{pmatrix}
\cos \alpha & \sin \alpha & 0 \\
-\sin \alpha & \cos \alpha & 0 \\
0 & 0 & 1
\end{pmatrix},
\]

(9.44)

\[
R_y(\beta) = \begin{pmatrix}
\cos \beta & 0 & -\sin \beta \\
0 & 1 & 0 \\
\sin \beta & 0 & \cos \beta
\end{pmatrix}.
\]

(9.45)

The updated orientation of spin \( i \), \( S'_i \) is

\[
S'_i = R^{-1}_x(\alpha) R^{-1}_y(\beta) S''_i,
\]

(9.46)

\(^{10}\)Or a cyclic state in anomalous cases.
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where

\[ S_i'' = (\sin \theta \cos \phi, \sin \theta \sin \phi, \cos \theta) , \]  
\[ \theta = \Omega(T) \eta_1 , \]  
\[ \phi = 2\pi \eta_2 . \]  
\[ (9.47) \]
\[ (9.48) \]
\[ (9.49) \]

A useful form\(^\text{11}\) of \( \Omega(T) \) may be

\[ \Omega(T) = \pi \tanh(\xi T) . \]  
\[ (9.50) \]

With this form of \( \Omega(T) \), we can adjust \( \xi \) so that the acceptance rate is around 50%. Note the two extreme limits: \( \Omega(T) \to 0 \) as \( T \to 0 \) and \( \Omega(T) \to \pi \) as \( T \to \infty \).

Overrelaxation Technique

The overrelaxation technique is used in combination with the Metropolis algorithm for improving the rate of relaxation to the equilibrium configuration, especially at low temperatures. Let us assume that we are treating a system of isotropic continuous spins in the Heisenberg model, with the Hamiltonian

\[ \mathcal{H} = \sum_{ij} J_{ij} S_i \cdot S_j . \]  

In the overrelaxation method, a spin is precessed about the full interaction field, which it can do without any change of energy. When the angle of precession, \( \theta \), is as large as \( \pi \), this alteration of the spin structure can promote quicker changes in the orientation of other spins during subsequent Monte Carlo steps.

At the site \( i \), the full interaction field for \( S_i \) is \( S_{nn} \equiv J_{ij} S_j \). The overrelaxed spin, \( S_i' \), is evaluated by successive rotations using rotations by Euler angles. One needs two angles of \( \alpha \) and \( \beta \) to define the direction of \( S_{nn} \):

\[ \alpha = \tan^{-1} \frac{S_{nn}^y}{S_{nn}^x} , \]  
\[ \beta = \cos^{-1} \frac{S_{nn}^z}{|S_{nn}|} , \]  
\[ S_i' = R_z^{-1}(\alpha) R_y^{-1}(\beta) R_z(\pi) R_y(\beta) R_z(\alpha) S_i . \]  
\[ (9.51) \]
\[ (9.52) \]
\[ (9.53) \]

In an actual simulation, a single “hybrid” Monte Carlo update of the spin configuration could include a Metropolis sweep and two overrelaxation sweeps.

Equilibration (Thermalization)

Repeating the hybrid Monte Carlo steps, spins are updated by sweeping the whole spin lattice, eventually reaching the equilibrium configuration at a given \( T \). It is an important problem to minimize the number of Monte Carlo steps needed to obtain the fully relaxed equilibrium state. The optimization differs, however, depending on the specific problem or the simulation conditions such as temperature and lattice size, for example. It is essential to perform calculations of simple quantities like susceptibilities or magnetization, and test if

\[^{11}\text{The form of } \Omega(T) \text{ given here is just the simplest example. One may introduce a more complicated form to give a more desirable performance.}\]
the system is in the equilibrium state. In more advanced validations, one may check the autocorrelation time. Substantial gains in efficiency are often found for hybrid Monte Carlo simulations. For the square lattice Heisenberg model, the nonhybrid Monte Carlo method typically requires at least $O(10^4)$ steps at the temperature range of $O(0.1)$, whereas the hybrid method requires $O(10^3)$ steps.

### 9.2.4 Time Evolution of Spins

For a system of spins in the Heisenberg model, the equation of motion for the spin degrees of freedom is

$$\frac{\partial S_i}{\partial \tau} = -S_i \times \frac{\partial H}{\partial S_i} = -JS_i \times \sum_{j \neq i} S_j . \quad (9.54)$$

The $N$ spins are coupled to each other, and the time-evolution of the spins is obtained by solving the coupled differential equations. There are many integration algorithms to solve the differential equations. Here we introduce the Suzuki-Trotter decomposition, which is especially suitable for the lattice problem. Taking the example of the simple square lattice with the nearest-neighbor spin coupling, one can divide the lattice into two sublattices $A$ and $B$ by checkerboard decomposition. Employing the Suzuki-Trotter decomposition up to $O(d\tau^5)$,

$$|S_i(\tau + d\tau)| = e^{(A+B)\tau} |S_i(\tau)| , \quad (9.55)$$

and $e^{(A+B)\tau}$ is decomposed using

$$e^{(A+B)\tau} = \prod_{i=1}^5 e^{p_i A \tau/2} e^{p_i B \tau/2} e^{p_i A \tau} + O(d\tau^5) , \quad (9.56)$$

with

$$p_1 = p_2 = p_4 = p_5 = p = 1/(4 - 4^{1/3}) \quad (9.57)$$

and

$$p_3 = 1 - 4p . \quad (9.58)$$

Here $A$ and $B$ are the rotation generators of the sublattices $A$ and $B$, with fixed $\{S_{iB}\}$ and $\{S_{iA}\}$, respectively.

The following algorithmic explanation may be more clear.

$e^{A\tau} |S_i| :$ time evolution of sublattice $A$

for $i = 1$ to $N$

when $i \in A$

$$S_{nn} = \sum_{j \neq i \in B} S_j$$

The Suzuki-Trotter decomposition is particularly useful for lattice problems, allowing efficient simulation of systems with spin interactions. The decomposition is applied up to $O(d\tau^5)$, which provides a balance between accuracy and computational cost.
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\[ e^{A\delta \tau} [S_i] = \left[ (S_{nn} \cdot S_i) S_{nn} / |S_{nn}|^2 \right] \left( 1 - \cos(|S_{nn}| \delta \tau) \right) + S_i \cos(|S_{nn}| \delta \tau) + \left( |S_{nn} \times S_i| / |S_{nn}| \right) \sin(|S_{nn}| \delta \tau) \]

\[ e^{B\delta \tau} [S_i] : \text{time evolution of sublattice } B \]

for \( i = 1 \) to \( N \)

when \( i \in B \)

\[ S_{nn} = \sum_{j \neq i \in A} S_j \]

\[ e^{B\delta \tau} [S_i] = \left[ (S_{nn} \cdot S_i) S_{nn} / |S_{nn}|^2 \right] \left( 1 - \cos(|S_{nn}| \delta \tau) \right) + S_i \cos(|S_{nn}| \delta \tau) + \left( |S_{nn} \times S_i| / |S_{nn}| \right) \sin(|S_{nn}| \delta \tau) \]

In this algorithm, the time evolution of spins is performed by combinations of \( e^{-A\delta \tau} \) and \( e^{-B\delta \tau} \), just as given by the Suzuki-Trotter decomposition.

9.2.5 Observables

Inelastic neutron scattering, especially experiments on single crystal samples, can probe directly the spin-spin correlation function \( S(q, \omega) \) that describes the dynamics of spins

\[ S(q, \omega) = \frac{1}{2\pi N^2} \sum_{ij} e^{i q \cdot (r_i - r_j)} \int_0^{\tau_{\text{max}}} e^{-i \omega \tau} \langle S_i(\tau) \cdot S_j(0) \rangle \, d\tau. \tag{9.59} \]

This \( S(q, \omega) \) can be calculated by Monte Carlo simulation. \( S(q, \omega) \) is obtained by a simple Fourier transformation of the time-dependent correlation function

\[ C_i(\tau) = \langle S_i(\tau) \cdot S_j(0) \rangle, \tag{9.60} \]

which is readily evaluated in a Monte Carlo simulation. Pole structures of \( S(q, \omega) \) and \( \omega(q) \), can provide information on magnetic excitations and relaxations, which are fundamental to understanding the spin systems. Incidentally, the quasi-elastic response \( S(q, 0) \) tells us the kinds of magnetic fluctuations (correlations) that dominate at a given \( T \). For instance, if \( S(Q, 0) \gg S(0, 0) \) with say \( Q = (\pi, \pi) \), we conclude that antiferromagnetic exchange interactions are dominant over ferromagnetic ones at the given \( T \).

For powder or polycrystal samples, the measured spectra are in the angle-integrated form

\[ S(q, \omega) = \frac{2}{N^2} \sum_{ij} \frac{\sin(|q||r_i - r_j|)}{|q||r_i - r_j|} \int_0^{\tau_{\text{max}}} e^{-i \omega \tau} \langle S_i(\tau) \cdot S_j(0) \rangle \, d\tau. \tag{9.61} \]

In some experiments, or in some stages of data analysis, scattered neutrons with a rather wide distribution of momenta are collected into a single energy bin. For such data, the resulting spectra \( S(\omega) \) probes the local responses of spins
as
\[
S(\omega) = \int S(q, \omega) \, dq = \frac{1}{2\pi N} \sum_i \int_{0}^{\tau_{\text{max}}} e^{i\omega\tau} \langle S_i(\tau) \cdot S_i(0) \rangle \, d\tau.
\] (9.62)

The quasi-elastic local response, \(S(0)\), is also an interesting quantity. It is directly related to the spin-lattice relaxation rate \(T_1\) of a local probe as in nuclear magnetic resonance (NMR) or Mössbauer spectrometry.

### 9.2.6 Comments on Quantum Monte Carlo Simulations

An essential difference between classical and quantum Monte Carlo simulations is in how the spin configuration is updated, that is, how the Metropolis algorithm is implemented. The classical Monte Carlo method is easier in that the equilibrium probability has the proportionality
\[
P(|\{S_i\}|) \propto e^{-\beta H(|\{S_i\}|)}
\] (9.63)

for the particular configuration \(|\{S_i\}|\), that is: \(H(|\{S_i\}|) = E(|\{S_i\}|)\). On the other hand, for the quantum Heisenberg model, for which
\[
H = H_0 + V
\] (9.64)

with
\[
H_0 = \sum_{ij} S_i^z S_j^z, \quad V = \sum_{ij} (S_i^x S_j^x + S_i^y S_j^y),
\] (9.65) (9.66)

\[
[H_0, V] \neq 0,
\] (9.67)

we apply the Trotter formula\(^{12}\)
\[
e^{-\beta H} \approx \left[ e^{-\beta H_0/m} e^{-\beta V/m} \right]^m, \quad Z = \sum_{\alpha} \langle \alpha | e^{-\beta H_0} | \alpha \rangle, \quad Z = \sum_{\alpha} \prod_{k=1}^{m} \langle \alpha_k | e^{-\beta V/m} | \alpha_k' \rangle \langle \alpha_k' | e^{-\beta V/m} | \alpha_{k+1} \rangle,
\] (9.68) (9.69) (9.70)

where \(|\alpha\rangle\) can be the eigenstate of \(H_0\) and \(|\alpha_1\rangle = |\alpha_{m+1}\rangle\). Then the matrix element \(\langle \alpha | e^{-\beta V/m} | \alpha' \rangle\) is evaluated classically, leading to \(e^{-\beta V(\alpha,\alpha')/m}\). A

\(^{12}\)cf. Suzuki-Trotter decomposition for the time evolution integrator
9.3. INTERACTIONS BETWEEN THERMAL EXCITATIONS OF ELECTRONS AND PHONONS

A $d$-dimensional quantum spin problem therefore always corresponds to an effective $(d+1)$-dimensional problem.\textsuperscript{13}

Besides the quantum (localized) spin model, another important problem is that of electrons with itineracy. One may introduce the Hubbard model as the simplest example,

$$\mathcal{H} = t \sum_{ij} \sum_\sigma c^\dagger_{i\sigma} c_{j\sigma} + U \sum_i n_{i\uparrow} n_{i\downarrow}. \quad (9.71)$$

For this problem, we need to apply another kind of quantum Monte Carlo method that incorporates the path integral formalism. In this method, the itinerant degrees of freedom of electrons are completely integrated out by the path integral formalism\textsuperscript{14} and the remaining problem is then cast as an Ising spin problem in $(d+1)$-dimensions.

9.3 Interactions between Thermal Excitations of Electrons and Phonons

9.3.1 Formalism of Phonon-Phonon Interactions

The full Hamiltonian of the crystal, $H$, is

$$H = H_n + H_e + H_{ep} \quad (9.72)$$

where $H_e$ and $H_{ep}$ are the contributions from the electron energy (including electron-electron interactions and thermal electronic excitations), and from electron-phonon interactions (EPI). To understand phonon-phonon interactions, we consider in detail the Hamiltonian for nuclear motions $H_n$, extending the Taylor series of Eq. 8.49

$$H_n = \Phi_0 + \frac{p^2}{2m} + \frac{1}{2!} \sum_{\kappa \alpha \kappa' \alpha'} \Phi_{\kappa \alpha \kappa' \alpha'} u_{\kappa \alpha} u_{\kappa' \alpha'}$$

$$+ \frac{1}{3!} \sum_{\kappa \alpha \kappa' \alpha' \kappa'' \alpha''} \Phi_{\kappa \alpha \kappa' \alpha' \kappa'' \alpha''} u_{\kappa \alpha} u_{\kappa' \alpha'} u_{\kappa'' \alpha''}$$

$$+ \frac{1}{4!} \sum_{\kappa \alpha \kappa' \alpha' \kappa'' \alpha'' \kappa''' \alpha'''} \Phi_{\kappa \alpha \kappa' \alpha' \kappa'' \alpha'' \kappa''' \alpha'''} u_{\kappa \alpha} u_{\kappa' \alpha'} u_{\kappa'' \alpha''} u_{\kappa''' \alpha'''}) \quad (9.73)$$

where we have simplified the development to one atom per unit cell, so $u_{\kappa \alpha}$ is the displacement of atom $\kappa$ along Cartesian direction $\alpha$.

\textsuperscript{13}One additional dimension comes from the Trotter decomposition.

\textsuperscript{14}The basic idea is to transform the Hubbard model into a quadratic form by introducing the additional Ising-type bosonic field through the Trotter decomposition and the Hubbard-Stratonovich transformation. Fields of the quadratic action can be always integrated out, that is, $\text{Tr} [ e^{\sum_{ij} \lambda_{ij} c^\dagger_{i\sigma} c_{j\sigma}} ] = \det (1 + e^{-A})$, where $A$ carries the Ising-type auxiliary field.
The transformation to normal coordinates for an infinite periodic crystal is

$$\vec{U}_{\vec{k}} = \frac{1}{\sqrt{N}} \sum_{\vec{r}_e} \vec{u}_{\vec{k}} e^{-i\vec{k} \cdot \vec{r}_e}, \quad (9.74)$$

and by Fourier inversion

$$\vec{u}_{\vec{k}} = \frac{1}{\sqrt{N}} \sum_{\vec{r}_e} \vec{U}_{\vec{k}} e^{i\vec{k} \cdot \vec{r}_e}. \quad (9.75)$$

Substituting Eq. 9.75 into Eq. 9.73, we identify the Fourier transform of $\Phi_{\alpha\alpha'}\kappa\kappa'$, and use it to define the dynamical matrix of Eqs. 8.58 and 8.59. Our interest now is in the higher-order terms of Eq. 9.73, which cause anharmonic behavior when the atoms are vibrating with substantial amplitude. Using Eqs. 9.73 - 9.75 we can assess phonon-phonon interactions with a classical approach, which is valid at high temperatures.

There is, however, a clarity to be gained in working with the quantized phonon field (a displacement field in the crystal), especially when considering excitations of nonlocal electron states simultaneously with phonon excitations. The phonon field operator is

$$A_{\vec{k}} = a_{\vec{k}} + a^+_{-\vec{k}} = A^+_{-\vec{k}}. \quad (9.76)$$

The raising and lowering operators $a^{\dagger}_{\vec{k}}$ and $a_{\vec{k}}$ create and annihilate the phonon $\vec{k}_i$. They are constructed from the momentum and position operators of the Hamiltonian when a plane wave solution is considered for one $\vec{k}_i$. The reader should have references showing the properties of $a^\dagger$ and $a$ for a simple harmonic oscillator (40), and the $a^{\dagger}_{\vec{k}}$ and $a_{\vec{k}}$ work the same way for a single normal mode of a harmonic crystal. The formalism is now in the “occupation number representation,” where $n$ phonons of wavevector $\vec{k}$ are created with $n$ raising operations as $\left(a^\dagger_{\vec{k}}\right)^n |0\rangle_{\vec{k}} = (n!)^{-1/2} |n\rangle_{\vec{k}}$. The phonon field operators are related to the displacements (now operators) as Fourier coefficients

$$\vec{u}(\vec{r}_e) = \sum_{\vec{k}} \sqrt{\frac{\hbar}{2Nm\omega_{\vec{k}}}} \vec{c}(\vec{k}) e^{i\vec{k} \cdot \vec{r}_e} A_{\vec{k}}. \quad (9.77)$$

Note the similar roles of the phonon field operator $A_{\vec{k}}$ in Eq. 9.77 and the vector $\vec{U}_{\vec{k}}$ in Eq. 9.75.

By substituting Eq. 9.77 into Eq. 9.73, the dynamical matrix is recovered from the quadratic term, with prefactors such as $\hbar/(4\omega_{\vec{k}}\omega_{\vec{k}'}^\prime)$. We define simi-
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larly the transformed higher-order terms, so Eq. 9.73 becomes

\[
\hat{H}_n = \Phi_0 + \sum_{\mathbf{k}_i} \hbar \omega_{\mathbf{k}_i} \left( a_{\mathbf{k}_i}^\dagger a_{\mathbf{k}_i} + \frac{1}{2} \right) + \sum_{\mathbf{k}_i} \sum_{\mathbf{k}_j} V(\mathbf{k}_i, \mathbf{k}_j) A_{\mathbf{k}_i}^\dagger A_{\mathbf{k}_j}^\dagger A_{\mathbf{k}_j} A_{\mathbf{k}_i} + \sum_{\mathbf{k}_i} \sum_{\mathbf{k}_j} \sum_{\mathbf{k}_k} V(\mathbf{k}_i, \mathbf{k}_j, \mathbf{k}_k) A_{\mathbf{k}_i}^\dagger A_{\mathbf{k}_j}^\dagger A_{\mathbf{k}_j} A_{\mathbf{k}_k}^\dagger A_{\mathbf{k}_i} + ... ,
\]

(9.78)

where the first sum on the right-hand side includes both the kinetic energy and the harmonic part of the potential energy (following standard development (41)), and the phonon number operator is \( a_{\mathbf{k}_i}^\dagger a_{\mathbf{k}_i} \). The \( V \) are related to the \( \Phi \) as Fourier transforms. For example,

\[
V(\mathbf{k}_i, \mathbf{k}_j, \mathbf{k}_k) = \frac{1}{3!} \sqrt{\frac{1}{N m^3}} \left( \frac{\hbar}{2} \right)^3 \frac{1}{\omega_{\mathbf{k}_i} \omega_{\mathbf{k}_j} \omega_{\mathbf{k}_k}} \delta(\mathbf{k}_i + \mathbf{k}_j + \mathbf{k}_k - \mathbf{g}) \times \sum_{\mathbf{r}_\kappa} e^{i(\mathbf{k}_i + \mathbf{r}_\kappa) \mathbf{r}} e^{i(\mathbf{k}_j + \mathbf{r}_\kappa) \mathbf{r}} e^{i(\mathbf{k}_k + \mathbf{r}_\kappa) \mathbf{r}} \Phi_{a,\alpha,\alpha',\alpha''},
\]

(9.79)

where the factor \( \delta \) signifies that the sum \( \mathbf{k}_i + \mathbf{k}_j + \mathbf{k}_k \) equals a reciprocal lattice vector, \( \mathbf{g} \). For terminology, the terms with \( V(\mathbf{k}_i, \mathbf{k}_j, \mathbf{k}_k) \) and \( V(\mathbf{k}_i, \mathbf{k}_j, \mathbf{k}_k, \mathbf{k}_l) \) in Eq. 9.78 give the energies of “phonon-phonon interactions” because they alter the phonon energies when larger vibrational displacements are present in the crystal.

Details of the phonon dispersions are immediately relevant when attempting to calculate anharmonic behavior, and several concepts have been developed in detail. For example, if the phonon dispersion relation \( \varepsilon(\mathbf{k}) \) were linear, it would be easy to add two wavevectors \( \mathbf{k}_i + \mathbf{k}_j = \mathbf{k}_l \), and simultaneously conserve energy \( \varepsilon + \varepsilon' = \varepsilon'' \). The number of three-phonon processes increases as \( k^2 \), at least for small \( k \). Many phonon dispersions \( \varepsilon(\mathbf{k}) \) are concave downwards, however, so simultaneous energy and momentum conservation is often impossible unless at least one phonon lies on a different branch. Phonon processes that satisfy momentum and energy conservation depend on the symmetry of the dispersion relations, and on the crystal structure. For larger phonon wavevectors \( \mathbf{k} \), momentum conservation is possible by adding a reciprocal lattice vector. The idea is that the momentum transferred to the entire crystal occurs with zero energy because of the large mass of the crystal. Such “umklapp” processes allow many more three-phonon interactions, but the phonon wavevectors must be of length comparable to the reciprocal lattice vector if the vector additions are possible.

An important question is how many terms are needed in Eq. 9.78 to account accurately for anharmonic behavior. We expect the higher-order terms in Eq.
9.73 to be progressively smaller, but the requirement of energy conservation and the factor $\delta$ for momentum conservation in Eq. 9.79 restricts the allowable three-phonon processes, owing to restrictions imposed by the phonon dispersions $\varepsilon(k)$. Fourth-order anharmonicity can therefore play a role as significant as the third. Another complexity is that four-phonon processes can be generated from three-phonon processes. The idea is that two three-phonon processes such as $k + k' \rightarrow k''$ and $k'' \rightarrow k''' + k''''$ can be combined through intermediate states $[k'']$ to become a four-phonon process in second-order perturbation theory, which involves matrix elements as $\sum_{k'''} \langle k''', k'''' | H'''' | k''', k'''' \rangle$. With effort, this field theoretic approach has been used for some physical predictions as described below. Approximate summations of the perturbation series have been performed, and some results for anharmonic heat capacity were reported for solids with Lennard-Jones potentials (42). This approach also has conceptual value for understanding the meaning of anharmonicity.

### 9.3.2 Formalism of Electron-Phonon Interactions

The large difference in energy scales of phonons and electrons can motivate the separation of the Hamiltonian of the solid into a term with nuclear coordinates for the phonons $H_n$, and a term with electron coordinates $H_e$ (Eq. 9.72). The energy of the crystal deformation caused by a phonon originates with the electrons, of course, but although this potential energy of deformation is electronic in origin, it transfers to kinetic energy in the motion of the nuclei.\(^{15}\) If the electrons were always in their ground states, all this energy of deformation would be associated with $H_n$ alone (i.e., the phonons). Treating the electron system and the phonon system as two independent thermodynamic systems becomes inconsistent at finite temperature, however, because the presence of phonons alters the thermal excitations of electrons.

Very generally, the electron-phonon interaction (EPI) requires the coordinates of the electrons $\{r^e_{\lambda}\}$, and coordinates of the nuclei $\{r^n_j\}$

$$H_{ep} = \sum_{\lambda, j} V(r^e_{\lambda}, r^n_j).$$  \hspace{1cm} (9.80)

First assume the nuclear motions are slow enough so the electron levels adapt continuously to the evolving structure, i.e., the Born–Oppenheimer approximation. This “adiabatic approximation” does not allow the nuclear kinetic energy to alter the electron states.\(^ {16}\) The adiabatic EPI increases with the number of phonons, but it also scales with the thermal excitations of electrons. (Effects

\(^{15}\)For a harmonic phonon, the electron-phonon interaction affects equally the energies of the electrons and the phonons.

\(^{16}\)The “non-adiabatic” electron-phonon interaction accounts for how the electronic states are altered by the nuclear kinetic energy, not the potential energy of displaced nuclei as for the adiabatic case. The non-adiabatic EPI does require thermal activation, and can be responsible for superconductivity.
from how the interatomic potential depends on phonon populations alone are already counted as phonon-phonon interactions.)

In this adiabatic picture, unique electron eigenstates exist for a snapshot of the nuclear thermal displacements, but these evolve continuously as the nuclei move. Taking for reference a ground state with the nuclear positions of a perfectly periodic crystal and the electronic structure at 0 K, we seek to account for how the displaced nuclear positions alter the thermally-excited electronic energy levels (changing the electronic excitations induced by temperature).

For a given electronic structure, the adiabatic electron-phonon interaction is proportional to the number of phonons, \( n(\epsilon, T) + \frac{1}{2} \), times the difference of electron occupancy with respect to the ground state, \( f(T) - f(0) \), where \( f(T) \) is the Fermi-Dirac distribution. When there are sharp features in the electronic DOS near the Fermi level, there will be other temperature-dependences associated with the thermal sampling of the excited electron states.

How the electron eigenstates change with nuclear displacements is a problem that generally requires considerable detail about the states of electrons and phonons. A simple approach is to consider an electronic band formed from isotropic \( s \)-electrons, and a uniform dilation as may be associated with longitudinal phonons of long wavelength,

\[
H_{ep}^{D} = -\sum_{\lambda} D \Delta(\rho_{\lambda}^{3}) .
\]

(9.81)

where \( \Delta(\rho_{\lambda}^{3}) \) is the fractional change in volume at \( \rho_{\lambda}^{3} \), and \( D \) is a “deformation potential,” typically a few eV.\(^{17}\) This simple approach is convenient because the \( \Delta(\rho_{\lambda}^{3}) \) is related to the displacement \( \vec{u} \) as through its divergence: \( \Delta(\rho_{\lambda}^{3}) = \vec{\nabla} \cdot \vec{u} \), so

\[
H_{ep}^{D} = -D \sum_{\lambda} \vec{\nabla} \cdot \vec{u}(\rho_{\lambda}^{3}) .
\]

(9.82)

For longitudinal phonons

\[
\vec{e}(k_{l}) = \frac{\vec{k}_{l}}{k_{l}} ,
\]

(9.83)

and by taking the divergence of Eq. 9.77 we have, now in the occupation number representation

\[
\hat{H}_{ep}^{D} = -iD \sum_{\rho_{\lambda}^{3}} \sum_{k_{l}} \sqrt{\frac{\hbar}{2N m \alpha_{k_{l}}}} |\vec{k}_{l}| e^{i\vec{k}_{l} \cdot \rho_{\lambda}^{3}} A_{k_{l}}
\]

(9.84)

\(^{17}\)It is not typical to have electron energies shifted by several eV under high compressions, so the \( D \) typically pertain to small elastic distortions over a finite volume of crystal.
Consistent with the second quantization formalism for phonons, fermion field operators are used for the electrons

$$\Psi^\dagger(\vec{r}_n^\lambda) = \sum_{\vec{k}_i} C_{\vec{k}_i}^\dagger e^{-i\vec{k}_i \cdot \vec{r}_n^\lambda}, \quad (9.85)$$

which when applied to a multi-electron state, places an electron in the state $\vec{k}^\lambda$ (the operator $C_{\vec{k}_i}$ removes an electron from this state). When small, $\hat{H}^{D}_{\text{ep}}$ in Eq. 9.84 is a perturbation that mixes electronic states as $\langle \Psi_f | \hat{H}^{D}_{\text{ep}} | \Psi_i \rangle$. This integral can be evaluated by using the fermion field operators $C$ and $C^\dagger$ (Eq. 9.85), where the exponential phase factors cancel the $k$-space integration unless there is a conservation of wavevector. This forces the same total $\vec{k}^\lambda + \vec{k}_i$ for the creation operators as for the annihilation operators

$$\hat{H}^{D}_{\text{ep}} = -iD \sqrt{\frac{\hbar}{2N m}} \sum_{\vec{k}_i} \sum_{\vec{k}_i} \sqrt{\frac{\vec{k}_i}{c_L}} A_{\vec{k}_i} C_{\vec{k}_i+\vec{k}_i}^\dagger C_{\vec{k}_i}^\dagger, \quad (9.86)$$

plus an analogous term with $A_{\vec{k}_i}^\dagger C_{\vec{k}_i} + C_{\vec{k}_i}^\dagger$ for phonon creation. For Eq. 9.86 we also used a linear dispersion relationship $\omega_L(\vec{k}_i) = c_L \vec{k}_i$, appropriate for long-wavelength, longitudinal acoustic phonons. In accounting for the effects of phonons on the electron states, it is usually necessary to consider a more general case than the isotropic deformation potential of Eq. 9.81 as

$$I \propto \langle \vec{k}_n^\lambda + \vec{k}_n^\lambda | \vec{e}_j^\lambda(\vec{r}_n^\lambda) \cdot \nabla_{\vec{r}} \varphi_1(\vec{r}_n^\lambda, \vec{r}_j^\lambda) | \vec{k}_n^\lambda \rangle, \quad (9.87)$$

where the gradient is with respect to the nuclear coordinates.\(^{18}\)

In general, there are two lower order terms that are used to describe the electron-phonon interaction. The first is a generalization of the previous result Eq. 9.86

$$\hat{H}^{1}_{\text{ep}} = \sum_{\vec{k}_i} \sum_{\vec{k}_i} \nu^{\text{ep}}(\vec{k}_i, \vec{k}_i^\lambda) A_{\vec{k}_i} C_{\vec{k}_i+\vec{k}_i}^\dagger C_{\vec{k}_i}, \quad (9.88)$$

It accounts for the processes where an electron is excited from state $\vec{k}^\lambda$ to $\vec{k}^\lambda + \vec{k}_i$, simultaneously with the annihilation of a phonon in state $\vec{k}_i$, or creation of a phonon in state $-\vec{k}_i$. Momentum conservation restricts significantly these three-phonon processes. The second low-order term for electron-phonon coupling is $\hat{H}^{2}_{\text{ep}}$, and it includes a factor from second-order perturbation theory,

\(^{18}\)“Frölich coupling” is another example of an EPI, where longitudinal optical phonons create an electric field in an ionic solid. In this case, the Fourier transform of a Coulomb potential gives a factor of $4\pi k^{-2}$, and with two ions involved, $\hat{H}^{F}_{\text{ep}}$ goes as $k^{-1}$ (instead of the $k^{1/2}$ for $\hat{H}^{D}_{\text{ep}}$ in Eq. 9.86). Transition rates, or scattering rates, go as the square of these quantities.
Further Reading

The contents of the following are described in the Bibliography.


Chapter 10

Chopper Spectrometers

10.1 Concept of a Chopper Spectrometer

Enrico Fermi was unusual. It is a remarkable and profound honor that the name “fermion,” with a lower case f, is standard terminology. The field of experimental neutron scattering offers him a smaller honor with the term “Fermi chopper” (Fig. 10.1). A generic Fermi chopper is depicted in Figure 10.2. The ARCS instrument is in fact a Wide Angular Range Direct Geometry Fermi Chopper Spectrometer.\(^1\)

The Fermi chopper is discussed at length in Section 10.4. For now we note that it works as a fast shutter. By aligning its slot at the right instant after a neutron burst leaves the moderator, it selects a bunch of neutrons that have a particular velocity. We know the time of the neutron burst and the distance between the neutron source and the Fermi chopper, so we therefore know the neutron velocity and its energy, \(E = \frac{1}{2}mv^2\). The idea is best illustrated with the time and distance diagram in Figure 10.3.

The slopes of the lines in Figure 10.3 are velocities of neutrons. Neutrons of many velocities are emitted from the moderator. The figure depicts two neutron pulses from the moderator, separated in time by perhaps 1/60 sec. The Fermi chopper selects a narrow range of velocities, corresponding to a time window of tens of microseconds. Of course the narrower this window, the more neutrons are blocked by the chopper, and the lower the flux of neutrons on the sample. On the other hand, the narrower this window, the more precisely selected are the velocities, and hence better energy resolution is achieved. The Fermi chopper therefore controls the incident energy, the intensity, and the energy resolution of the neutrons incident on the sample.

The sample usually transmits most of the incident neutrons without scattering, and most of the scattered neutrons are scattered elastically. There is a strong tendency for the lines in Figure 10.3 to have unchanged slopes through

\(^1\)Unfortunately for Enrico Fermi, the acronym ARFCS is less euphonious than ARCS.
CHAPTER 10. CHOPPER SPECTROMETERS

Figure 10.1: Enrico Fermi works with an electronic control for a neutron chopper during his Argonne days.

Figure 10.2: Fermi chopper, comprising a spinning cylinder with a slot in it. Neutrons are transmitted only when the slot is aligned properly along the path of the beam. The chopper works by scattering neutrons out of the forward beam, or by absorbing them by the inner surfaces of the slot. The ARCS choppers work primarily by absorption.
10.1. CONCEPT OF A CHOPPER SPECTROMETER

Figure 10.3: Distance–time diagram for inelastic scattering by a direct geometry chopper spectrometer. Positions of the moderator, chopper, sample, and detectors are marked on the vertical axis. Two moderator pulses occur at points on the $x$-axis. (Real pulses are not instantaneous, and this issue is discussed below.) When neutrons reach the position of the chopper, they are transmitted, absorbed, or scattered.
It is straightforward to know the total distance from the neutron source to the sample, and from the sample to the detector tube. Given the velocity selected by the Fermi chopper, we can figure out the time of arrival of the elastic neutrons at the detectors. Experimentally, we observe an intense elastic peak in the time spectrum at any of the detectors. This is typically quite close to the predicted arrival time, but the experimental time is used to identify the elastic scattering. Those neutrons that are scattered inelastically, however, have lines with kinks at the sample, and after the sample their slopes in Fig. 10.3 are either steeper or shallower than for the incident beam. The neutrons that arrive earlier have gained energy from the sample, and those that arrive later have lost energy to the sample. The inelastic spectrum is obtained from the histogram of neutron arrival times. A Fermi chopper spectrometer works entirely by timing.

### 10.2 Neutron Sources

#### 10.2.1 Spallation

It is a challenge to produce “free neutrons,” meaning neutrons that are “free of the nucleus.” "Spallation" is one way to make them. The “spallation” process got its name by analogy to using a hammer to “chip” pieces off a heavy stone. Here the hammer is the particle beam, the chips are neutrons, and the stone is the nucleus. Other things come out of the nucleus besides neutrons, especially $\gamma$ radiation. High-energy protons (of order 1 GeV) are preferred for the particle beam because protons produce more neutrons, and less heat and photons than are produced by electron beams. The major component of a spallation neutron source is therefore a high-energy, high-current proton accelerator. It often includes a linear accelerator followed by a buncher ring to compress the proton pulse into a short burst in time. Neutron yields are largest for nuclei of high atomic number, since these are neutron-rich. Tungsten and mercury are good choices in practice. Uranium is even better, but it tends to give problems at higher power levels.

#### 10.2.2 Moderation and Moderator Spectrum

The neutrons fresh from a spallation reaction have energies of order MeV, but the neutrons used in inelastic scattering have energies of order 100 meV. The excessive energies of the spalled neutrons need to be “moderated.” The

---

2 Elastic scattering will change the direction of the neutron, but this cannot be depicted on Fig. 10.3.

3 An excess of neutron over protons is required for stability. Too many protons means too much Coulombic repulsion, but neutrons can help overcome this instability and keep the nucleus together. Nevertheless, too many neutrons will “drip” out of the nucleus, and neutron-rich isotopes near the neutron “drip line” are good candidates for the target material.
“moderator” makes the transition between the neutron target and the neutron instrument by delivering a useful spectrum of neutrons to the instrument.

How does the moderator reduce the neutron energy by a huge factor of $10^7$? By inelastic collisions with nuclei in the moderator. In an inelastic collision between a neutron and a nucleus of the same mass, hydrogen of course, up to half of the kinetic energy of the collision can be transferred to the hydrogen nuclei. The hydrogen in the moderator can be in various chemical forms such as liquid hydrogen, water, or solid methane, depending on the desired temperature and density, for example. The number of collisions required for moderation, $n$, is

$$2^n \approx 10^7, \quad (10.1)$$

$$n \approx 23. \quad (10.2)$$

For the moderator of the ARCS spectrometer, the process of moderation therefore involves a relatively small number of collisions, and is therefore a statistical process. As an approximation it is often assumed that the neutrons leaving the moderator consist of two components. The first is a fully moderated spectrum of thermal neutrons. These neutrons have the Maxwell-Boltzmann spectrum, with probabilities based on Boltzmann factors of $\exp\left(\frac{-m_p v^2}{kT}\right)$, where $T$ is the temperature of the moderator. The second subspectrum is called the “epithermal” neutron spectrum. It is a broad spectrum with a tail that goes to very high energies. Epithermal neutrons have not undergone enough interactions with the moderator to acquire a thermal distribution. This approximation of two subspectra has some semblance of the truth.

A better moderator spectrum can be calculated by Monte Carlo neutron transport codes, such as the one that produced the spectra of Figure 10.4a. The peak intensity for the “decoupled, poisoned moderator” is around 40 meV, but there is useful intensity over a broad range of energies beyond 1 eV. Figure 10.4 shows similar calculations for a moderator that uses liquid hydrogen. Its spectrum is peaked at a much lower energy, perhaps 3 meV. The total number of neutrons is somewhat better, but by bunching up their energy into a narrow range, the usable intensity is much higher. “Cold neutrons” are a new frontier in neutron scattering research.

### 10.2.3 Decoupled, Poisoned Moderator

An important consideration for moderator design is “coupling,” meaning the connection between the target that is hit by the proton beam, and the material of the moderator. After slowing down in the moderator, the slower neutrons can be absorbed by materials such as Cd and Gd. Putting a layer of Cd or Gd between the moderator and the target can suppress the transmission of the slower neutrons back into the target. This is a “decoupled” moderator. (On the other hand, the fast neutrons from the target are not absorbed as they enter the moderator.) In contrast, a “coupled” moderator allows transmission of all neutrons between the target and the moderator, and less absorption.
Figure 10.4: Monte Carlo simulations of intensity spectra and time spectra for two SNS moderators. ARCS is on the decoupled, poisoned water moderator. For comparison a cryogenic hydrogen moderator is also shown. (a) Total intensity. (b) Mean emission time.
A coupled moderator produces more neutrons, but it has a disadvantage in the time structure of its neutron pulse. The slow neutrons traveling around the coupled moderator take some time before entering the neutron instrument. This time delay degrades the energy resolution of the Fermi chopper spectrometer, which works by timing.

Another trick to producing short neutron bursts from the moderator is to put a neutron absorber such as Cd or Gd at some depth inside the moderator itself. Again, the idea is to absorb the slow neutrons that do not follow an efficient path out of the moderator. This is called “poisoning” the moderator. A “poisoned, decoupled moderator” generates neutron bursts that are short in time, and this is the moderator of choice for the ARCS instrument, for example. For neutrons having energies below 100 meV, the moderator for ARCS emits neutron bursts of approximately 20 microseconds. In contrast, an “unpoisoned, coupled moderator” generates more neutrons in each burst, although these neutrons are emitted over a longer time.

An additional complexity is that the neutrons of different energies are emitted from the moderator at different times. Of course the neutrons with highest energies, which do not undergo enough collisions with the nuclei in the moderator, are emitted in the shortest times after the proton pulse hits the target. In general, the lower-energy neutrons leave the moderator at later times, with a broader spread in their emission times. The details of this time-energy correlation are not simple, however, and are best understood by Monte Carlo simulations and experimental measurements. Nevertheless, the neutron emission times affect the energy resolution of the spectrometer, as discussed in Section 10.4. Figure 10.4b shows this energy dependence of the pulse emission times for two SNS moderators. It also shows clearly that the emission time is shorter for the decoupled and poisoned ARCS moderator than for the coupled, unpoisoned hydrogen moderator.

10.3 Neutron Guides

10.3.1 Geometrical Optics

Here we develop the scattering of neutrons at interfaces between two homogeneous media. In a homogeneous potential, a neutron wavefunction propagates forward without deflection, but it has a wavelength that depends on the potential. This is much like the propagation of light through glass, for example. The only deflections occur at interfaces, or at changes in the “density.” With geometrical optics we can readily utilize the familiar constructions of light optics, scaled appropriately for neutrons. In the present analysis we justify the “geometrical optics” approach to analyzing neutron scattering from macroscopic objects, especially mirrors.

Recall from (1.74) and (1.75), or (6.39) the integral form of the Schrödinger
equation in the Born approximation:

\[ \Psi_{sc}(\vec{Q}) = -\frac{m}{2\pi\hbar^2} \int V(\vec{r}') e^{i\vec{Q} \cdot \vec{r}'} d^3r' , \]  

(10.3)

where we have ignored the standard form of the outgoing spherical wave that properly multiplies (10.3) (and sets the relationship between \( f(\vec{Q}) \) and \( \Psi_{sc}(\vec{Q}, \vec{r}) \)). For nuclear scattering we use the “Fermi pseudopotential” of (6.40), which places all the scattering potential at a point nucleus at \( \vec{r} \):

\[ V_{\text{nuc}}(\vec{r}) = 4\pi\frac{\hbar^2}{2m} \bar{b} \delta(\vec{r}) , \]  

(10.4)

where \( \bar{b} \) is a simple constant (perhaps a complex number). The next step is to place numerous Fermi pseudopotentials at the positions of all \( N \) nuclei in the material, \( \{\vec{r}_i\} \):

\[ V(\vec{r}) = 4\pi\frac{\hbar^2}{2m} \bar{b} \sum_{\vec{r}_i} \delta(\vec{r} - \vec{r}_i) , \]  

(10.5)

where \( \bar{b} \) is the average scattering length per nucleus (assuming a mix of isotopes or elements). Substituting (10.5) into (10.3), we notice the handy cancellation of many constant prefactors:

\[ \Psi_{sc}(\vec{Q}) = -\bar{b} \int \sum_{\vec{r}_i} \delta(\vec{r} - \vec{r}_i) e^{i\vec{Q} \cdot \vec{r}_i} d^3r_i . \]  

(10.6)

In a homogeneous medium we can consider wave motion only in the forward direction, for which \( \vec{Q} = 0 \). In the forward direction, \( e^{i\vec{Q} \cdot \vec{r}_i} = 1 \), simplifying (10.6):

\[ \Psi_{sc}(\vec{Q}) = -\bar{b} \int \sum_{\vec{r}_i} \delta(\vec{r} - \vec{r}_i) d^3r_i , \]  

(10.7)

\[ \Psi_{sc}(\vec{Q}) = -N\bar{b} . \]  

(10.8)

Note how the details of the positions \( \{\vec{r}_i\} \) are lost for forward scattering. We can obtain the same equation (10.8) if, instead of placing \( \delta \)-functions at all nuclei, we use a homogeneous potential throughout the entire material. Instead of using (10.5), equation (10.8) can be obtained by using the following homogeneous potential for \( V(\vec{r}) \) in (10.3):

\[ v_0(\vec{r}) = 4\pi\frac{\hbar^2}{2m} \bar{b} \rho , \]  

(10.9)

where \( \rho \) is the number of nuclei per unit volume.
This is an important observation. Using (10.9) instead of (10.5), we make the transition from individual scatterings by atomistic Fermi pseudopotentials to geometrical optics. The neutron wave is now considered to travel through a homogeneous potential. The neutron is treated as propagating without scattering, although with a different wavevector depending on the “density,” $\bar{b} \rho$.

### 10.3.2 Total Reflection

When a neutron of energy $E$ enters a region where the potential energy $v_0(\vec{r})$ is positive, its kinetic energy is reduced and its wavelength is increased (see Fig. 10.5). The kinetic energy cannot go below zero, of course, and a consequence is that some neutrons may not have enough energy to enter a material having a positive $v_0(\vec{r})$ (except for some surface penetration). Substituting typical numbers into (10.9), we find that $v_0(\vec{r}) = 3 \times 10^{-4}$ meV, corresponding to a neutron wavelength of 500 Å. Reflection will be total, since the neutron cannot penetrate into the solid, but the neutron is conserved. This result of 500 Å pertains to neutrons arriving normal to a surface. Neutrons of wavelength longer than this critical wavelength will be reflected by the surface. A critical wavevector is shown in Fig. 10.6.

It is possible to get total reflection of more energetic neutrons if they arrive at shallow angles to the surface. The effect is analogous to the case of total internal reflection of light, where a light ray moving in a medium of high index of refraction can be totally reflected at an interface with a medium of low index of refraction, if it reaches this interface at an angle below a critical angle. Such a case is shown in Fig. 10.6. The neutron travels a bit slower in the Ni than in the vacuum. In the Ni, the neutron has a higher potential energy, a lower kinetic
energy, and a longer wavelength. Notice the matching of the wave crests across the interface. This continuity of the neutron wavefunction forces a change in direction of the wavevector across the interface – this allows for differences in wavelengths in the vacuum and in the Ni.

The relationship between the neutron wavelength and the critical angle for a homogeneous potential $v_0(\vec{r})$ can be derived from the Schrödinger equation (10.14) by separation of variables. Start with the neutron wavefunction for a neutron moving as a plane wave in the $x$-$z$ plane, where $\hat{z}$ is normal to the interface between, say vacuum and nickel metal. (For neutrons, the nickel metal has the lower index of refraction than the vacuum.)

\begin{align*}
\Psi_{sc}(k, r) &= e^{i\vec{k} \cdot \vec{r}}, \\
\Psi_{sc}(k, x, z) &= e^{ikx} e^{ik_z z}, \\
\Psi_{sc}(k, x, z) &= \psi_x(x) \psi_z(z).
\end{align*}

Substitute (10.13) into the Schrödinger equation (1.53):

\begin{equation}
\frac{\hbar^2}{2m} \left( \frac{\partial^2 \psi_x}{\partial x^2} + \frac{\partial^2 \psi_z}{\partial z^2} \right) + V(z) \psi_x(x) \psi_z(z) = E \psi_x(x) \psi_z(z),
\end{equation}

where we have assumed the potential varies only along $z$, changing only at the interface. Dividing through by $\psi_x(x) \psi_z(z)$ and rearranging:

\begin{align*}
-\frac{\hbar^2}{2m} \frac{\partial^2 \psi_z}{\partial z^2} + V(z) - E &= + \frac{\hbar^2}{2m} \frac{\partial^2 \psi_x}{\partial x^2}, \\
&= -\varepsilon.
\end{align*}

We have separated the $z$-dependence from the $x$-dependence. The left-hand side of (10.15) depends only on $z$, the right only on $x$, but both $z$ and $x$ can change independently. This means that both sides can only be equal to
10.3. NEUTRON GUIDES

a constant, which we denoted $-\varepsilon$ in (10.16). The two equations for $\psi_x(x)$ and $\psi_z(z)$ become:

\[
\frac{\hbar^2}{2m} \frac{\partial^2 \psi_x}{\partial x^2} = -\varepsilon \psi_x, \quad (10.17)
\]

\[
\frac{\hbar^2}{2m} \frac{\partial^2 \psi_z}{\partial z^2} = \left[ \varepsilon - E + V(z) \right] \psi_z, \quad (10.18)
\]

To obtain the required a plane wave solution for a propagating neutron, $\psi_x = e^{i(\sqrt{2m}\varepsilon)x/\hbar}$, we see that $\varepsilon$ must be positive. Our neutron is moving mostly along the $x$-direction at a glancing angle to the surface, with kinetic energy $E$ when it is in the vacuum. We therefore know that $\varepsilon - E$, although negative, must be rather small. It is therefore possible for a small positive $V(z)$ to switch the sign of the right-hand side of (10.17) from negative to positive. The consequence is interesting. The solution for $\psi_z(z)$ changes from a propagating wave to a damped exponential function. When $V(z)$ is sufficiently positive, the neutron wavefunction therefore does not propagate into the nickel. It is instead reflected from the interface. Of course the $V(z)$ for nickel is fixed, but we can alter the incident angle to get the same effect as shown in Fig. 10.5. By reference to (10.12) and (10.13), for example, we can see that the change in sign of (10.17) occurs when we select $k_z$ so that:

\[
k_{z,\text{crit}} = \frac{\sqrt{2m\nu_0}}{\hbar} = \sqrt{4\pi \frac{\overline{b}}{\rho}}. \quad (10.19)
\]

10.3.3 Critical Angle

The critical angle for total reflection, $\phi_{\text{crit}}$, is the ratio of $k_{z,\text{crit}}$ to the wavevector along $x$, $k_x$, (the magnitude of $k_x$ is essentially the same as the magnitude of the incident wavevector $k$):

\[
\phi_{\text{crit}} = \frac{\sqrt{4\pi \overline{b}\rho}}{k} = \frac{\lambda}{2\pi} \sqrt{4\pi \overline{b}\rho}, \quad (10.20)
\]

\[
\phi_{\text{crit}} = \lambda \sqrt{\frac{\overline{b}\rho}{\pi}}. \quad (10.21)
\]

The critical angle increases in proportion to the wavelength of the neutron. Lower-energy neutrons can be reflected from a surface at higher angles than higher-energy neutrons. For natural Ni metal, the evaluation of (10.21) gives the wavelength-dependence of the critical angle, $\phi_{\text{crit}}$, here converted from radians to degrees:

\[
\phi_{\text{Ni, crit}}[^{\circ}] = 0.0991 \lambda [\text{Å}]. \quad (10.22)
\]

Finally, neutron reflectivity can be used for making spin-polarized neutron beams. The coherent scattering lengths are different when neutrons of up spin
or down spin are scattered by a magnetic moment having a component along \( \hat{z} \). For Co, in fact, the scattering lengths, \( b_{\uparrow}^{\text{Co}} \) ad \( b_{\downarrow}^{\text{Co}} \), are of opposite sign. A polarizer can be built by choosing incident angles where one of the neutron polarizations is transmitted into the magnetic material, and the other reflected.

### 10.3.4 Guide Design

For neutrons incident on a surface below the critical angle, simple ray diagrams can be used to develop conceptual designs of neutron guides. Their characteristics are analogous to cylindrical optical fibers for light, but neutron guides are usually made of four long mirrors with a rectangular cross section of order \( 0.1 \times 0.1 \text{ m}^2 \). A section of the ARCS guide is shown in Fig. 10.7. The four mirrors are seen end-on in the drawing at the top left.

The critical angle for Ni metal is a reference standard, but today a “super-mirror” can be prepared from multiple layers of metals, giving higher critical angles by a factor \( m \), where \( \phi_{\text{crit}} = m \phi_{\text{Ni crit}} \). Today the upper limit to \( m \) has reached 7. Since these angles are quite small for neutrons of meV energy, guides tend to be quite long. Nevertheless, focusing guides can be designed with parabolic or elliptical surfaces, for example, and these are efficient for transporting neutrons over long distances.

Efficient neutron transport is usually the main function of a guide. With a guide in its incident flight path, the instrument can be placed a good distance away from the moderator. This allows more space around the instrument, which can be important when instruments are crowded around a small neutron target. The separation from the moderator also allows the instrument to be
10.3. NEUTRON GUIDES

placed in a location with lower levels of background radiation. Two other
issues are useful for understanding the use of guides for neutron transport.

- By giving the mirrors of the guide a slight curvature along the beam
direction, the direction of the neutron beam can be bent away from its
original straight-line path out of the moderator. This makes it possible
for the specimen to be out of a line-of-sight path from the moderator, and
therefore less subject to background from fast neutrons.

- Most importantly, the guide reduces the usual attenuation of neutron
flux with distance. Instead of the $r^{-2}$ fall-off of intensity, a good guide
should cause minimal loss of neutrons that enter the guide below the
critical angle. (Note that those entering at higher angles would often be
blocked by collimators before the specimen anyhow.) A simple straight
mirror guide essentially takes the numbers and divergences of neutrons
entering the guide, and translates this distribution to the exit of the guide.

10.3.5 Brightness

Inelastic scattering measurements require beams on samples that have:

- High neutron current [neutrons/sec]
- Small size [cm$^2$]
- High flux [neutrons cm$^{-2}$ s$^{-1}$]
- Low divergence [$\alpha$, radians] (This requirement is unimportant if the $Q$-
  dependence is not of interest.)

Even if the neutron guide optics were perfect, and did not absorb neutrons
or cause unnecessary spread in divergence, for example, compromises will
always be required for meeting all these criteria of a good incident beam. These
compromises are quantified in the present section.

A fundamental problem with neutron optics is the size of the moderator.
With a cross section of order $0.1 \times 0.1$ m$^2$, the moderator is far from being a
point source of neutron emission. This gives a fundamental limitation to the
“brightness,” $\beta$, which is depicted with the three sources shown at the top of
Fig. 10.8. All three moderators in Fig. 10.8 emit the same neutron current,
and they send the same flux (current density) into the guides, which focus the
rays on the sample below. The sources to the left have the higher brightness,
and sources with higher brightness are better for making the smallest neutron
beams on the sample. The focused spot on the specimen is, in fact, an image of
the source itself, so it should be easiest to form a small spot when the source is
small.

The source of Fig. 10.8c has the lowest brightness. Nevertheless, the focused
beams in Figs. 10.8b and 10.8c have the same size. To make a small spot on the
specimen with the low brightness source of Fig. 10.8c, however, the guide in Fig.
10.8c must provide stronger focusing, i.e., a larger angle of convergence. (Good focusing with a large angle of convergence requires higher quality optics.)

More quantitatively, the brightness of the source, \( \beta \), is defined as the flux per solid angle \( \left[ \text{neutrons/s cm}^2 \text{ sr} \right] \), measured at the source of the neutrons. Brightness is a valuable concept because brightness is a conserved quantity when the subsequent optics are ideal. For example, after a guide focuses the beam as in Fig. 10.8c, the width of the focused neutron beam is reduced by a factor of two compared to the source, but the angle of convergence is increased by a factor of 2. In other words, the flux has increased by a factor of 4, and the solid angle has increased by a factor of 4, leaving unchanged the flux per solid angle (the brightness is conserved). Where the focused beam hits the specimen:

\[
\beta = \frac{j_0}{\alpha_x \alpha_y}. \tag{10.23}
\]

Here \( j_0 \) is the flux (neutrons/cm\(^2\)) in the beam on the specimen, \( \alpha_x \) and \( \alpha_y \) are the the angles of beam convergence in the \( x \)- and \( y \)-planes. We can relate the beam size to the brightness of the source and the convergence angles of the guide, assuming perfect guide optics. The beam width, \( d_0 \), is related to the total neutron current, \( I_p \), by the relationship between current and flux:

\[
I_p = d_x d_y j_0. \tag{10.24}
\]

For simplicity, assume square cross-sections for the guide and moderator, so
10.4. FERMI CHOPPERS

\[ d_x = d_y = d \text{ and } \alpha_x = \alpha_y = \alpha. \]

Substituting (10.23) into (10.24) and solving for \( d_0 \):

\[ d_0 = \frac{\sqrt{\frac{I_p}{\beta}}}{\alpha}. \tag{10.25} \]

Equation 10.25 shows that the beam width \( d_0 \) improves (becomes smaller) in proportion to the product \( \alpha \sqrt{\beta} \), as suggested by the previous discussion of Fig. 10.8.

The compromises needed in guide design and experimental setups are therefore clear. We have to balance beam width against divergence. A high divergence impairs the \( Q \)-resolution of the instrument, but for a moderator of fixed brightness, a high divergence is required for a high neutron flux on a small sample. The trade-off is one of inverse proportionality. If \( Q \)-resolution is not an issue, however, small samples become more appropriate.

- To Do: plots of \( I(E) \) for ARCS

## 10.4 Fermi Choppers

A first glance at Figure 10.1 shows electronic equipment that looks a bit complicated, and in fact it is. The electromechanical control of a Fermi chopper is not simple, and requires specialized technology. The energy resolution of a Fermi chopper neutron spectrometer depends on timing, and much timing precision is demanded from the Fermi chopper. It must be open at a precise time after the proton pulse hits the neutron target. Microsecond precisions are needed for Fermi chopper timings, as we now show from an elementary calculation.

A neutron of 200.0 meV energy has a velocity of 6,185 m/s. It travels down a 11.6 m flight path in 1,876 \( \mu \)sec, where it encounters the ARCS Fermi chopper. If the energy resolution is to be 1%, the velocity resolution needs to be 2%. The Fermi chopper should therefore be open to the beam for 37 \( \mu \)sec. To ensure that all neutrons passing through the Fermi chopper have energies of 200 meV, this opening must occur at a reproducible time delay after each proton pulse hits the neutron target. Variations in this time delay correspond directly to an energy broadening (with no gain in intensity). The electromechanical control system for the Fermi chopper should ensure that the chopper is at the same angle of rotation after the required time delay, here 1,876 \( \mu \)sec. This “phasing accuracy” needs to be 2 to 4 \( \mu \)sec to be negligibly smaller than opening window of 37 \( \mu \)sec in our example.

This is a stringent demand – the slot in a rotating cylinder must be at the same orientation after each neutron pulse, with only a 2\( \mu \)sec margin for error. For the ARCS spectrometer, the SNS proton pulse on target has a pulse frequency of 60 Hz, but has some drift over time associated with drift in the Tennessee power grid. An electromechanical feedback control system ensures synchronization of the Fermi chopper rotor with the proton pulse. Today these
electronic units are somewhat smaller than the one being caressed by Enrico Fermi in Figure 10.1, but they are far more precise and reliable than his.

Another stringent demand can be understood from the opening of the slot in the rotor. The slot has an effective width that corresponds to perhaps $5^\circ$ of the rotor circumference. How fast does the rotor need to spin for the opening time of 37 $\mu$sec in our example? This rotational frequency is $(5/360)/(48 \mu\text{sec}) = 375 \text{ Hz}$. Such fast rotations present mechanical challenges for bearings and heat dissipation.\(^4\) In practice, Fermi choppers have magnetic bearings, and their rotors spin in a partial vacuum with some helium gas for heat transport. An obvious issue with our calculation of a 375 Hz rotational frequency is that it is not an integral multiple of the 60 Hz proton pulse frequency of the Spallation Neutron Source. In practice, we might select a 360 Hz rotational frequency so that the chopper can be open at a fixed time delay for every proton pulse. Note that the chopper will be open six times for each neutron pulse.\(^5\)

There are different opinions on what to do with the other five pulses that could pass through the Fermi chopper for each proton pulse on target. The relationship between neutron energy and neutron velocity is:

$$E_n = 5.2276 \times 10^{-6} v_n^2.$$  \hspace{1cm} (10.26)

For our hypothetical rotor spinning at 360 Hz (six rotations per proton pulse at 60 Hz), with a phase delay of 1,876 $\mu$sec to select a 200.0 meV energy, chopper openings will occur at time delays, $\tau$, of:

$$\tau = 1,876 + n \times 2,778 [\mu\text{sec}],$$ \hspace{1cm} (10.27)

where $n = \{0, 1, 2, 3, 4, 5\}$, and 2,778 $\mu$sec is one-sixth of $1/60$ sec. For these opening times and a 11.6 m flight path, the five incident energies are 200.0, 25.9, 9.6, 5.0, 3.0 and 2.0 meV. It might be possible to acquire additional inelastic scattering spectra from these five additional incident energies. This practice is called “repetition rate multiplication.” It should be noted that in the simple Fermi chopper configuration proposed here, the energy resolution becomes impractically narrow for the lower incident energies, however.

The ARCS spectrometer includes a second chopper, a $T_0$-chopper, located closer to the moderator than the Fermi chopper. This $T_0$-chopper is much like a Fermi chopper, and is phased to block the incident neutrons that would arrive at the Fermi chopper during our five other openings of the Fermi chopper. The $T_0$-chopper also serves to block the flash of $\gamma$-radiation and fast neutrons that are emitted by the moderator when the proton pulse hits the target.

The slot through a Fermi chopper has a gradual curvature to accommodate the time for neutron passage through the chopper rotor. This idea is shown in

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\(^4\) Disk choppers are an alternative to cylindrical Fermi choppers. For a neutron beam of moderate width, the disk needs to be large, and centrifugal stresses become excessive at 375 Hz.

\(^5\) There are actually twelve openings per proton pulse if you count the half-rotations where the back side of the rotor is facing the incident beam. For these cases, however, the rotor is upside-down in the figures below, and the curvature of the slats impedes the neutron transmission.
10.5. DETECTORS

Figure 10.9: Fermi chopper rotor with two slats. The structure at left is an exploded view of the layered slits and slats that fill the curved slot through the rotor.

Figure ?? for a neutron depicted as moving from left to right. The positions of the neutron and the chopper are shown for three snapshots in time, with $t_1 < t_2 < t_3$.

The slot through the rotor of a Fermi chopper contains a stack of slits and slats, which allow the Fermi chopper to have good energy resolution while passing a wide neutron beam. The structure is shown in Figure 10.9 for a simple case with two “slats.” The slats typically contain boron, a strong absorber of neutrons. The neutrons traversing the Fermi chopper must travel through the narrow gaps called “slits.” In practice these slits are plates of aluminum with many holes in them. The aluminum serves to maintain a precise spacing between the slats, a challenge at high rotational frequencies. In practice, a Fermi chopper may contain 10 to 40 slats. With many slats, it is possible for the Fermi chopper to pass a beam of 5 to 6 cm in width while maintaining good energy resolution. Not surprisingly, with narrower slits there is a loss of intensity that accompanies the improvement in energy resolution.

A drawing of the Fermi chopper used in the ARCS instrument is shown in Fig. 10.10. The neutron beam passes through the circular opening at the center of the housing. The electric motor, sensors, and magnetic bearings are above and below the rotor assembly. A corresponding pair of photographs is shown in Fig. 10.11. In the ARCS instrument, two identical chopper systems are mounted on a translation table so either can be moved into the neutron beam. The choppers can be moved while they are spinning, allowing quick changes to the energy of the incident beam of the ARCS instrument.

Rotor assemblies with curved slots, especially those with many slits and slats, are optimized to work at particular combinations of rotational frequencies, neutron velocities, and energy resolutions. These need to be selected in the planning stage of an inelastic neutron scattering experiment.

Finally, it is important to remember that Fermi choppers scatter neutrons out of the incident beam, generating a high intensity of scattered neutrons around a Fermi chopper. Good shielding is required, both for radiation protection for the experimenters, and for minimizing background in the instrument itself. The ARCS $T_0$ chopper helps alleviate this problem by minimizing the number of neutrons that are scattered out of the beam by the Fermi chopper.

10.5 Detectors

Neutron detectors usually rely on a nuclear transmutation reaction with a large cross-section. Widely used is $^3$He, which absorbs a neutron and undergoes
Figure 10.10: Exploded view of ARCS Fermi chopper, showing housing and rotor at left. The slit packages are held into the rotor with two pins, as shown at right.

Figure 10.11: Left: photograph of ARCS Fermi chopper, showing housing, windows, and water cooling. Right: photograph of the slit package, with aluminum slits seen end-on.
The \( {^3}\text{He} \) gas-filled detector depicted in Fig. 10.12 can be modified to locate the position of the neutron arrival along the length of the detector. The modifications include the use of a resistive wire as the anode, and preamplifiers at both ends of the anode wire. By comparing the electron charges collected on the capacitors at the two ends of the detector, the position of the event can be determined. A neutron that ionizes the gas at one end of the detector tube

\[
n + {^3}\text{He} \rightarrow {^1}\text{H} + {^3}\text{H} + 0.76\text{MeV}.
\]  

(10.28)
produces a larger pulse in the preamplifier connected to that end. Such linear position-sensitive detectors require that the resistivity of the anode wire be steady with time, and not affected by contamination from the detector gas, for example. A two-dimensional array of side-by-side $^3$He linear position sensitive detector tubes is often used as a 2-dimensional area detector. Modules of linear position-sensitive detector tubes are shown in Fig. 10.13, and the detector array of the spectrometer was completed by adding more detectors to the right of those shown.

### 10.6 Energy Resolution

Energy resolution is an important figure-of-merit for the design of direct geometry chopper spectrometers, and can be the most important figure-of-merit for incoherent scattering. The primary flight paths at the SNS are quite long, and the moderator pulses are short on flight paths 17 and 18, promoting good energy resolution. In evaluating instrument configurations, the secondary flight path is the parameter for adjusting energy resolution. When selecting a distribution of detector positions around the specimen, a spherical locus of detector positions provides energy resolution that is uniform at all angles around the sample.

Basic considerations for energy resolution were presented in the discussion of Fig. 10.3. Here we consider the issue further, starting with the moderator. The moderator does not produce an instantaneous pulse of neutrons, but the
10.6. ENERGY RESOLUTION

Figure 10.14: Distance–time diagram for a direct geometry chopper spectrometer. The figure is much the same as Fig. 10.3, but there is a spread in times when the neutrons leave the moderator. Positions of the moderator, chopper, sample, and detectors are marked on the vertical axis. Here the Fermi Chopper is open for only a minimal time.

The energy resolution of a Fermi chopper spectrometer is often analyzed in terms of the moderator performance and the chopper performance. For analytical calculations, it is typical to assume that both the moderator and the chopper cause time smearing that has a Gaussian shape about an average time. The advantage to this approach is that one can convolute the different effects of time smearing to get another Gaussian function, whose width is obtained by adding in quadrature the widths from independent broadenings. The resolution is then obtained in terms of the following variables for time, energy, and distance:

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6The paths from moderator to detectors in Fig. 10.14 are often compared to the ray diagram of a pinhole camera, where the pinhole is at the position of the chopper.
Chapter 10. Chopper Spectrometers

- $L_0$...distance from the moderator to the Fermi chopper
- $L_1$...distance from the sample to detector
- $L_2$...distance from the Fermi chopper to the sample
- $t_r$...time spread of neutrons passing through the Fermi chopper
- $t_m$...time spread of neutrons from the moderator
- $\delta_m$...distance the neutrons travel in time $t_m$
- $E_0$...incident energy selected by Fermi chopper
- $E_1$...energy of the neutron leaving the sample

It is typical to express the energy resolution as a fraction of the incident energy, $\Delta E/E$. The Gaussian analysis performed by Windsor [C.G. Windsor] gives:

$$\frac{\Delta E}{E} = \frac{2\delta_m}{L_0} \left[ \left(1 + \left(\frac{E_1}{E_0}\right)^{3/2} \frac{L_2}{L_1}\right)^2 + \left(\frac{t_r}{t_m}\right)^2 \left(1 + \left(\frac{E_1}{E_0}\right)^{3/2} \frac{L_0}{L_1} \left(1 + \frac{L_2}{L_0}\right)^2\right) \right], \quad (10.29)$$

Energy resolution can be presented in various ways, and it is important to be careful when comparing plots for different instruments. Perhaps the most important criterion is the neutron flux on the sample for a given energy resolution, calculated for conditions that optimize the intensity. Figures 10.15 and 10.16 show such plots for the ARCS instrument. The first figure shows the performance for incident neutrons of 63 meV energy, approximately at the peak of the moderator brightness. These neutrons have a longer time spread for emission from the moderator, however, and this limits the ultimate energy resolution to about 2.5% of the incident energy. Figure 10.16 shows the same plot for neutrons of 250 meV incident energy. The ultimate resolution is better, perhaps 2% of the incident energy, in large part owing to moderator performance. The intensity is lower at 250 meV for the modest resolution of 4%, however. The neutron guide is not so effective at 250 meV, and the moderator spectrum is weaker at this energy.

10.7 $Q$ Resolution

Optimization is quite different for $Q$ resolution, however. This has been less of a focus of the ARCS instrument design, however, because historical trends have emphasized the design of instruments for work with polycrystalline samples. Measuring dispersive excitations in single crystals forces the consideration of
Q resolution. Specifically, it is suggested that the instrument be optimized for fractional Q resolution. The figure-of-merit, $R_Q \equiv \Delta Q/Q$, is analogous to the figure-of-merit, $\Delta E/E$ for E resolution.

The parameter space for optimization is broad, and optimization depends on the type of measurement to be performed. Various types of excitations must be considered. The extreme cases are treated below as: 1) elastic scattering, 2) highly dispersive inelastic scattering and 3) dispersionless inelastic scattering.

### 10.7.1 Summary of Q Broadening

Four sources of Q broadening are considered:

- The finite size of the moderator, even over a long primary flight path, provides a significant angular divergence of neutrons on the specimen. This divergence is about 0.4° without a guide, and will be larger for thermal neutrons that have been reflected along the guide. If the excitations in the specimen do not select among these incident $\vec{k}_i$, the resolution of the experiment will be dominated by incident divergence.

- A 0.2° mosaic spread of a good crystal will cause an angular spread of 3.5 cm at 10 m. This is comparable to detector pixelation along a diagonal direction, which is approximately $2.5 \sqrt{2} = 3.5$ cm.

- The energy resolution leads to a coupled smearing in Q. Since the energy resolution is quite good for all flight paths, this source of Q smearing is
CHAPTER 10. CHOPPER SPECTROMETERS

not the biggest problem for single crystal work, especially in the forward direction.

- The sin $\theta$-dependence of $Q$ conspires with finite detector pixel size to give a divergent $l_3$ in the forward direction. A $R_Q$ of 1.0 % is not possible for scattering angles, $2\theta$, smaller than 14° if a 10 m secondary flight path is used with 2.54 cm detectors. The problem is proportionately worse for shorter flight paths.

10.8 Optimization for $\Delta Q/Q$ in Elastic Scattering

It is easiest to first analyze the $Q$ resolution for elastic scattering, and some of the issues for elastic scattering pertain directly to inelastic scattering.

10.8.1 Incident Divergence

The finite size of the moderator provides an angular divergence of incident neutrons on the sample. This is approximately 10 cm over 1350 cm, or about 0.4°. This incident divergence may or may not contribute to the divergence in scattered beams, depending on the process involved. For elastic Bragg scattering, for example, the incident divergence will allow intense Bragg peaks from all parts of a sample misoriented to within about 0.4°. Although incident divergence is a potentially serious problem, a perfect crystal will still provide sharp Bragg diffractions. This incident divergence can be the dominant effect on $Q$-resolution for polycrystalline samples, however.

10.8.2 Mosaic Spread

It is important to consider the limit to $Q$ resolution caused by the sample itself – the instrument need not be much better than the intrinsic resolution of the sample. One issue is that a finite sample subtends a non-zero angle over the secondary flight path, $l_3$. This can be ignored for a typical 1 cm sample, which is much smaller than the expected size of detector pixels.

The “mosaic spread,” denoting the mean mutual misorientations of different subvolumes in a crystal, is an important figure-of-merit for single crystal samples. A good single crystal sample may have a mosaic spread of 0.2°, for which the angular blurring of an elastic beam at a detector located 10 m away from the sample is 3.4 cm. This is only slightly larger than typical detector pixel resolutions. Better instrument resolution would be a reasonable request, except for the fact that a 10 m sphere of detectors around the sample is prohibited by constraints of cost and space. Nevertheless, we note that a long secondary flight path, $L_1 = 10$ m, could be justified for single crystals of high quality in cases where the incident divergence does not dominate the $Q$ resolution.
10.8. **OPTIMIZATION FOR $\Delta Q/Q$ IN ELASTIC SCATTERING**

### 10.8.3 Locus of Constant $\Delta Q/Q$

We seek the locus of detectors that provides a constant $\Delta Q/Q$, defined as the resolution, $R_Q$:

$$R_Q \equiv \frac{\Delta Q}{Q}.$$  \hspace{1cm} (10.30)

By definition

$$Q \equiv 4\pi \frac{\sin(\theta)}{\lambda}, \text{ so}$$  \hspace{1cm} (10.31)

$$\Delta Q = 4\pi \frac{\cos(\theta)}{\lambda} \Delta \theta,$$  \hspace{1cm} (10.32)

giving

$$R_Q = \cot(\theta) \Delta \theta.$$  \hspace{1cm} (10.33)

When $R_Q$ is plotted in polar coordinates, the detector locus is defined. This is shown in Fig. 10.17. This shape is quite incompatible with the spherical solution for constant $\Delta E/E$. At low angles, the detector placements required for constant $\Delta E/E$ and constant $\Delta Q/Q$ are exactly orthogonal.

The resolution $R_Q$ diverges at small angles, because at small angles $Q \to 0$, so very small $\Delta Q$ is needed to maintain a constant $\Delta Q/Q$. A small angular spread for $\Delta Q$ is achieved only when the detector pixels are very small or are placed at very large $L_1$. At the other extreme, the detector distance for constant $\Delta Q/Q$ is infinitesimal when the scattering angle, $2\theta$, is $180^\circ$, since small variations in angle have no effect on $Q$ for direct backscattering.

### 10.8.4 Coupling of $Q$ Resolution to $E$ Resolution

One obvious incompatibility of the $\Delta Q/Q$ optimization and the $\Delta E/E$ optimization is that the secondary flight path, $L_1$, becomes vanishingly small at $\theta = 90^\circ$. (This is strictly true only for the elastic scattering. For a fixed energy transfer, however, there will be a particular angle of inelastic scattering for which $\Delta Q = 0$ in Eq. (10.33).)

Poor energy resolution will cause a smearing in $Q$, increasing $R_Q$. For elastic scattering:

$$Q = k_f - k_i = \frac{\sqrt{2mE}}{\hbar} 2 \sin \theta,$$  \hspace{1cm} (10.34)

$$\frac{dQ}{dE} = \frac{\sqrt{2m}}{\hbar \sqrt{E}} \sin \theta,$$  \hspace{1cm} (10.35)

$$\frac{dQ}{dE} = \frac{Q}{2E} \sin \theta,$$  \hspace{1cm} (10.36)

$$\frac{dQ}{Q} = \frac{dE \sin \theta}{E \frac{1}{2}},$$  \hspace{1cm} (10.37)

so $\Delta Q/Q$ and $\Delta E/E$ are comparable for modest angles.
The obvious solution is to use a spherical detector locus for high scattering angles. The 3.0 m flight path of ARCS gives energy resolutions as small as $\Delta E/E = 1\%$. At $2\theta = 40^\circ$ this contribution to $\Delta Q/Q$ is less than 0.2 %, and this is even smaller at lower angles.

10.9 Optimization of $\Delta Q/Q$ for Inelastic Scattering

10.9.1 Ewald Spheres and Incident Divergence

Three important cases are depicted in Fig. 10.2. Case 1, elastic scattering, was the topic of the previous Sect. 10.8. Case 2, dispersive inelastic scattering, is considered first. This problem is presented in two parts, the case for the first Brillouin zone, and the case for higher Brillouin zones. Case 3 of Fig. 10.2 is presented last, and it is more straightforward.

For a fixed energy loss, the value of $|\vec{\epsilon}_f|$ is constant, although $\vec{\epsilon}_f$ may take different orientations. The kinematics of this scattering are understood most conveniently by use of the Ewald sphere construction, which is shown for elastic scattering in Fig. 10.3a. The Ewald sphere construction is useful for analysis of diffraction conditions because it identifies the relationship between the wavevector transfer, $\vec{Q}$, and the reciprocal lattice vector, $\vec{\tau}$, showing when the Laue condition, $\vec{Q} = \vec{\tau}$, is satisfied for diffraction. In inelastic scattering the momentum of the excitation, $\vec{q}$, plays a similar role to $\vec{\tau}$, especially in the first
10.9. OPTIMIZATION OF $\Delta Q/Q$ FOR INELASTIC SCATTERING

Figure 10.2: Three types of scattering processes (1) elastic scattering, (2) dispersive inelastic scattering, and (3) non-dispersive inelastic scattering.

Figure 10.3: (a) Ewald sphere construction for elastic scattering, showing allowed $\vec{Q}$ for Bragg diffraction. (b) Ewald sphere construction for inelastic scattering, showing allowed $\{\vec{Q}\}$.

Brillouin zone where $\tau$ is zero.

For the inelastic scattering process shown in Fig. 10.3b, all $\{\vec{k}_f\}$ have the same length (as for the elastic case of Fig. 10.3a), since the energy $E_f$ is identical for all orientations of $\vec{k}_f$. In both Figs. 10.3a and 10.3b, the allowed $\vec{k}_f$ make a sphere, which when placed at the tail of $\vec{k}_i$, defines the allowed $\{\vec{Q}\}$ as shown in the figure (where as usual $\vec{Q} \equiv \vec{k}_f - \vec{k}_i$).

The advantage of the Ewald sphere constructions of Fig. 10.3 is in analyzing tilts of the incident beam, as occurs for incident divergence, for example. The cases of Fig. 10.4 show conditions for elastic scattering before and after tilt. It is evident that the condition for momentum conservation (the Laue condition)

$$\vec{Q} - \vec{\tau} = 0$$

is generally violated by tilting, because tilts of $\vec{k}_i$ by the angle $\phi$ cause $\vec{Q}$ to
Figure 10.4: Ewald sphere construction for elastic scattering (a) before, and (b) after tilt of $\vec{k}_i$. Notice that $Q$ no longer touches the reciprocal lattice vector $\vec{\tau}$ after tilt.

Tilt by this same angle $\phi$. The consequence of this violation of momentum conservation is that coherent elastic scattering is altered, and Bragg diffractions may be eliminated, for example. Those neutron trajectories with improper $\vec{k}_i$ will not contribute to the Bragg diffraction.

10.9.2 First Brillouin Zone – Soft Dispersions

One case for inelastic scattering is shown in Fig. 10.5a. This case is a scattering from the first Brillouin zone, where the reciprocal lattice vector, $\vec{\tau} = 0$. The diameters of the two circles are set by the energy and momentum transfer to the excitation. In Fig. 10.5a, $|\vec{k}_f| = \sqrt{2}|\vec{k}_i|$, indicating that the neutron has lost half its energy to the solid. For the present case where the excitation is of energy $E_i/2$, we further assume a relatively soft dispersion where the momentum of the excitation $q$ is assumed the same as the momentum of the neutron, $k_f$. A circle is used for the locus of acceptable $\vec{q}$ (a circle is not expected for anisotropic crystals, of course). This circle has a large radius, as may be expected if a dispersive excitation has a soft dispersion relation.

The effect of tilt on the scattering condition is shown in Fig. 10.5b. It is evident by geometry that the tilt of $\vec{k}_i$ and $\vec{q}$ are by the same angle for this case of $\vec{q}$ lying in the first Brillouin zone. This could affect the ability of the neutron to excite the dispersive mode, at least if the dispersive mode is sharp in $\vec{q}$. Conditions where highly anisotropic dispersions have a delicate contact with the locus of $Q$ can also be envisioned.

10.9.3 Higher Brillouin Zones – Stiff Dispersions

The case of a high energy excitation is shown in Fig. 10.6. In this case the dispersion is very steep, and the energy is so high that the energy of the excitation is large, even when $q$ is small. In this particular case, there are no excitations in
Figure 10.5: Ewald sphere construction for inelastic scattering (a) before, and (b) after tilt of $\vec{k}_i$. Notice that although $\vec{Q}$ has the same magnitude, it changes orientation by the same tilt angle, $\phi$, as the incident beam.
Figure 10.6: Ewald sphere construction for inelastic scattering (a) before, and (b) after tilt of $\vec{k}_i$. In this case the excitation has a large $E$ for a small $q$. Notice that although $\vec{Q}$ has the same magnitude, it changes orientation strongly after tilt of the incident beam.

the first Brillouin zone because $q < Q_{\text{min}}$.\footnote{Experimentally, it might be prudent to use a larger $E_i$ and $k_f$ so that excitations in the first zone are allowed. In this case the result of Sect. 10.9.2 is recovered, that is the tilt angle of $\vec{k}_i$ equals the tilt angle of $\vec{q}$. The present results from higher Brillouin zones remain valid, however.} The excitation can occur in a higher Brillouin zone with the help of a reciprocal lattice vector, $\vec{\tau}$ so that

\[ \vec{Q} - \vec{q} - \vec{\tau} = 0. \] (10.39)

This case of a high-energy excitation is a higher Brillouin zone is interesting because a tilt of $\vec{k}_i$ does not induce a tilt of $\vec{Q}$ by the same angle. Figure 10.6 shows that small tilts have big effects on the orientation of $\vec{q}$. In essence, the reciprocal lattice vector can amplify the tilt of the incident beam on the rotation of $\vec{q}$. In this case it seems plausible that for sharp, stiff excitations, only a $\vec{k}_i$ of the correct orientation can generate the excitation. The mosaic spread of the sample will be able to pick these acceptable $\vec{k}_i$ from the incident beam, and the $Q$ resolution of the experiment will originate with the mosaic spread of the sample and not the incident divergence.
10.9.4 Non-Dispersive Excitations

The case of non-dispersive excitations in Fig. 10.2 is straightforward to analyze with Figs. 10.5 and 10.6. The point is that the circle of \( q \) can be of arbitrary radius, since the energy of the excitation does not depend on \( q \). For this reason, at a fixed energy transfer equal to that of the excitation, there will always be an excitation with an appropriate \( q \) to satisfy momentum conservation. All orientations of \( \vec{k}_i \) will be useful for generating the excitation, and the incident divergence will dominate over the crystal mosaic spread in setting the \( Q \) resolution. A similar result pertains to inelastic incoherent scattering.

10.10 Background

Much of the work in commissioning a new inelastic neutron spectrometer is eliminating sources of background in the measurements. Finding and exercising the different types of background contributions is a dark art, but some general guidance is possible.

The usual sources of background originate with neutrons that 1) were once good neutrons in the incident beam, or 2) are bad neutrons that entered the instrument uninvited. Unwanted neutrons can enter the instrument from the experiment hall around the instrument. A heavy shielding of borated hydrocarbons around the detector chamber can help protect humans in the experiment hall, but it is at least as important for keeping neutrons out of the instrument.

Perhaps a bigger problem comes from fast neutrons generated in the proton target region. Not all of these are moderated to reasonable velocities, and some of these fast neutrons enter the instrument and cause trouble. Concerning Fast Neutrons:

- Even with a heavy \( T_0 \) chopper made from iron and nickel, some 0.1% to 1% of the fast neutrons typically pass through. They then scatter in a number of ways from materials in the sample chamber and detector chamber, and can lose enough energy to be detected efficiently by \( ^3\text{He} \) detectors.

- On the way towards the detector chamber, either before or after the \( T_0 \) chopper, fast neutrons can interact with materials in the primary flight path, such as concrete shielding, and lose energy. Background problems from these “moderated” neutrons are tricky to diagnose because they can have a time structure similar to the wanted neutrons in the incident beam. They may, however, scatter from parts of the instrument that are nominally protected from the incident beam and cause unexpected effects.

Neutrons wanted in the incident beam can scatter in unwanted ways. Concerning Neutrons in the Incident Beam:

- The first concern in designing a neutron scattering experiment is to keep all materials out of the incident beam except for the sample itself. Obviously anything exposed to the incident beam (sample support, heating
elements, pressure cell) will scatter neutrons in much the same way as the sample.

- Peculiar effects can occur when neutrons scattered from the sample find their way into another material in the sample chamber or detector chamber, and scatter a second time. Multiple scattering is a problem inside thick samples themselves, but it is even trickier to sort out if a neutron scattered by the sample is again scattered by a pressure cell and then goes into the detectors of the instrument. Calculating multiple scattering is the topic of Section 12.2, but avoiding multiple scattering should be a goal of experiment design.

- There is often a “small-angle elastic scattering” from small-scale density variations in the sample or in the sample environment equipment. The forward beam is broadened as it leaves the sample, and the albedo of the beam may not all enter the “get lost” pipe and beam stop. These neutrons may scatter from the forward-mounted detectors, giving a source of neutrons that shines back into the spectrometer. Radial baffles can help hide detectors from scattering from other detectors across the instrument. A radial collimator around the sample itself also proves effective in suppressing scattering diametrically across the detector array.

Evidently there are numerous risks of background from neutrons that enter the instrument either invited or uninvited. An older viewpoint is that since the energies and directions of these neutrons are hard to understand, they can be modeled as a gas with a wide spectrum of energies. In other words, these unwanted neutrons go everywhere. Covering everything with absorbing material is good practice.

A more refined practice is to use a radial collimator around the sample so that the scattered neutron beams from the sample must take a straight line to the detector array. Neutrons that are scattered by other materials away from the sample position are absorbed by the walls of the collimator and do not reach the detector. (Usually this collimator oscillates by a small angle so it does not cast a shadow, and its absorption characteristics are averaged across the detectors.) Very large improvements in background are possible with an oscillating radial collimator, as shown in Sect. 12.3.4.

Last but definitely not least, another important practice to reduce background is to eliminate parts of the incident neutron beam that cannot illuminate the sample. If the sample is small and intercepts only a fraction of the area of the incident beam, it is often a good idea to eliminate the rest of the incident beam by upstream collimation before the beam reaches the sample. This is less of a concern when the sample is isolated by being hung on a thin wire or mounted at the end of a thin stick, for example. In the case of an isolated sample, the outer neutrons that miss the sample will go straight into the get lost pipe and beam stop. However, if there is anything around the sample that could be hit by the incident beam, the best practice is to remove this extra material, or collimate the incident beam so incident neutrons do not hit it.
10.11 Sample Design

Inelastic neutron scattering experiments require careful attention to the sample itself. Of course the sample must be in the correct state, satisfying basic criteria such as magnetic polarization, chemical composition, crystal structure, crystallographic quality or texture, temperature and pressure. Criteria for these sample states must not be compromised to the detriment of the science. The in-situ control of temperature, pressure, and magnetic field often leads to some compromises in experiment design, and expertise with the available “sample environment equipment” is essential in planning an experiment.

A common problem for inelastic scattering experiments is a sample that is too small. There is always a background in the spectrum from stray neutrons in the instrument, and inelastic neutron scattering experiments may be ruined by poor signal-to-noise ratios whenever this background has an intensity comparable to the scattering from the sample itself. Robust scattering from the sample usually means having a large quantity of sample. Experimenters are often dismayed that inelastic scattering measurements typically two orders-of-magnitude more sample than is typical for neutron diffraction measurements. The ideal sample for inelastic scattering will be as wide as possible so that it fills the cross section of the incident beam. With a wide sample, all neutrons in the beam will be candidates for scattering, and no parts of the beam will pass directly into the beamstop or generate unnecessary background.

The thickness of the sample in the direction of the beam needs to be chosen with care. It is possible to have a sample that is too thick. There are two limits that set upper bounds on the thickness of samples, and we consider each in turn below. The sample should be as thick as possible, provided it does not exceed limits imposed by:

- Multiple scattering. If the scattering cross-section is large and the sample is thick, too many neutrons will be scattered multiple times inside the specimen. Each scattering has its own energy spectrum, and the convolution of these spectra is difficult to sort out in the measured data.

- Absorption. Some nuclei are strong absorbers of neutrons. For thick samples of strong absorbers, few neutrons can leave the samples and enter the detectors.

10.11.1 Scattering and Attenuation

As a neutron beam passes through a material, there is a reduction in the number of neutrons in the forward beam. At the depth $x$, the increment of thickness of a material, $dx$, scatters a number of neutrons, $dI$, removing them from the beam. The number of scattered neutrons, $-dI(x)$, equals the product of 1) the increment of thickness, $dx$, 2) the number of neutrons present at $x$, $I(x)$, and 3)
a material coefficient, $s$:

\begin{align}
-dI(x) &= s I(x) \, dx, \\
\frac{dI(x)}{dx} &= -s I(x), \\
I(x) &= I_0 e^{-sx}.
\end{align}

(10.40)

(10.41)

(10.42)

If scattering is the only process that removes neutrons from the incident beam as it traverses a sample of thickness $t$, the loss from the beam equals the intensity of the scattering:

\[ I_{\text{scat}}(t) = I_0 \left( 1 - e^{-st} \right). \]

(10.43)

The product in the exponent, $sx$ or $st$, must be dimensionless, so $s$ has dimensions of $[\text{cm}^{-1}]$. When $sx$ is small, it equals the fraction of neutrons removed from the incident beam. From Fig. 1.4 we know that this fraction also equals $N\sigma/A$, so:

\[ s = \frac{N\sigma}{Ax} = \frac{N}{V}\sigma, \]

(10.44)

where $N/V$ has units $[\text{atoms cm}^{-3}]$ and $\sigma$ is the scattering cross-section with units $[\text{cm}^2]$.

8Since density varies with the type of material, “mass attenuation coefficients” are often normalized as ratios $s/\rho$. Here the density, $\rho$, has units $[\text{g cm}^{-3}]$, so the coefficients $s/\rho$ have units $[\text{cm}^{-1}] [\text{g cm}^{-3}] = [\text{cm}^2 \text{ g}^{-1}]$. Exponents in (10.42) are products $(s/\rho) \times \rho \times x$, and are, of course, dimensionless.

9Often the next largest fraction of neutrons are scattered elastically, and the smallest fraction are scattered inelastically.

10.11.2 Multiple Scattering Criterion

It may seem curious that in a well-planned experiment, most of the neutrons are transmitted through the sample without scattering. Using a very thick specimen can cause the data to be uninterpretable, as we now show.

Consider the probability of inelastic scattering, $p_i$, and elastic scattering, $p_e$, through a thin layer of material. We set $p_e + p_i = p$, where $p$ is the total

\[ \langle s \rangle = \sum_i f_i s_i. \]

(10.45)
probability of scattering in the layer. For thin samples of \(n\) layers, each of thickness \(x\), we have the relationship between numbers of layers and the bulk scattering coefficient, \(s\):

\[
\langle s \rangle x = np.
\] (10.46)

For consideration of multiple scattering, it is most convenient to work with the numbers of layers and scattering probabilities.

A challenge for multiple scattering calculations is the three-dimensionality of the sample. Neutrons can be scattered initially upwards, followed by a second scattering to the side, and a third scattering out to the detectors. A computer program such as MSCATT or MCViNE is required for accounting for such possibilities. Here we provide an approximate analysis in one dimension.

We assume that all neutrons entering the layered sample eventually leave the back of the specimen and are observed. The probability, \(p'\) for a neutron being scattered \(j\) times is the product of \(j\) of the layers doing a scattering, and \(n - j\) of the layers doing no scattering:

\[
p' = p^j (1 - p)^{n-j}.
\] (10.47)

We are not keeping track of which particular layer has done the scattering. We keep track of the fractions of neutrons that are scattered \(j\) times in our \(n\) layers when the scattering probability in each layer is \(p\). The number of ways of arranging the \(j\) scatterings over the \(n\) layers is therefore the binomial coefficient. The total fraction of neutrons that are observed in our example is 1, and should equal the sum of all scatterings, including \(j = 0\) for no scattering:

\[
1 = \sum_{j=0}^{n} \frac{n!}{(n-j)!j!} p^j (1 - p)^{n-j}.
\] (10.48)

We have implicitly assumed \(p \ll 1\), so the number of scatterings \(j \ll n\), and each layer scatters at most once. (This also ensures that the terms for which \(j \sim n\) are negligible in (10.48).) We can always satisfy this assumption \(p \ll 1\) by dividing our sample into finer and finer layers, because (10.46) shows us that for a fixed sample, \(np\) is a constant. Write out the first few terms of (10.48) in the binomial expansion:

\[
1 = (1 - p)^n + np(1 - p)^{n-1} + \frac{n(n-1)}{2} p^2 (1 - p)^{n-2} + \frac{n(n-1)(n-2)}{6} p^3 (1 - p)^{n-3} + ...
\] (10.49)

We simplify by using our condition of large \(n\) and small \(p\),

\[
1 = (1 - p)^n + (np)^1 (1 - p)^{n-1} + \frac{1}{2} (np)^2 (1 - p)^{n-2} + \frac{1}{6} (np)^3 (1 - p)^{n-3} + ...
\] (10.50)
The first term in (10.50) is the probability of zero scattering, the second is the probability of one scattering, the third – two scatterings, etc. We are interested in knowing the ratios of the different terms in this series. Since $1 - p \approx 1$, and using (10.46), the ratios are:

$$1 : sx : \frac{1}{2}(sx)^2 : \frac{1}{6}(sx)^3 \ldots$$  \hspace{1cm} (10.51)

The parameter $sx$ is dimensionless, and for experiment design we seek $sx \approx 0.10$. In other words, we seek a sample that scatters 10% of the incident neutrons. For this particular case, (10.51) gives the ratio of double scattering to single scattering of $1/2sx = 0.05$. Consider the effect of this double scattering on the inelastic spectrum. The scattering probability is the sum of an inelastic and elastic probability:

$$p = p_e + p_i,$$  \hspace{1cm} (10.52)

$$np = np_e + np_i,$$  \hspace{1cm} (10.53)

$$\langle sx \rangle = \langle s_e \rangle x + \langle s_i \rangle x.$$  \hspace{1cm} (10.54)

For this sample that scatters 10% of the incident neutrons, the ratio of the amount of inelastic single scattering to the amount of inelastic double scattering will be 5% – the same ratio as for the total scattering.

The effect of multiple scattering is to smear out the measured inelastic energy spectrum, and spread it out over twice the energy range of the single-scattering spectrum. Suppose the first scattering has the double-differential cross section of (6.52). Suppose a particular inelastic scattering causes an energy loss $\epsilon = \hbar \omega$. If this scattered neutron with altered energy and wavevector now undergoes a second inelastic scattering, the process may involve another double-differential cross section of (6.52), but with somewhat different parameters. An energy spectrum associated with this second scattering will be associated with every energy loss from the first scattering. Approximately, the energy spectrum for double scattering is the convolution of the single-scattering energy spectrum with itself. A similar type of argument applies to momentum distribution, but this is complicated by the vectorial aspects of the momentum transfer. Today the accepted practice is to ignore multiple scattering by assuming it is a weak background underneath the measured data, and does not contain significant structure. So long as it represents only 5% of the spectral area, this shouldn’t be a problem, should it?

A more insidious problem occurs when the sample is near other material that can scatter neutrons, such as a heater or a cold finger for sample temperature control. Depending on geometry, neutrons can be scattered from the sample, to this support material, to the detector. The result may be a new feature in the inelastic spectrum, such as a spurious peak from an excitation in the support material. What is so insidious is that by removing the sample, this spurious peak will disappear. It is therefore tempting to assign this peak to an excitation in the sample, not in the support material. This problem was discussed qualitatively in Sects. 10.10, and a specific example is calculated quantitatively in Sect. 12.3.4.
10.11. SAMPLE DESIGN

10.11.3 Absorption Criterion

When the sample contains a strong absorber of neutrons, the analysis is a bit more complicated than for (10.42). The loss of neutrons from the incident beam is now

\[ -dI(x) = (s + a) I(x) \, dx, \quad (10.55) \]

\[ I(x) = I_0 e^{-(s+a)x}. \quad (10.56) \]

where the parameter \( a \) is the absorption coefficient, causing neutrons to “disappear,” rather than scatter. The \( I(x) \) in (10.56) is the number of neutrons at depth \( x \) into the specimen, but Eq. 10.43 does not give the number of scattered neutrons. Furthermore, the scattered neutrons can be absorbed on their way out of the sample. With a one-dimensional model, for a sample of thickness \( t \), the path out has a length \( t - x \), so an exponential function with this argument attenuates the outgoing beam. For the scattered neutron intensity, \( I_{\text{scat}} \) from the increment \( dx \) is

\[ dI_{\text{scat}} = I(x) e^{-a(t-x)} \, s \, dx. \quad (10.57) \]

Substituting (10.56) into (10.57), rearranging, and integrating \( dI_{\text{scat}} \):

\[ I_{\text{scat}} = I_0 s e^{-at} \int_0^t e^{-sx} \, dx, \quad (10.58) \]

\[ I_{\text{scat}} = I_0 e^{-at} \left(1 - e^{-st}\right). \quad (10.59) \]

We can use (10.59) to calculate the scattering for any combination of absorption coefficient \( a \) and scattering coefficient \( s \). Suppose that \( a = 0 \), and there is no absorption. For a sample that scatters 10% of the incident neutrons, we obtain (10.43). Incidentally, for such a thin scatterer we can expand the exponential in (10.43) to obtain \( I_{\text{scat}} = I_0 s \, t \), which is perhaps more intuitive.

Now consider the other limit where absorption is strong. A sample that is too thick will have no scattered intensity, since all neutrons are absorbed. There is an optimal thickness \( t' \) for maximum scattering that we find by the analysis:

\[ \frac{dI_{\text{scat}}}{dt} \bigg|_{t'} = 0 \quad (10.60) \]

\[ t' = \frac{1}{s} \ln \left(\frac{s}{a} + 1\right). \quad (10.61) \]

For strong absorbers, \( a \gg s \), and we obtain from (10.61):

\[ t' = \frac{1}{a} \quad \text{optimal thickness for strong absorbers.} \quad (10.62) \]

10.11.4 Hydrogen Criterion

Finally, we consider the inelastic scattering from hydrogen, a unique element. Phonon scattering cross sections are proportional to \( a/m \). Hydrogen has a
uniquely large $\sigma$ and an uniquely low $m$, making it a stronger inelastic scatterer than Zr, by a factor of 1100. A trace of hydrogen in Zr would contribute a large amount of the inelastic scattering, and Zr has a tendency to absorb hydrogen and retain it in a modest vacuum. In many cases the spectral contributions from hydrogen are at high frequencies, and can perhaps be separated from the modes from Zr, for example. Nevertheless, it is important to know the hydrogen concentration in a sample for inelastic scattering experiments, and the best practice is to eliminate hydrogen from the sample.

### 10.12 Sample Design: Worked Example of LiFePO$_4$

Here are some sample calculations that were used for obtaining a thickness and mass of a typical sample for inelastic scattering. The neutronic properties of the elements of lithium iron phosphate are listed in Table 10.1.

The first step is to calculate the mass, $M$, of a sample that is a 10% scatterer. We do the calculation for 1 cm$^2$ of sample area, so the total cross section should be 0.1 cm$^2$

$$\sigma_{\text{tot}} = 0.1 \text{ cm}^2 = [1.37 + 11.62 + 3.31 + 16.93] \times [10^{-24} \text{ cm}^2] \left[ \frac{1 \text{ mole}}{158 \text{ g}} \times \frac{6.02 \times 10^{23} \text{ atoms}}{\text{mole}} \right] M \text{.} \quad (10.63)$$

$$M = 0.79 \text{ g} \text{.} \quad (10.64)$$

To fill the 5x5cm beam of the ARCS instrument, the sample mass should be 19 g.

The second check is for absorption. The total absorption for a 1 cm$^2$ sample is:

$$\sigma_{\text{abs,tot}} = [70.5 + 2.56 + 0.176] \times [10^{-24} \text{ cm}^2] \left[ \frac{1 \text{ mole}}{158 \text{ g}} \times \frac{6.02 \times 10^{23} \text{ atoms}}{\text{mole}} \right] 0.79 \text{ g} \text{.} \quad (10.65)$$

$$\sigma_{\text{abs,tot}} = 0.219 \text{.} \quad (10.66)$$

This is small enough to be ignored, but if it were much larger we would have to resize the sample to be consistent with (10.62), for example.
Finally, we consider the risk from some residual water in the sample. One H$_2$O molecule has a ratio of $\sigma_{\text{scat}}/m = 170$ for its hydrogen atoms. The $\sigma_{\text{scat}}/m$ in Table 10.1 are nearly 1000 times smaller. We should seek a mole fraction of water that is $10^{-4}$ or less. Perhaps some water could be removed by heating the sample slightly in the vacuum of the ARCS spectrometer.

Through data analysis procedures described in Chapter 6, it is possible to convert a measured inelastic spectrum to a representation of the phonon DOS. The problem of “neutron weighting” in LiFePO$_4$ can be seen from the last line in Table 10.1 in the values of $\sigma_{\text{scat}}/m$. For this material, the vibrational modes involving large amplitudes of motion for Li atoms will be much more prominent in the extracted DOS than the modes involving large amplitudes for O atoms. This neutron weight problem is one that can be addressed by lattice dynamics calculations, for example.

10.13 Sample Design: Sachets for Powders

Powder samples can be challenging to mount and accommodate in furnaces and cryostats. Here is an illustrated example of a successful design for a sachet (a small bag with a French name) filled with powdered SrO. It was designed to be heated to 1100°C in a vacuum furnace. Prime concerns were:

- The sample must not spill from the container. Spills of powder inside neutron instruments are serious issues, and require professional decontamination. After exposure to neutrons, many materials become “activated,” meaning that they are radioactive. If the powder is also chemically toxic, the spilled powder is called “mixed waste,” which is one of the highest categories of hazardous materials. Do not spill powders of samples used for neutron scattering research.

- The sachet should be made of thin sheet material so it does not contribute significantly to the inelastic scattering background. Neutron facilities often supply vanadium sample containers because these are well suited for neutron diffraction. Although vanadium has no coherent scattering (suppressing diffraction peaks), vanadium still has considerable inelastic scattering. For inelastic scattering, niobium can be a good choice for high temperatures, and aluminum is good for low temperatures.

- For experiments in furnaces, the sample should not react with the sachet at high temperatures. In some cases it is possible to examine a phase diagram to see if any compound made from elements in the container and elements in the sample has a low melting temperature. If so, the highest temperature of the experiment should be set with caution. It is usually a good idea to test the combination of the sachet and the sample in a furnace well before going to the neutron experiment.

- The sample thickness should be approximately uniform, and remain stable with handling and mounting. Shaping loose powders into flat plates
Figure 10.7: A partially-complete sachet. A sheet of thin niobium was folded in half, and the edges were folded multiple times in ever sharper triangle shapes. A pair of pliers will be used to crimp the edges.

is challenging, and a technique is shown below.

Figure 10.7 shows a sachet constructed of thin (50 micron) niobium sheet. The edges comprise many thicknesses of material, but they will be shielded from the incident beam. It is useful to draw the shape of the sample sachet in the lab notebook. An important trick shown in Fig. 10.7 is that the opening for the sample is formed around a plate of appropriate thickness. In this case a ruler proved the right thickness, and was used to shape the sachet.

Figure 10.8 shows that it usually takes two pairs of hands to fill the sachet with powder. It takes a bit of skill to ensure that the sachet does not bulge excessively in the middle, or to ensure that the powder does not settle to the bottom with further handling. Figure 10.9 is a reminder to weigh every sachet both before and after loading with powder.

Figure 10.10 shows a boron nitride frame around the sample sachet, which serves to block the incident beam from hitting the edges of the sachet. The frame should face the incident beam, looking upstream from the sample. For this experiment, the sample plane normal was tilted $+45^\circ$ with respect to the beam, so the normal to the sample pointed to detectors at $+45^\circ$, and the plane of the sample intersected detectors at $+135^\circ$, giving a low scattered intensity at this angle. It is also appropriate to adjust collimation slits in the incident beam to confine the height and width of the beam to the central region of the sample.

Further Reading

The contents of the following are described in the Bibliography.
Further Reading

Figure 10.8: Loading powder into the sachet. For valuable powders it is appropriate to do the loading over clean aluminum foil so that any spilled powder can be recovered and reused.

Figure 10.9: Remember to write the mass of the empty and filled sachet in the lab notebook.
Figure 10.10: Completed sample assembly, ready to mount in the neutron beam.
Further Reading


Chapter 11

Essential Data Processing

This chapter explains procedures for the reduction of data from a time-of-flight inelastic neutron spectrometer. To date the practice of reducing inelastic scattering data from time-of-flight instruments has been poorly-documented, dull, tedious, and error-prone. We hope this chapter transcends these traditional shortcomings.

The present chapter explains the rationale behind the data reduction modules of DANSE (Chapter 7 better explains their use). After an overview of a typical data reduction process, the individual steps are explained in detail. Approximately, analysis steps that are further from the raw data are more closely-related to the dynamic processes in the sample, and tend to be more computationally-intensive. These steps (such as corrections for absorption, multiple scattering, and multiphonon scattering) are described later in this chapter. Although the data analysis steps described first are somewhat independent of the previous chapters, understanding some of the later steps (such as corrections for multiphonon scattering) may be easier after reading Chapters 2 and 3.

The treatment in this chapter presupposes a “direct geometry” configuration (monochromation before the sample) and a two-dimensional, pixellated detector system, composed of an array of linear, position-sensitive detectors. This configuration is used in the Pharos and ARCS spectrometers, for example.

The first task is to take a raw set of data from an inelastic neutron scattering experiment (with a powder sample), and convert it into physical units such as meV and Å⁻¹ for energy and momentum transfer. Such basic steps are needed before the data can be compared to predictions from theory, or even compared to results from other inelastic spectrometers. Although the specific steps may depend in part on the scientific issues being studied, some data processing steps are common for nearly all experimental work, and are required to make an inelastic neutron scattering data set useful. There may be others steps as well, but most experiments follow most of the steps presented here.

Some notation is listed in Table 11.1. Fig. 11.1 shows the position of the detector tubes with respect to the sample, S, in the plane of the spectrometer,
Table 11.1: Notation

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$k_i$ ($k_f$)</td>
<td>modulus of the initial (final) neutron wavevector</td>
</tr>
<tr>
<td>$\vec{Q} = \vec{k}_i - \vec{k}_f$</td>
<td>momentum transferred from the neutron to the sample</td>
</tr>
<tr>
<td>$\phi$</td>
<td>scattering angle ($Q = \sqrt{k_i^2 + k_f^2 - 2k_i k_f \cos \phi}$)</td>
</tr>
<tr>
<td>$E_i$ ($E_f$)</td>
<td>incident (final) neutron energy</td>
</tr>
<tr>
<td>$\hbar \omega = \Delta E = E_i - E_f$</td>
<td>Energy transferred from neutron to sample</td>
</tr>
<tr>
<td>$l_2$</td>
<td>distance from the sample to the center of a detector</td>
</tr>
<tr>
<td>$d$</td>
<td>labels a specific detector</td>
</tr>
<tr>
<td>$h$</td>
<td>the height of a pixel in a detector</td>
</tr>
</tbody>
</table>

Fig. 11.2 the position perpendicular to the plane.

Typically one wants to know the dynamic structure factor, or scattering law, $S(Q, \omega)$. The coherent nuclear scattering of (7.94) can be summarized in simplified form:

$$\frac{d^2 \sigma}{d\Omega dE}_{\text{coh}} = \frac{\sigma_{\text{coh}}}{4\pi} \frac{k_i}{k_i} N S_{\text{coh}}(\vec{Q}, \omega),$$

(11.1)

with similar expressions for incoherent scattering or magnetic scattering. Owing to several complications originating with how the measurement is made, the measured data are actually:

$$\frac{d^2 \sigma}{d\Omega dH}_{\text{coh}} = \Phi \left[ \frac{\sigma_{\text{coh}}}{4\pi} \frac{k_i}{k_i} NS_{\text{coh}}(d, h, t) \times T(d, h, t) \times \epsilon(d, t) \right. \left. + S_{\text{nuis}}(d, h, t) + b(d, h, t) \right].$$

(11.2)

Here $\Phi$ is the incident flux, and often must be known by an independent measurement with a beam monitor, for example. The data are initially histogrammed as a function of neutron time-of-flight (TOF) rather than neutron energy, where the two are related by $E = 1/2m_n v^2 = 5.227(l/t)^2$ (the second equality holds for speeds in mm/µs or equivalent). Background $b$ and nuisance scattering from the sample, $S_{\text{nuis}}$, pollute the interesting signal.

Other factors modify the interesting term involving $S_{\text{coh}}$. The sample may absorb neutrons, so only a fraction $T$ of neutrons are transmitted through the

---

1. The introduction of new data acquisition hardware and software might alleviate this by histogramming directly into energy and momentum. That may simplify some tasks, for example eliminating the conversions from TOF to energy and from TOF and angle to momentum. However, this may complicate other tasks such as background subtraction.

2. This list includes anything that does not interest the experimenter. For example, in a phonon measurement this may include all magnetic scattering, multiphonon and multiple scattering effects.
Figure 11.1: Layout of the secondary flight path of a direct geometry time-of-flight spectrometer. The detectors are arrayed in a circle of radius $l_2$ around the sample (at $S$). The beam is incident on the sample from the left. The $x$ and $z$ axes show the coordinate system used in the discussion of binning into rings (Sect. 11.1.2), the $y$-axis of that system points out of the page. $\psi$ is the angle from the $z$-axis to the center of a given detector (in the $x$-$z$ plane).

Figure 11.2: Labeling some of the distances in the secondary flight path of a direct geometry time-of-flight spectrometer. The center of the detector is $l_2$ from the sample (at $S$), and a given pixel is at height $h$ from the center. The total distance from the sample to a pixel is $l_2 + b(h) = (l_2^2 + h^2)^{1/2}$. 
sample, and this varies with scattering angle and neutron energy. The efficiency $\epsilon$ of the detectors varies with energy and from one detector to another. The jobs ahead are:

- Account for incident flux.
- Remove background.
- Compute incident energy.
- Convert from time to energy.
- Correct for detector efficiency.
- Correct for absorption.
- Subtract similarly treated empty can data.
- Bin into rings of constant scattering angle $\phi$.
- Subtract additional nuisance scattering (multiple scattering and so on).
- Convert from angle to momentum.

Errors must be propagated. This is usually done according to a few simple rules, such as when data sets are added or subtracted, the errors add in quadrature; the error for the raw histogram is the square root of the counts in any given bin (156).

While many experiments could be treated by a similar process, not all of them can. In the later case, we have to know when departures from the routine require intervention in the reduction process. The sequence in which these tasks are presented here is that in which they are usually performed. As a rule, these operations do not commute, and we have to figure out what needs to be adjusted when re-ordering things. For instance, if the background is known as a function of time, either it should be subtracted before the data are transformed into energy, or the background must be transformed into energy as well. The sequence for other tasks, such as conversion to momentum units, is motivated by efficiency (one needs to think about energy when thinking about momentum, but not vice versa, so one might as well do the energy conversion first, followed by the momentum conversion).

### 11.1 Steps to Transforming Data into a Function of Energy and Momentum

#### 11.1.1 Operations and Data Structures

The basic data structure is a histogram. This implies sets of arrays: the array to store the histogram values *per se*, a similar array for errors, and associated arrays to store the axis values for each dimension.
Operations can be divided into those that change the structure of some of the arrays, and those that change only the contents of the arrays. Those that change the structure include energy rebin (§ 11.1.2), sum into rings (§ 11.1.2), and momentum rebin (§ 11.1.2); those that don’t change the array structures include subtracting various artifacts (§ 11.1.2, 11.1.2) and multiplicative corrections (§ 11.1.2, 11.1.2). The structure-changing operations require the creation of one or more new arrays; the rest can be done in place (assuming one is confident of the correction or if the correction can be easily undone). In the sections ahead, changes to the structure of the data will be noted.

11.1.2 A Closer Look at Each Task

Initial Data

The starting point is an array representing $I_0(d, h, t)$, the raw counts in each detector, at each detector position, for each TOF bin. The error is assumed to be the square root of the intensity: $\sigma_0 = \sqrt{I_0(d, h, t)}$. The TOF bins in this histogram usually have a constant time width, so all the bins are specified by some initial time $t_0$, the bin-width $\Delta t$, and the number $N$ of bins.

Initially there are five major arrays representing:

- The actual data. Three dimensional array; if there are $T$ time bins, $D$ detectors, and $H$ positions per tube, the array dimensions are $D \times H \times T$.
- The error. Same dimensions as the data.
- The time of flight. A one dimensional array of length $T$.
- The detector angles. A one dimensional array of length $D$.
- The pixel heights. A one dimensional array, length $H$.

Those arrays are stored in a histogram.

Some additional parameters are needed. These include the incident neutron energy, $E_i$, and the geometry of the instrument, especially the length $l_2$ from the sample to each detector, and the angle of each detector. We assume these additional parameters are known.

Normalize by Incident Flux

Typically one divides by the total counts $m$ in a peak in a beam monitor, which gives a proportional measure of the incident flux. Other measures may also be considered in unusual circumstances, for example, microamp-hours of proton

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3One might allow for the possibility that the pixel heights vary from one tube to another. The pixel position array becomes a two dimensional array, with size $D \times H$. 
beam delivered to the target. Regardless of the source, one still divides each element of the input array by some scalar:

\[ I_1(d_j, h_k, t_i) = I_0(d_j, h_k, t_i)/m, \]

for all \(i, j, k\), and similarly for the error.

### Subtract Background

Physically, the background is another complex subject; computationally, it is not. It can have different origins, from cosmic rays hitting the detectors (probably a small contribution) and neutrons from other experiments, to more serious causes, such as fast neutrons thermalizing in the instrument shielding, or slow neutrons scattering from poorly-masked beamline components. At times it can be difficult to determine what is background and what isn’t, especially when looking for diffuse scattering.

A common way to estimate the background is to subtract a constant (in time) background determined from the data. This may have some justification for removing the background that originates with radiations from neighboring instruments, or neutrons from far enough away that any time structure is lost. Another approach subtracts the scattering measured with an empty can. This is better done at a later stage, and is described below.

Because the background could vary with time and detector position, we denote it as \(b(d, h, t)\). Thus, we have

\[ I_2(d_j, h_k, t_i) = I_1(d_j, h_k, t_i) - b(d_j, h_k, t_i). \]

For each element of \(I_2\) one must know about one element of \(I_1\) and one element of \(b\), and the same for the error.

### Convert from Time to Energy

This step changes the structure of the data arrays. Prior to this step, the structures are the same as the initial arrays. There are still some more corrections to be made, but those corrections can be done more easily in terms of energy. This is computationally more intricate than the previous steps. In fact, it may be done in two steps.

First, one creates a new set of histogram bins whose boundaries are determined by \(t_2\), the sample-to-detector TOF, and \(l_2\), the distance from sample to detector, via the classical relation

\[ \hbar \omega = E_i - E_f = 5.227\left( v_t^2 - \left( \frac{l_2}{t_2} \right)^2 \right). \]

Note that pixels at different positions along the length of the detector tube are at different distances from the sample.\(^4\) The effect is to coarsen the energy

\(^4\)For a nominal \(l_2\) of 4 m (Pharos), this discrepancy gets as large as about 0.03 m (generally, the extra path length is \(b = l_2[(1 + \hbar^2/l_2^2)^{1/2} - 1] \approx \hbar^2/2l_2^2\)).
resolution. An extra 3 cm is a little more than the diameter of the detector tube, so this makes a contribution to the resolution similar to the size of the detector, or the sample size. For 35 meV neutrons and \( l_2 = 4 \text{ m} \) the difference in energy is about 1.5\%. One way to account for this is to write a separate set of histogram energy bin boundaries for each height in the detector tube:

\[
\hbar \omega = E_i - E_f = 5.227 \left[ \frac{v_i^2}{v_i^2} - \left( \frac{l_2^2 + h^2}{l_2^2} \right) \right].
\]  

(11.6)

In making this transition, we need to multiply by the time bin-width and divide by the energy bin-width:

\[
I_3(d, \hbar, \omega(h)) = I_2(d, h, t) \left| \frac{dt}{d\omega(h)} \right|.
\]  

(11.7)

At this step, the array describing the time bins has been replaced by a two-dimensional array describing the energy bins. Here is a list of the arrays at this stage of data reduction:

- Data array. Still \( D \times H \times T \).
- Error array. Still \( D \times H \times T \).
- Intermediate energy bin boundaries array. Two dimensional, \( H \times T \) or so.
- Detector angles array, one-dimensional, length \( H \). Same as before.
- Pixel position array. Same as before.

The energy binwidth is changing from bin to bin, which makes it hard to think about in plotting, fitting, etc. Also, the energy of any given bin is a function of the position of the pixel height. This is why we usually take the second step, rebinning into constant energy bins. Rebinning will also remove the artifact of each detector position having a unique set of energy bin boundaries.

For the second step one creates another set of histogram bin boundaries, spaced uniformly in energy. One then assigns counts from the old bins to the new bins by prorating them. The rules are simple: if an old bin lies entirely within a new bin, one puts all of the old counts into the new bin; in doing so, multiply by the old bin width and divide by the new bin width. If an old bin overlaps two new bins, assign counts to the first new bin based on the fraction of old bin overlapped by the first, and assign counts to the second new bin according to the fraction of the old bin overlapped by the second bin (and multiply by the old bin width while dividing by the new). The same holds true if the old bin overlaps many new bins. One can of course think of this the other way around: each new bin asks “what does each old bin owe me”.

\footnote{This is because the histogram reflects counts per some unit (\( \mu \text{s} \) or meV); we have the constraint that the \textit{integral} of the cross section be independent of the unit. If the histogram contains counts instead of counter-per-some-unit, then this factor is not necessary.}
One could write:

$$I_3(d, h, \omega) = \sum_i M(\omega, \tilde{\omega}(h)) \times I_3(d, h, \tilde{\omega}(h)) \times \left| \frac{d\tilde{\omega}}{d\omega} \right|, \quad (11.8)$$

where $M(\omega, \tilde{\omega}(h))$ is a matrix with the overlaps between old bins and the bins. One could get clever and drop the $|dt/d\tilde{\omega}|$ in Eq. 11.7 if one remembers to use $|dt/d\omega|$ instead of $|d\tilde{\omega}/d\omega|$ in Eq. 11.8. One typically needs to know something about several (but not all) elements of the old array before he can learn something about one element of the new array.

At the end of this step, three arrays have changed:

- Data array. Now $D \times H \times N_E$.
- Error array. Now $D \times H \times N_E$.
- Energy bin values array. One dimensional, $N_E$.
- Detector angles array, one-dimensional, length $H$. Same as before.
- Pixel position array. Same as before.

**Detector Efficiency**

Detector efficiency varies with energy. These data are provided by the manufacturer. One simply divides by a pre-existing array of numbers:

$$I_4(d_j, h_k, \omega) = \frac{I_3(d_j, h_k, \omega)}{\epsilon(d_j, \omega)}. \quad (11.9)$$

The efficiencies are typically known as a function of final neutron energy, so an additional step is required to convert $\epsilon(E_f)$ into $\epsilon(\omega)$.

**Absorption correction**

The absorption by the sample varies with both neutron energy and mean path length through the sample. For a single energy, the transmission through a material with unit cell volume $V$ and absorption cross section per unit cell $\sigma_{abs}$, the transmission probability is $T = e^{-l \sigma_{abs}/V}$. For a wise choice of units for $V$ and $\sigma_{abs}$, the path length $l$ will be in cm. The mean path length through the sample varies with angle, so one needs to compute $T(\phi, \omega)$.

In a real experiment, the length through the sample depends on where in the sample the neutron scatters, to which direction it scatters, and the change in the neutron energy (recall that the absorption cross section per unit cell varies as inversely with neutron speed, so when the speed changes, the probability of absorption per unit length changes). It does not depend on momentum transfer.

---

$^6$ $N_E$ is the number of new energy bins
11.1. STEPS TO TRANSFORMING DATA INTO A FUNCTION OF ENERGY AND MOMENTUM

One must calculate this dependence; let us assume this has been done. Then one divides each element of the old array by one element the array representing $T(\phi, \omega)\)

$$I_7(\phi_i, \omega_j) = I_6(\phi_i, \omega_j)/T(\phi_i, \omega_j) \quad \text{(11.10)}$$

Section 11.3 of this chapter takes a more detailed look at the absorption correction.

**Subtract empty can data**

Having accounted for absorption by the sample, it now makes sense to subtract the scattering measured from the empty can, if the data from the empty can can be brought through a similar treatment chain to this stage.

**Bin into Rings**

For a powder sample, the useful spatial information is the scattering angle $\phi$. That is to say, for a given $Q, E_i,$ and $\omega$, a powder scatters into a cone of angle $\phi$. So the two labels $d$ and $h$ can be condensed into the scattering angle $\phi$.

The first step is to identify the $\phi$ for each pixel. A conventional instrument coordinate system has the sample at the origin and the transmitted beam forming the $z$-axis. The $x$-axis runs horizontally from the sample (for Pharos, that’s toward FP-15) and perpendicular to the beam, and the $y$-axis, determined by right-handedness, points upward. (See Fig. 11.1.) It is natural to think of the detector as lying on a cylinder, with the axis of the cylinder coincident with the $y$-axis just described. So consider two vectors: $\vec{p}$, which runs from the sample to the pixel, and $\vec{a}$, with length $l_2$ running from the sample in the direction of the transmitted beam. In the instrument coordinate system, $\vec{a} = l_2 \hat{z}$, and $\vec{p} = -l_2 \sin(\psi)\hat{x} + h\hat{y} + l_2 \cos(\psi)\hat{z}$. $\psi$ is the angle between the transmitted beam and the center of the detector; it lies in the $x$-$z$ plane. The angle between $\vec{a}$ and $\vec{p}$ is given by $\vec{p} \cdot \vec{a} = |\vec{p}| |\vec{a}| \cos \phi = a_p z$:

$$\phi = \cos^{-1} \left( \frac{a_p z}{|\vec{p}| |\vec{a}|} \right) = \cos^{-1} \left( \frac{p_z}{|\vec{p}|} \right) = \cos^{-1} \left( \frac{l_2 \cos(\psi)}{\sqrt{l_2^2 + h^2}} \right)$$

$$= \cos^{-1} \left( \frac{\cos(\psi)}{\sqrt{1 + h^2/l_2^2}} \right). \quad \text{(11.11)}$$

Having found the angle $\phi$ for each pixel, an efficient method to combine pixels with similar angles is desirable. The simplest method uses three nested for

---

7Not an identical chain: hopefully your empty can didn’t need an absorption correction. (If it was made out of vanadium, it might need that!)
loops: one over the new angles, and two over the old detector indices. More abstractly:

\[ I_5(\phi, \omega) = \sum_{ij} N(\phi, d_i, h_j) \times I_4(d_i, h_j, \omega), \] (11.12)

where \( N \) is a matrix whose elements are 1 or 0 depending on whether \( \phi \) for the \( i^{th} \), \( j^{th} \) pixel is within the bounds defined for the \( k^{th} \) element of the new array. It may not be necessary to know all of the points of the old array in order to find one element of the new array.

This operation depends on the instrument configuration (where the detectors are) and the set of angular rings which can be defined by users. After this step, all but one of the arrays have changed. Here is a list of the arrays after this step of data reduction:

- Data array: \( N_\phi \times N_E \).
- Error array: \( N_\phi \times N_E \).
- Energy bin values array: \( N_E \).
- Phi-bin values array, \( N_\phi \)

**Subtract Nuisance Scattering**

Other nuisance scattering includes all scattering from the sample that is either non-interesting or confusing. This usually includes higher-order scattering processes, and is generally difficult to compute (especially since one has now applied a number of multiplicative factors which must be accounted for). Computationally, this is simply a matter of subtracting two arrays, element-by-element, which brings us to:

\[ I_6(\phi, \omega) = I_5(\phi, \omega) - S_{\text{nuis}}(Q, \omega) \] (11.13)

**Convert to Momentum Transfer**

Ideally, there would be a prorating scheme like that used in going between time and energy. Another approach frequently used is interpolation. Essentially one recognizes that the regularly-spaced angles at which one has measured are an irregularly-spaced array of momentum transfers, \( Q \). This momentum rebinning is typically performed with a convenient interpolation algorithm in, for example, IDL. Like the energy rebinning, one could write this process as a matrix operator acting on a vector:

\[ S(Q, \omega) = \sum_i M(Q, \phi_i, \omega_i) I_7(\phi_i, \omega_i) \] (11.14)

The arrays have changed again. After this step in data reduction, the arrays are:
• Data array: 2D, $N_E \times N_q$.
• Error array: 2D, $N_E \times N_q$.
• Energy bin values array: 1D, $N_E$.
• Q-bin values array: 1D, $N_q$.

At this point we have isolated the scattering of interest, and converted it into real physical units. This was considered a real achievement in the 1990’s. For some experimental work the data analysis can be declared complete, and graphs of the results prepared for publication.

### 11.2 Transformations and Information

Information is lost in the course of data reduction – the word “reduction” itself implies a loss. Some loss of information is both inevitable and necessary with data from a direct geometry chopper spectrometer – with $10^5$ detector pixels of $10^4$ time bins each, a data set of $10^9$ elements defies human comprehension. As described in the previous sections, the full experimental information on the arrival of each neutron at a specific detector pixel at a specific time is put through a number of transformations to improve comprehension, but also to improve counting statistics by summing events that are expected to be physically equivalent. It should also be noted that many data elements are usually uninteresting, or hold minimal counts, but even this information can be useful for removing extraneous scattering in the “good” data elements.

This section discusses the different physical scattering processes that can be measured experimentally, the possible transformations that can be performed on them, and whether these transformations are warranted, given the information required to do the transformation properly. For example, if the Q-resolution is lost by doing experiments on an incoherent scatterer like hydrogen, by performing measurements on a filter-difference spectrometer, or by rebinning all the Q-dependent data into energy, there is no path back to Q information by direct analysis of data alone. Fortunately, there may be a route back to the Q information through the use of a theoretical model or a simulation, and this is an opportunity provided by DANSE software.

Theory and computation play a bigger role when the experimental data require more supplementary information to enable the transformation. For data containing minimal information, it is possible to obtain detailed elementary excitations with theory alone – this is of course what has been done for many years by computational condensed-matter physicists. This dominance of theory over experiment was not the original intent of DANSE, but since DANSE provides such tools it will be interesting to see the path taken by future users of the DANSE system. Some community standards will have to be considered, but it is already clear that some transformations are better described as computational science rather than experimental science, even if experimental data is part of
the effort. For example, consider data on elastic incoherent scattering from a polycrystalline sample. Very little information is contained in such data on excitations in solids, but these data could be augmented by ab-initio theory to make predictions of the inelastic coherent scattering from a monocrystal. In this example most of the work would be done by theory, and the experimental data would add little value. For this reason, the DANSE software does not support this transformation. The present section lists all possible data transformations, and explains why approximately half of them are supported by DANSE.

11.2.1 Categorization of Transformations and Information

Table 11.2 lists the fundamental possibilities for a neutron scattering measurement. A real experiment may include combinations of coherent and inelastic scattering processes, but we consider these independently in what follows, because this is the approach used in data analysis. The categories of energy, momentum, and sample are orthogonal, and make the natural diagram of Fig. 11.3. Figure 11.3 is structured so that it the data of maximum detail is near the origin in the box labeled “inelastic coherent mono.” (Note: with the word “inelastic,” elastic scattering is also assumed included.) This inner cube contains enough information so that reprocessing allows one to move in any of the other three directions using experimental data alone (assuming the data are complete, of course).

As examples, consider the reprocessing of data from the inner box labeled “inelastic coherent mono”:

1. Averaging the scattering over all crystal orientations (from \( \vec{Q} \) to \( Q \)) allows reprocessing into the box above it, “inelastic coherent poly,”
2. Averaging over \( Q \) allows transformation to incoherent scattering, to the box in back, “inelastic incoherent mono,”
3. Discarding the inelastic contribution moves to the box at right “elastic coherent mono,” which is useful for single-crystal diffraction patterns.

These three statements are discussed further below, as are other transformation paths. Some are less obvious than others – statement 2 above is less obvious than 1 and 3, and it has been discussed in the literature. Nevertheless, it should be almost intuitive that considerable information is lost in going to the remote cube, “elastic incoherent poly.” Returning from such data to the origin is impossible.

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8Such analysis is possible by the ab-initio tools in DANSE, but DANSE does not offer this as a transformation of experimental data.
11.2. TRANSFORMATIONS AND INFORMATION

Table 11.2: Scattering Processes

<table>
<thead>
<tr>
<th>Energy</th>
<th>Momentum</th>
<th>Sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>inelastic</td>
<td>coherent</td>
<td>monocrystal</td>
</tr>
<tr>
<td>elastic</td>
<td>incoherent</td>
<td>polycrystal</td>
</tr>
</tbody>
</table>

Figure 11.3: Eight possibilities for scattering processes in a neutron experiment. Each of the eight can be calculated independently in a simulation or modeling process. Some can be obtained from another by transformation. In general, information is lost as one moves away from the origin, as explained in the text.
11.2.2 Coherent – Incoherent

Forward

An energy spectrum can be obtained by integrating over the \( Q \) coordinate of \( S(Q,E) \). With some care to account for thermal factors, multiple scattering, and multiphonon scattering, often before integrating over \( Q \), this can be converted to a phonon energy spectrum or a density of states. This approach is rigorous if all values of \( Q \) are accounted for, even if the sample is a coherent scatterer with sharp dispersion information. An integration over all \( Q \) accounts for all phonons, at least in principle. Some of the issues concerning the incoherent averaging over coherent scattering have been discussed in the literature:


Nevertheless, the forward transformation from “inelastic coherent” to “inelastic incoherent” can be done reliably, especially for data from direct geometry chopper spectrometers that provide a wide range of \( Q \). This transformation is offered by the DANSE software.

It should be mentioned that for many years, phonon DOS information has been obtained from phonon dispersion curves measured in special crystallographic directions on a triple-axis spectrometer. These dispersions are not good averages over \( Q \), and energies are not measured for all phonons in the solid. Nevertheless, after fitting the dispersion curves to a lattice dynamics model, the energies of all other phonons can be calculated. This approach has never been challenged for measurements on pure elements – it seems to work well because the high amount of information in the dispersion curves cause the interatomic force constants to be well determined. This method is not appropriate for disordered alloys, at least when the virtual crystal approximation is employed.

Reverse

In the same sense that it takes a lattice dynamics model to go from coherent inelastic scattering data from a triple-axis instrument to a good sampling of the incoherent inelastic scattering over all phonons, a model is always required for the reverse transformation. The positions of the Van Hove singularities in the phonon DOS are effective in constraining the dispersion curves, but there are
only three of them in a phonon DOS of a pure element. It is not yet clear how reliably the reverse transformation can be performed, but it is certainly a useful capability that is offered in the DANSE software.

11.2.3 Monocrystal – Polycrystal

Forward

Transforming a complete data set from a monocrystalline sample to a polycrystalline average is simple in principle, and has the most physical interpretation of all the transformations in this section. In essence, the single crystal data need to be averaged over all crystallographic orientations by “rotating” the monocrystalline data to produce a polycrystalline average. This amounts to transforming $\vec{Q}$ to $Q$. The results can be similar to the transformation of coherent to incoherent scattering, and in fact proves identical when the crystals have isotropic scattering. DANSE supports these transformations.

Reverse

Except in special cases, the reverse transformation from polycrystal data to monocrystal data requires a model. The question of importance is how reliably can the polycrystalline average of, for example phonon dispersions, be used to define the single crystal phonon dispersions along specific crystallographic directions. The answer is not fully known today, but in the case of crystals such as tungsten, which are elastically isotropic, it is expected that the amount of information is essentially the same for data from polycrystalline and monocrystalline scattering experiments. In the case of anisotropic crystals it is expected that the dispersive information in $S(Q,E)$ will have intensity variations from anisotropic effects, and perhaps important information can be obtained from the intensity variations within the dispersion curves if the data are of good statistical quality. DANSE supports these transformations, using both phenomenological and ab-initio models. It is hoped that the neutron scattering community will eagerly assess these capabilities of DANSE, because they free the experimenter from acquiring single crystals.

11.2.4 Inelastic – Elastic

Forward

A good diffraction pattern should be possible to acquire using white-beam mode on a direct-geometry chopper spectrometer, and the interpretation of the data would be the same as for a total scattering diffractometer. Here, however, we consider operation with a monochromatic beam as in the normal operation of a direct-geometry chopper spectrometer such as ARCS. The coherent elastic scattering is measured as part of the spectrum, so ignoring the scattering with energy transfers greater than the elastic peak width is essentially the same as
performing a diffraction experiment. There are some defects in this approach—the spread of incident wavelengths broadens the diffraction peaks in $Q$. Nevertheless, useful information about the sample can often be obtained by simply ignoring the inelastic part of the data. Transformations to do this are provided by DANSE.

**Reverse**

Taking elastic data and transforming it to inelastic data is not impossible. Unfortunately, direct transformations are not robust. Robust approaches require substantial additional information. Phonon effects are evident in pair distribution functions, but care must be taken to separate these from atomic size effect diffuse scattering, which can be of the same order. There have been some efforts to obtain phonon dispersions from diffraction patterns, and the reader is welcome to explore these:


At the inception of the DANSE project, the methods in these reference were placed outside the scope of the effort.

Nevertheless, DANSE does offer methods for reverse transformation. These are robust in that they give stable answers, but they require substantial computational effort. The approach is to obtain the lattice dynamics for the structure. From the lattice dynamics the inelastic scattering can be obtained. If the crystal structure or molecular structure is obtained from a diffraction measurement, for example, phonon dynamics can be calculated in one of two ways.

- A model of the lattice dynamics for the structure can be used to calculate the phonons. This requires input on interatomic force constants. They may be guessed, or obtained from other sources. From the dynamics, the inelastic scattering can then be calculated.

- Using the unit cell obtained from a diffraction pattern, or a molecular structure obtained by other means, an ab-initio electronic structure calculation can be performed to optimize the positions of the atoms in the structure. These may not be the real atom positions, but relaxed positions are needed for the following computational step. Next, the atom positions are displaced by a controlling software package such as PHON, and
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the ab-initio code is run again for the distorted unit cell. This permits calculation of tensorial force constants that are used in the lattice dynamics. From the dynamics, the inelastic scattering can be calculated.

Without such theoretical input, however, attempting a reverse transformation from elastic to inelastic would not be robust, and with the DANSE software it is an error to do so. Note that the transformation of structural information into inelastic inelastic scattering information requires perhaps the most sophisticated tools offered by DANSE. An increasing loss of experimental information requires an compensating sophistication in the theoretical tools to reverse the situation.

11.2.5 All Specific Cases

The previous subsections discussed general issues when transforming data forward or backwards between elastic and inelastic, coherent and incoherent, and monocrystal and polycrystal. For each of these three cases, and six general transformations, there are four specific cases. For example, Fig. 11.3 shows that between the monocrystal and polycrystal samples (up in the graph), there are a total of eight forward and reverse transformations:

- inelastic coherent (mono\leftrightarrow poly),
- inelastic incoherent (mono\leftrightarrow poly),
- elastic coherent (mono\leftrightarrow poly),
- elastic incoherent (mono\leftrightarrow poly).

Similar forward and reverse transformations between adjacent cubes are also possible along the elastic\leftrightarrow inelastic and coherent\leftrightarrow incoherent directions of Fig. 11.3, giving a total of 24 such transformations between adjacent cubes in the figure (along the \langle100\rangle directions). In addition, a total of 12 transformations are possible between scattering data in cubes in Fig. 11.3 that are not immediately adjacent (along the \langle110\rangle directions), and a total of 8 possibilities for transformations between cubes located diagonally in the figure (along \langle111\rangle directions). Tables ?? – ?? list all 44 possible transformations, and identify the specific transformations supported by the DANSE software. As of early 2007, the plan for DANSE supports 24 of these possible transformations.

Sections 11.2.2 - 11.2.4 provided general reasons for the why transformations along the main axes of Fig. 11.3 are not provided by DANSE. Eight transformations are possible along each of these axes, however. Nearly all cases in Table ?? can be understood with the rules of Sects. 11.2.2 - 11.2.4, with the following exceptions:

- Transformations from elastic coherent polycrystal data (powder diffraction patterns) to elastic coherent monocrystal data (single crystal diffraction data) are not supported in the rebinning operation in the first release of DANSE. These capabilities are possible with Rietveld refinement.
methods (or PDF modeling), which are available as methods outside the scattering kernel.

- Elastic incoherent scattering pertains to the isotopic incoherence of neutron scattering, for example, and not to incoherence from atomic disorder (which is treated in methods for elastic coherent scattering). DANSE therefore does not support well transformations involving elastic incoherent scattering from either monocrystals or polycrystals.

The more complex transformations of two or three data characteristics, listed in Tables ?? and ??, are generally consistent with the previous rules and individual transformations. More transformations are possible by chaining individual transformations from Table ?? than are listed in these Tables ?? and ?? . These latter tables list algorithms that do both transformations in an integrated way. The emphasis is on data from inelastic scattering, since this field was first to exploit this method of direct experiment simulation.

11.3 Absorption

With an absorbing sample, one observes \( \frac{d^2\sigma}{d\Omega d\omega} \times T(\Omega, \omega) \) rather than just \( \frac{d^2\sigma}{d\Omega d\omega} \), where \( T(\Omega, \omega) \) is the probability of a neutron being transmitted through the sample in a given direction (described by the solid angle \( \Omega \)) and with a given final energy (final energy = incident energy − energy transfer, or \( E_f = E_i - \hbar\omega \)). All that needs to be done is to calculate the transmission for each final energy and for each detector, and then divide the observed scattering by the transmission probability. For an experiment in which all the observed neutrons scatter once, the calculation is straightforward, though somewhat computationally demanding.

We suppose that the sample is divided into a number of cells of equal volume; then a neutron observed in any given detector with a given final energy was scattered in any cell with equal probability. We compute the transmission probability for each scattering cell, then average over probabilities for all cells. The reason we have to be so careful is that the transmission probability depends on the distance the neutron travelled through the sample, and this depends on where in the sample the scattering occurred, and the outgoing direction of the neutron\(^9\).

For a neutron travelling across an absorbing medium, the probability of transmission is \(^{10}\)

\[
T = \frac{N}{N_0} = \exp(-l\sigma_{abs} \rho).
\]

(11.15)

Here, \( l \) is the length of the path through the sample, \( \sigma_{a} \) is the absorption cross section, and \( \rho \) is the number density of absorbers. If the neutron changes direc-

\(^9\)We assume the incoming neutrons are perfectly monochromatic and collimated.

\(^{10}\)Think \( dN/dx = -\sigma p \ldots \)
11.4. NEUTRON WEIGHTING

When a neutron scatters while traveling through the sample, the probability becomes

\[ T = \exp(-l_{inc}\sigma_{\text{abs}}\rho - l_f\sigma_{\text{abs}}\rho) \quad (11.16) \]

where \( l_{inc} \) is the path length from where the neutron enters the sample to the scattering spot, and \( l_f \) is the path length from the scattering spot to where the neutron exits the sample. There's one more difficulty: the absorption cross section depends on the neutron energy, so if the scattering is inelastic, there will be two absorption cross sections, \( \sigma(\omega_{inc}) \) and \( \sigma(\omega_f) \), and we have the transmission probability as

\[ T = \exp(-l_{inc}\sigma_{\text{abs}}(\omega_{inc})\rho - l_f\sigma_{\text{abs}}(\omega_f)\rho). \quad (11.17) \]

In what follows, we assume that the scattering is purely horizontal. This is an approximation that greatly reduces complexity of the computation. This is justifiable for several existing instruments including LRMECS and PHAROS. In fact, any horizontal scattering lies within about \( \pm 5^\circ \) of the horizontal on an instrument like Pharos,\(^{11} \) and is therefore negligible.\(^{12} \) This means each scattering cell needs two labels.

In considering a number of cells, the incoming path length depends on where the cell is located. The outgoing path length depends on where the cell is, and in what direction the neutron is heading. This direction can be determined if we know the detector \( d \) that the neutron is hitting. So we need four labels for the transmission from a given scattering cell to a given detector with a given final energy:

\[ T(d, \omega_f, i, j) = \exp(-l_{inc}(i, j)\sigma_{\text{abs}}(\omega_{inc})\rho - l_f(i, j)\sigma_{\text{abs}}(\omega_f)\rho). \quad (11.18) \]

In the last step, we’re going to average over all \( N \) cells for each detector and energy:

\[ T(d, \omega_f, i, j) = \frac{1}{N} \sum_{i,j} \exp(-l_{inc}(i, j)\sigma_{\text{abs}}(\omega_{inc})\rho - l_f(i, j)\sigma_{\text{abs}}(\omega_f)\rho). \quad (11.19) \]

It does not get much easier than that. Now all we need is a program that computes Eq. 11.19 for a number of interesting geometries.

### 11.4 Neutron Weighting

The phonon DOS obtained by the data reduction methods of this chapter is not the true phonon DOS, except in the case of pure elements. It is instead

\(^{11}\)the detectors are at most 0.5 m above the scattering plane, while the middle of the detector is 4 m from the sample, so the maximum angle is about \( \pm 7^\circ \). However, the top and bottom 10 cm or so of the tubes are thrown out...

\(^{12}\)This assumption has not been sufficiently examined. Unless the effect is very small (\( \ll 1\% \)), it should be included in the calculation when that becomes computationally affordable. On Pharos, the extra path length is about 0.5%. Is this important? For instruments with larger vertical divergence (ARCS), this will be necessary anyway, so might as well get used to it.
the “neutron-weighted DOS.” The neutron-weighted DOS accounts for the different efficiencies of phonon scattering from different chemical elements or isotopes. (If the sample is chemically homogeneous, this is usually not a problem.) The neutron-weighted phonon DOS is rigorously defined (Sect. 6.3.3) as:

\[ g_{NW}(E) \propto \sum_d g_d(E) \exp(-2W_d) \exp(2W) \frac{\sigma_d}{m_d} \]  

(11.20)

where \( \exp(-2W_d) \), \( \sigma_d \) and \( m_d \) are the Debye-Waller factor, total scattering cross-section and mass of atom d. The Debye-Waller factor is an explicit function of \( g_d(E) \). The term \( \exp(2W) \) is the average Debye-Waller correction; this is calculated from the self-consistent neutron-weighted DOS. The factor \( \exp[2(W - W_d)] \) is approximately unity. For the case where \( \sigma_d/m_d \) is the same for all species \( d \), \( g_{NW}(E) \approx g(E) \).

Obtaining the true phonon DOS from the neutron-weighted phonon DOS requires a full analysis of the lattice dynamics. This can be performed by simulational procedures described in a later chapter. The neutron-weight correction as well as other approximations involved in the data analysis can be overcome by fitting a dynamics model to the neutron-scattering data directly. Although this approach is both scientifically and computationally demanding, we foresee no better method for extracting the vibrational dynamics from inelastic neutron scattering measurements.

### 11.5 Calculation of Multiphonon Scattering

The multiphonon expansion was developed with some rigor in Section 7.3.2. Here we explain how it works in practice. The total measured spectrum, \( S(E) \), is the sum of components, \( \sum_{n=0}^\infty S_n(E) \), from neutrons scattered after creating different numbers, \( n \), of phonons in the sample.\(^{13}\) Only the 1-phonon scattering is useful for obtaining a phonon DOS, so it is important to have an understanding of the higher-order terms for planning an experiment, or if one seeks quantitative corrections of experimental data. Performing a multiphonon expansion is done in two steps.

- First, the weights of the different \( n \)-components are calculated independently with input information on \( Q \) and atom displacements \( u \) at the temperature \( T \).
- Second, the spectral shape of each component \( S_n(E) \) is obtained by sequentially convoluting the 1-phonon profile with itself a total of \( n - 1 \) times.

Finally, for a chopper spectrometer, detectors at different scattering angles \( \phi \) provide energy spectra \( S_\phi(E) \) where \( Q \) varies with \( E \) across the spectrum. The

\(^{13}\)Phonon annihilation is handled by extending the range of \( E \) to negative numbers for each \( S_n(E) \).
11.5. Calculation of Multiphonon Scattering

Multiphonon corrections must take into account the kinematical relation of $Q$ vs. $E$ at each scattering angle $\phi$ if data are analyzed from each detector bank. The constant-$Q$ multiphonon analysis is simpler to understand, and is applicable for data from triple-axis spectrometers, or for data from chopper instruments that have been reduced to $S(Q, E)$.

Intensities of $n$-Phonon Spectral Components

Start with the Debye–Waller factor, which attenuates the elastic scattering $S_0$, to a fraction of the total scattering $S$:

$$S_0 = S \exp(-2W), \quad (11.21)$$

which we rearrange and expand:

$$S = S_0 \exp(+2W), \quad (11.22)$$

$$S = S_0 \sum_{n=0}^{\infty} \frac{(2W)^n}{n!}. \quad (11.23)$$

We now have a series of terms in an expansion, but the next step of substituting (11.21) into (11.23) amounts to nothing more than writing $\exp(-2W) \exp(+2W) = 1$:

$$S = S \exp(-2W) \sum_{n=0}^{\infty} \frac{(2W)^n}{n!}. \quad (11.24)$$

$$1 = \sum_{n=0}^{\infty} \frac{(2W)^n}{n!} \exp(-2W). \quad (11.25)$$

The terms in (11.25) are associated as 0 for elastic scattering, 1 for 1-phonon scattering, 2 for 2-phonon scattering, etc. What is special about (11.25) is that the $n^{th}$ term in the series is exactly the fraction of the $n$-phonon spectral component in the total scattering (elastic plus inelastic).

To calculate the fraction of any multiple phonon term (e.g., the amount of 2-phonon scattering), all we need is $2W$. Recall the physical origin of $2W$:

$$2W = \langle Q^2 u^2 \rangle, \quad (11.26)$$

where $\langle u^2 \rangle$ is the mean-squared atom displacement. Equation (11.26) shows that $\exp(-2W)$ becomes much less than 1 when the atom displacement becomes comparable to $1/Q \sim \lambda/2\pi$, the number of wavelengths associated with the scattering angle. This is consistent with $2W$ originating from the destructive interference of scattered wavelets. The $Q^2$ and $u^2$ are related to the energy of recoil, and the thermal energy, respectively. The recoil energy, $E_R$ is

$$E_R = \frac{\hbar^2 Q^2}{2M}, \quad (11.27)$$
where $M$ is the mass of the atom that is scattering, in units of the neutron mass. (The units of $M$ are very similar to the atomic weight.) The thermal energy is:

$$k_B T = \frac{1}{2} M \omega^2 \langle u^2 \rangle$$

(11.28)

for one mode of frequency $\omega$, and is a quantity that depends on the phonon DOS for a real material. Sadly, an exact evaluation of (11.28) is not simple, in part because the phonon states are not fully occupied at modest temperatures, but also because of the commutation relationships discussed in Section 7.2. The result is summarized as $\gamma_0$, where:

$$\gamma_0 = \int_0^\infty \coth(E/2k_B T) \frac{g(E)}{E} dE .$$

(11.29)

In the limit of high temperatures, where $\coth(E/2k_B T) \to 1$, it is clear from (11.29) that $\gamma_0$ increases as $1/E$ (because the amplitudes of atom motions become larger in low-energy modes as predicted by (11.28)). The low-energy modes are even more important at low temperatures, where they have a larger phonon occupancy. The Debye–Waller factor of (11.26) is, rigorously:

$$2W = \gamma_0 E_R ,$$

(11.30)

In performing a multiphonon expansion for one $T$ and one phonon DOS $g(E)$, the value of $\gamma_0$ is computed once with (11.29). The $2W$ from (11.30) for appropriate $Q$ is used in (11.38) to get the fractions of all $n$-phonon scatterings. We therefore rewrite (11.25) as

$$S(Q) = \sum_{n=0}^\infty S_n(Q) , \text{ where :}$$

$$S_n(Q) = \frac{(2W(Q))^n}{n!} \exp(-2W(Q)) .$$

(11.31)

(11.32)

**Shapes of $n$-Phonon Spectral Components**

It remains to get the spectral shape of each order of the multiphonon scattering. The energy spectrum for 1-phonon scattering was discussed in Sections 6.3 and 7.3.2. The spectrum for one-phonon scattering weights more heavily the low-energy modes because they have larger amplitudes of motion, providing a factor of $g(E)/E$. The number of phonons is the Planck distribution $1/\{\exp(E/k_B T) - 1\}$, so the two factors provide the shape of the 1-phonon profile, $A_1(E)$:

$$A_1(E) = \frac{g(E)}{E \gamma_0} \frac{1}{e^{E/k_B T} - 1} .$$

(11.33)
11.5. CALCULATION OF MULTIPHONON SCATTERING

Each phonon created has the profile $A_1(E)$ of (11.33). When two phonons are created simultaneously, the total spectrum of energies is the convolution of the 1-phonon profile with the 1-phonon profile. The 2-phonon spectrum is:

$$A_2(E) = A_1 * A_1 = \int_{-\infty}^{\infty} A_1(E - E') A_1(E') \, dE' ,$$  \hspace{1cm} (11.34)

and the $n$-phonon profile is the convolution of another 1-phonon profile with the $(n-1)$-phonon profile:

$$A_n(E) = A_1 * A_{n-1} = \int_{-\infty}^{\infty} A_1(E - E') A_{n-1}(E') \, dE' .$$  \hspace{1cm} (11.35)

Starting with $A_1$, we can generate the spectral shapes of all the orders of multiphonon scattering by the systematic application of (11.35). The total scattering is the sum of these spectral profiles, weighted by the corresponding terms of (11.31):

$$S(Q, E) = \sum_{n=0}^{\infty} S_n(Q) A_n(E) ,$$  \hspace{1cm} (11.36)

$$S(Q, E) = \sum_{n=0}^{\infty} \frac{(2W)^n}{n!} \exp(-2W) A_n(E) .$$  \hspace{1cm} (11.37)

If we define each term in the sum as $S_n(E)$:

$$S(E) = \sum_{n=0}^{\infty} S_n(E) .$$  \hspace{1cm} (11.38)

Examples of Multiphonon Scattering from a Chopper Spectrometer

The analysis presented here is for inelastic energy spectra obtained from a group of detectors centered about the scattering angle, $\phi$. The energy transfer, $E$, is obtained simply and reliably from the arrival times of the neutrons at the detector. On the other hand, for each detector bank at $\phi$, the value of $Q$ varies with $E$ and the energy of the incident neutron $E_{\text{inc}}$ as:

$$Q[\text{Å}^{-1}] = 0.6947 \sqrt{2E_{\text{inc}} - E - 2 \sqrt{E_{\text{inc}}(E_{\text{inc}} - E) \cos \phi} ,$$  \hspace{1cm} (11.39)

where $E$ and $E_{\text{inc}}$ have units [meV]. Curves showing relationships $Q(E)$ are presented in Figure ???. The curves cannot extend to the right beyond $E_{\text{inc}}$.

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14 Consider each phonon excitation to be a random variable with a probability distribution of $A_1$. The sum of two random variables has a distribution that is the convolution of the probability distributions $A_1 * A_1$. 
because the incident neutron cannot lose a greater amount of energy to the sample. Notice that the curves are generally asymmetrical in ±E.

For calculating the multiphonon scattering at a particular scattering angle \( \phi \), it is necessary to evaluate the 2W and the Debye–Waller factor at each energy of the spectrum, because \( Q \) varies with \( E \) as in (11.39) and as shown in Figure ??a. Examples of such calculations are shown in Figure ??b. Figure ??a shows the phonon DOS curve from fcc nickel metal from which the subsequent curves were calculated. Figure ??b shows the total inelastic scattering calculated for a temperature of 500 K.\(^\text{15}\) The 1-phonon scattering is confined to the range ±37 meV, since this is the maximum phonon energy. With the excitation of two phonons, the energy range can be extended to \( 2 \times \pm 37 = \pm 54 \text{ meV} \), and this is the energy range of the 2-phonon scattering. In Figure ??b, the intensity at 40–50 meV is almost entirely 2-phonon scattering.

Notice the general trend in Figure ??b showing that the inelastic scattering increases with the scattering angle, owing to the factor \( 2W \) in (11.32). On the other hand, the scattering is suppressed by the Debye–Waller factor, \( \exp(-2W) \). At high \( Q \) or high \( T \) the Debye–Waller factor becomes increasingly important, and effects of this are seen in Figure ??c. First notice that although there are about twice as many phonons at 1000 K as at 500 K, the inelastic scattering of Figure ??c is nowhere near twice as large as in Figure ??b. Also notice the change in symmetry of the spectra in Figure ??c with detector angle. For a detector angle of 60°, Figure ??c shows an inelastic scattering that is approximately symmetrical in ±E. This is consistent with the gradual variation of \( Q \) with \( E \) in the kinematics as shown in Figure ???. On the other hand, for the larger detector angles of 90° and 140°, the value of \( Q \) is larger for \(-E\) than \( +E\). This asymmetry causes the Debye–Waller factor to attenuate the scattering at \(-E\) more severely than at \(+E\). When these effects are significant, it is obviously naive to interpret the asymmetry in \( E \) of the inelastic scattering in terms of detailed balance.

### 11.5.1 Multiphonon Correction – Iterative

The present section describes a correction procedure that can be used to remove higher-order multiphonon scattering from an experimental inelastic scattering spectrum, so the spectrum can be used to deduce a 1-phonon profile and a phonon DOS. It is based on the incoherent approximation, but this procedure has been used for many years with coherent multiphonon scattering data, and it is often expected to be reliable in such cases, at least for polycrystalline samples. (The idea is that averaging over crystalline orientations provides a broad sampling of reciprocal space, performing a good average over phonon modes. The incoherent approximation is adequate when all modes are sampled

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\(^{15}\)An experimental spectrum would contain a large peak at \( E = 0 \) from the elastic scattering. In principle, the area of this elastic peak would be the factor \( \exp(-2W) \) of the total integrated scattering (elastic plus inelastic).
this way. It is unlikely that this approach could be reliable for the analysis of coherent scattering from single crystals, of course.)

An example of multiphonon calculations and an extraction of 1-phonon scattering is shown in Figure ???. Cerium is a coherent scatterer, and the data were acquired at several values of \( Q \) to average over the different phonon modes. An inelastic spectrum for one value of \( Q = 3.924 \text{ Å}^{-1} \) is presented in the figure.\(^{16}\) Steps of an iterative procedure to refine the phonon DOS are shown in Figure ???. The first step of the iteration used a very crude approximation for the multiphonon scattering plus random background – a simple constant function of approximately 70 counts. After this constant was subtracted from the experimental data, the DOS curve, \( g(E) \) labeled “1” in Figure ??b was obtained from the 1-phonon profile from (11.33) with a Debye–Waller factor of (11.32). The 2–5 phonon scattering was calculated from this first iteration of a phonon DOS, scaled in height (plus a constant background was also fit to the data), and a second iteration of the DOS was generated by the same procedure. Notice that the second and third iterations of the phonon DOS in Figure ??b are rather similar. Even for this difficult case where the 2–5 phonon scattering is a large fraction of the total inelastic scattering, convergence is fairly quick because the 2–5 phonon scattering does not have much structure.

The multiphonon expansion assumes that the scattering is incoherent. Unfortunately, a better approximation would require a detailed simulation of the lattice dynamics to account for the \( \vec{Q} \)-dependence. To our knowledge, such a calculation has not been done as of 2004. Another case where the assumptions of this procedure are stretched is the application of the incoherent multiphonon correction procedure to inelastic spectra from alloys. One concern about alloys is that the Debye–Waller factors differ for the different atoms in the alloy, causing some phonons to be weighted more than others. Although the multiphonon correction procedure has been used without adapting to the different \( \langle U^2 \rangle \) of the atoms in the alloy, comparisons with other measurements have shown that the procedure is often acceptable, perhaps because the multiphonon scattering is essentially featureless.

11.5.2 Multiphonon Correction – Fourier Log

Another way to correct for multiphonon scattering is possible for energy spectra at constant \( Q \). Following the notation of Section 11.5, for fixed values of \( Q \) and temperature, the energy dependence of the various phonon contributions, \( S_n(Q, E) \), is

\[
S(Q, E) = \frac{(2W)^n}{n!} e^{-2W} A_n(E),
\]

(11.40)

\(^{16}\)A spectrum at one value of \( Q \) can provide a piece of the dynamical structure factor intensity, which we loosely call a “phonon DOS,” since the result would be a phonon DOS if the scattering were incoherent. Data were acquired from the triple-axis spectrometer HB3 at HFIR.
The measured scattering intensity, $S(Q, E)$, is the sum of intensities from all phonon processes:

$$S(Q, E) = \sum_n S_n(Q, E), \quad (11.41)$$

Our problem is to use the measured $S(Q, E)$ to isolate the single scattering profile, $A_1(E)$. An approach to extracting the phonon partial DOS from experimental data is called the “Fourier-log deconvolution method.” It was used for analysis of plasmon scattering measurements in electron energy-loss spectrometry [Spence]. This method is also used for incoherent inelastic phonon scattering [M. Y. Hu, W. Sturhahn, et al.]. It has the additional feature of being able to correct for the broadening of the elastic line, but only if the energy resolution is constant across the energy spectrum (not the case in TOF spectrometers). The zero-loss peak, $Z(E)$, is convoluted with the scattering from the specimen as:

$$S(Q, E) = Z(E) \ast \left( e^{-2W\delta(E)} + \sum_n S_n(Q, E) \right), \quad (11.42)$$

where the first term in (11.42) is the zero-loss peak, reduced by the Debye–Waller factor. Taking the Fourier transform, $F[]$, of (11.42) simplifies the convolutions of (11.35), which become multiplications in Fourier space:

$$F[S(Q, E)] = F[Z(E)]e^{-2W} \left( 1 + \sum_n \frac{(2W)_n^n}{n!} (F[A_1(E)])^n \right). \quad (11.43)$$

The term in parentheses () on the right side of (11.43) is recognized as the expansion of an exponential function:

$$F[S(Q, E)] = F[Z(E)]e^{-2W} \left( e^{2WF[A_1(E)]} \right). \quad (11.44)$$

Taking the logarithm of (11.44) and rearranging:

$$F[A_1(E)] = \frac{1}{2W} \ln \left( \frac{F[S(E)]}{F[Z(E)]} \right) + 2W. \quad (11.45)$$

The inverse Fourier transformation, $F^{-1}[]$, provides the single scattering profile, $A_1(E)$:

$$A_1(E) = \frac{1}{2W} F^{-1} \left[ \ln \left( \frac{F[S(E)]}{F[Z(E)]} \right) \right] + 2W\delta(E). \quad (11.46)$$

If we do not care about the normalization of the single scattering profile, and if we delete the zero-loss peak $Z(E)$ from the data (it is suppressed by the thermal correction anyway), we obtain:

$$A_1(E) = \frac{1}{2W} F^{-1} \left[ \ln \left( \frac{F[S(E)]}{F[Z(E)]} \right) \right]. \quad (11.47)$$
11.6. Corrections for Simultaneous Multiple Scattering and Multiphonon Scattering

11.6.1 Best Practices

Recursive computation

Multiple-scattering is naturally computed with a recursive algorithm, as implemented in MCViNE (102) or MCSCAT (72). For the purpose of this discussion, we differentiate between two types of multiple scattering: single-scatterer multiple scattering (SSMS) to describe the multiple scattering within one neutron scatterer and multiple-scatterer multiple scattering (MSMS) to describe multiple scattering among different neutron scatterers, such as between the sample and its holder. A neutron may scatter by different processes, and it may undergo these processes in different sequences and in different locations. The set of scattering processes (implemented as “scattering kernels”) used in MCViNE in 2014 is

- Incoherent elastic scattering
- Coherent elastic scattering from powder sample
- Incoherent inelastic single-phonon scattering
- Coherent inelastic single-phonon scattering from a powder sample
- Multi-phonon scattering
- Scattering from a dispersion surface where the dispersion relation and the dynamical structure factor are described by analytical functions of momentum transfer vector \( \mathbf{Q} \).

Figure 2 shows an example of SSMS in which a neutron is scattered three times before exiting a scatterer. When a neutron is scattered inside a homogeneous scatterer, the original incident neutron is split into two for computational efficiency. One neutron is propagated through the scatterer with its probability lowered by attenuation, while the other is scattered by one of the scattering kernels, chosen by random selection, at a point also randomly selected along the forward path of the incident neutron. This splitting process repeats for several scatterings, as illustrated in Fig. 2, until either the neutron probability is lower than a pre-selected limit, or the maximum order of multiple scattering is reached. At this point, the neutron that is still inside the scatterer is allowed to propagate out, with its probability attenuated appropriately.

Footnote: Sample components in some linear-chain-based MC ray tracing packages can support SSMS, but they do not have abstractions similar to composite scatterer or scattering kernel. As a result, the SSMS algorithm must be duplicated in these sample components.
Figure 3 shows an example of multiple scattering in a sample assembly with a simple “concentric” arrangement of the sample and the sample environment. Only five of an infinite number of possible multiple scattering paths are illustrated, and the splitting processes are not shown.

**Multiple scattering from an aluminum sample and environment**

In this example, we present experimental and simulated inelastic spectra from polycrystalline Al metal. Most interesting, perhaps, are the simulations that include subsets of the scattering processes. These can show how to optimize the experiment, or how to do the most important corrections for unwanted intensities. The experiment and simulations used a 60 mm × 60 mm × 4 mm polycrystalline aluminum (1100 alloy) plate in the ARCS spectrometer. A quick calculation using the total scattering cross section for Al shows that the sample scatters 5% of the incident beam. In the simulation and experiment, the sample was placed at approximately 135° to the beam direction, and the incident energy was tuned to 80.5 meV using a Fermi chopper slit package at 480 Hz. The simulated sample assembly contains only one homogeneous scatterer for the aluminum plate. Different simulations made use of different combinations of scattering kernels. All phonon-related scattering kernels used phonon energies and polarization vectors computed on a regular grid in a Brillouin zone from a BvK model [67].

Figure ?? displays different I(Q, ω) for the aluminum plate. The experimental result is in panel (a) and panels (b)–(e) show simulated data. In (b) only the incoherent elastic and incoherent single phonon scattering are included (The intensity was enhanced to show detail – the values on the colorbar give an indication of the scaling). In (c) only the coherent elastic (powder diffraction) and the coherent single-phonon inelastic scattering are included. In (d), all of the kernels in (b) and (c) with the addition of a multiphonon kernel using the incoherent approximation. In (e), all of the kernels in (d) are used with multiple scattering turned on.

Overall, the features shown in the experimental data (a) and the simulated data (e) agree very well. Comparison of (b) and (c) shows that coherent scattering gives rise to more features such as diffraction peaks and phonon dispersion curves. It is evident from comparing (c) and (d) that multiphonon scattering increases in intensity at higher Q. The most obvious difference in (d) and (e) is in the elastic line, which shows that multiple scattering of coherent elastic scattering seems to contribute similarly to incoherent scattering in the elastic line. The elastic lines in (a) and (e) suggest that the sample used in the experiment may contain traces of an additional phase, most likely from a surface layer of Al₂O₃. From quantitative comparisons between simulation and experiment, it was found that multi-phonon scattering plays a significant role in the observed intensity, whereas multiple scattering plays little role in this Q range (102). This is as expected because the sample is only a 5% scatterer.
11.6. CORRECTIONS FOR SIMULTANEOUS MULTIPLE SCATTERING AND MULTIPHONON SCATTERING

11.6.2 Approximate Corrections for Simultaneous Multiphonon and Multiple Scattering

 Corrections for multiple scattering have been performed in many ways, from subtracting a constant from the data [1], to full Monte-Carlo simulations [2]. At high temperatures, the former does not account for the slope of the scattering past the cutoff energy. The latter can be computationally intensive, and requires detailed information about the shape of the sample. Here we take an approach of intermediate complexity, reported previously [3]. For both multiple scattering and multiphonon scattering, a two-scattering profile involves a convolution of two single-scattering profiles. In either case, the idea is that an \( n \)-phonon-scattering profile, \( P_n(E) \), is related to the 1-phonon-scattering profile, \( P_1(E) \), through the recursion relation:

\[
P_n(E) = \int_{-\infty}^{\infty} P_{n-1}(E') P_1(E - E') \, dE'.
\]  

(11.48)

For multiple scattering processes, the \( n \)-phonon probability function has additional position and momentum dependencies, that do not appear for multiphonon scattering processes. Sears, et al. [1], argue that the integrals for multiple scattering are related to those for the multiphonon scattering through slowly varying functions of \( Q \) and \( E \). Here we take these functions to be constants, \( a_n \). In essence, we make the approximation that the position and momentum dependencies can be factored out. Thus,

\[
I(Q, E) = N' \sum_{n=1}^{\infty} (1 + a_n) S_n(Q, E),
\]  

(11.49)

where \( I(Q, E) \) is the experimentally-determined total scattering (including multiple scattering), \( S_n(Q, E) \) is the \( n \)-phonon scattering (both creation and annihilation), and \( N' \) is a normalization constant. Note that \( I(Q, E) \) is distinct from the scattering function, \( S(Q, E) \), which does not include multiple scattering [4]. (When we stripped the elastic peak from the data, the dominant multiple elastic scattering is removed, so the index \( n \) in Eq. 11.49 starts at 1 rather than 0.)

Consistent with this factoring of \( Q \) and \( E \) dependencies, we make the incoherent approximation [5]:

\[
S_n^{\text{coh}}(Q, E) = \frac{\sigma_{\text{coh}}}{\sigma_{\text{inc}}} S_n^{\text{inc}}(Q, E),
\]  

(11.50)

where we apply this equation to the 1-phonon terms as well as all higher orders. The last step in our procedure will be to assess any error this has introduced into our analysis.

Our next assumption is that \( a_n = C'_{\text{ms}} \) for all \( n \geq 2 \), where \( C'_{\text{ms}} \) is a single constant that relates the multiple scattering to the multiphonon scattering. Since the multiphonon scattering drops off rapidly with increasing \( n \), this approximation will only have a small effect on our results. The final normalization
is performed with the total scattering, so the factor \(1 + a_1\) is included in the normalization constant. We find:

\[
I(Q, E) = N \left[ S_{\text{inc}}^1(Q, E) + (1 + C_{\text{ms}})S_{\text{inc}}^{2+}(Q, E) \right],
\]

where \(N = N'(1 + a_1) \left( 1 + \sigma_{\text{coh}}/\sigma_{\text{inc}} \right)\) is the normalization constant, and \(1 + C_{\text{ms}} = (1 + C'_{\text{ms}})/(1 + a_1)\). Also, for notational convenience,

\[
S^{2+}(Q, E) \equiv \sum_{n, j} S^n(Q, E).
\]

For a cubic crystal, and a fixed value of \(C_{\text{ms}}\), we can now find the DOS by solving Eq. 11.51 in the manner described by Sears, et al. [1].

Since we do not know the value of \(C_{\text{ms}}\) a-priori, we generate a list of possible values, and solve for the DOS at each one. In the current study, values of \(C_{\text{ms}}\) between 0.0 and 2.0 were tested. It then remains to select the “best” DOS from those generated with the different \(C_{\text{ms}}\). This was done by minimizing a penalty function constructed to find the DOS that produced \(S(E)\) that best satisfied the following conditions:

1. \[
\frac{I(E)}{N} = S_{\text{inc}}1(E) + (1 + C_{\text{ms}})S_{\text{inc}}^{2+}(E),
\]

where the implied sum over \(Q\) allows us to compare the partially coherent scattering on the left with the totally incoherent scattering on the right.

2. The experimental noise at energy transfers near the incident energy oscillates about \((1 + C_{\text{ms}})S_{\text{inc}}^{2+}(E)\).

3. At energy transfers near the incident energy, the slope of a linear fit to the experimental noise matches the slope of a linear fit to \((1 + C_{\text{ms}})S_{\text{inc}}^{2+}(E)\).

These three criteria are correlated, but are not identical. For nickel at 300 K, these three contributions and their sum are shown in Fig. 11.4. Figure 11.5 shows the best fit to the normalized scattering, \(I(E)/N\) for nickel at 300 K, which had \(C_{\text{ms}} = 0.6\).

The DOS curves obtained this way were fit with a Born–von Kármán model, from which all phonon contributions to the scattering, both coherent and incoherent, were calculated. With these results, and with the final value for \(C_{\text{ms}}\), the calculation was checked against the measured scattering. It was our experience that this procedure worked well for the present case of nickel, and also works for cases of other BCC and FCC materials.


Figure 11.4: Penalty functions for nickel at 300 K, as defined in the text. The dash-dotted line (1) relates to the overall fit, the dotted line (2) relates to the noise near the incident energy, and the dashed line (3) relates to the slope near the incident energy. The solid line is the sum of these three contributions (offset).
Figure 11.5: Best fit to scattering for nickel at 300 K. The triangles are the normalized experimental scattering, $I(E)/N$. The solid line shows the fit, $S_{\text{inc}}^1(E) + (1 + C_{\text{ms}})S_{\text{inc}}^{2+}(E)$. The dashed line is the multiple scattering, $C_{\text{ms}}S_{\text{inc}}^{2+}(E)$. The dash-dotted line is the multiphonon scattering, $S_{\text{inc}}^{2+}(E)$. The dotted line is the sum, $(1 + C_{\text{ms}})S_{\text{inc}}^{2+}(E)$. 


Further Reading

The contents of the following are described in the Bibliography.


Chapter 12

Computational Scattering Science

12.1 Software for Inelastic Scattering

12.1.1 Overview of Capabilities

Data from inelastic chopper spectrometers can reveal the fundamental dynamical processes in materials or condensed matter, but not without substantial data analysis to produce even a basic graph of \( S(\vec{Q}, E) \), the intensity as a function of momentum and energy transfers. The elementary excitations in solids are a substantial topic in themselves, and are covered in the earlier chapters of this book. Making comparisons between the experimental \( S(\vec{Q}, E) \) and the underlying theory requires another level of software sophistication. This chapter describes the essential types of computational workflows that can be used to go from data to science.

A schematic overview of software for inelastic scattering is presented in Fig. 12.1. This figure, first presented as a roadmap in Sept. 2001, shows three paths for extracting scientific results from the raw data in the upper left corner. They are:

- Data reduction. The raw data are not interpretable by humans. They must be “reduced” to forms that display trends of intensity versus momentum and/or energy transfer between neutrons and the sample. Identifying features in these data may be publishable in their own right.

- Modeling. The intensity as a function of energy and momentum transfer, \( I(\vec{Q}, E) \), is a quantity that can be obtained from reduced data, and it can be calculated by theoretical models. The comparison between the models and the measurements can reveal new science.

- Direct Experiment Simulation. Although data reduction procedures make
a major effort to remove the quirks of and instrument from the experimental data, this practice is not perfect. For example, it is not always possible to correct accurately for the instrument resolution. Sometimes it is better to simulate the data obtained from an entire experiment, including the effects of the instrument and the sample.

These three paths to science are described in sequence below.

**12.1.2 Data Reduction**

The first path, horizontally across the top of Fig. 12.1, is the traditional approach to data reduction and visualization. The goal of this analysis is to obtain the intensity as a function of momentum transfer and energy transfer, \( S(\vec{Q}, E) \). To do so, the data arrays of counts acquired in terms of instrument parameters such as detector pixel and arrival time must be converted into normalized intensities with physical units such as Å and meV. Instrument backgrounds and other distortions must also be removed. It is often necessary to correct for other distortions caused by, for example, multiple scattering or multiple excitations. By implementing all software components in the interpreted Python language, the architecture can provide a set of components that can be replaced or rearranged to test different processing algorithms. It is also important that data streams can be piped into visualization windows for inspecting the data after the different steps of data processing. An example of a component for energy rebinning of data from a chopper spectrometer is described in the context of the data stream architecture in Sect. ?? of this chapter. Some of the required corrections were described in Sects. 11.3 – 11.6.
12.1. SOFTWARE FOR INELASTIC SCATTERING

12.1.3 Modeling

The second and third paths for extracting scientific results from experimental data are designed to connect experimental data to theory or analytical models.

The second path, the vertical chain in the center of Fig. 12.1, is for comparing experimental results to models of sample dynamics. This path is especially appropriate for analytical models with adjustable parameters. Consider its use for the “neutron-weighting problem” in phonon dynamics, which originates as follows. A measured $S(Q,E)$ from a polycrystalline sample of a pure element can often be converted into a phonon density-of-states using a thermal correction procedure. On the other hand, a phonon DOS from a binary compound cannot be obtained from the measured $S(Q,E)$ because the different elements in the compound do not scatter neutrons with equal efficiencies, causing a “neutron-weighting” of the experimental spectra. Without knowing the lattice dynamics of the compound, it is impossible to know the distortions of an experimental DOS obtained from a measured $S(Q,E)$ after a thermal correction procedure.

The phonon scattering efficiencies of the different atoms are well known, so a lattice dynamics model can be used to calculate an experimental spectrum. We do so with an iterative procedure where the force constants in the dynamics model are varied to obtain the best fit to the experimental data. We therefore can use a lattice dynamics model plus a “neutron weighting” correction to obtain the true phonon density-of-states from the measured $S(Q,E)$ of the compound. The force constants are obtained as parameters that give the best fit between calculation and experiment. This is now a routine procedure, and new experiments can be designed around this capability.

Besides phonon dynamics in ordered compounds, there are many dynamics models that can be compared to experimental data. Applications include data corrections, the determination of parameters such as force constants or exchange stiffnesses, or testing if the model is in fact consistent with the measured $S(Q,E)$. Several types of dynamics models have been included in systems for inelastic scattering software:

1. lattice dynamics with a Born–von Kármán model (periodic structure)
2. spinwave dynamics with a Heisenberg hamiltonian on a periodic structure
3. lattice dynamics on a disordered structure, using a moments analysis of the dynamical matrix
4. lattice dynamics with density functional theory on ordered structures, or disordered structures with supercell calculations
5. ab-initio molecular dynamics calculations with density functional theory on ordered structures, or disordered structures with supercell calculations
6. spin dynamics in a paramagnetic model
7. spin dynamics with density functional theory for chemically ordered structures

12.1.4 Direct Experiment Simulation

The third path from data to science is shown on the left of Fig. 12.1. This approach is a direct simulation of the data measured at the detectors. It is based on Monte Carlo codes that are used in the neutron community for simulation of instrument performance. These codes have been tested and validated against the performance of real instruments. These Monte Carlo simulations have been used with molecular dynamics simulations to perform direct simulations of experimental data. A number of molecular dynamics simulations are available to the theory community. These simulations are complementary to the analytical models of Sect. 12.1.3, and are sometimes advantageous. For example, no approximations are needed when implementing structural models of disordered solids, which are not handled well by the methods of Sect. 12.1.3 (except when the $Q$ information is ignored as in methods 3 and 6).

In these direct experiment simulations, the sample is used as a component in the Monte Carlo computations of neutron trajectories through the instrument, transforming an individual incident neutron into a neutron scattered into the detector array. Typically, simulation results for the primary flight path are stored and used for several simulations. For lattice dynamics simulations, a large four-dimensional ($Q_x, Q_y, Q_z, E$) data structure of dispersive excitations can be pre-calculated, and the velocity-velocity correlation function of the atoms in the sample can be sampled. This same path could be followed for classical spin dynamics simulations.

12.2 Simulation of Neutron Scattering Experiments with MCViNE

12.2.1 General Features of Monte Carlo Ray Tracing Simulations

Except when interacting with atomic-scale periodicities in the sample, much of the simulation of a neutron scattering experiment can be done with the assumption that neutrons are particles that move in approximately straight lines. Monte Carlo (MC) ray tracing simulations of neutron scattering spectrometers with support of multiple scattering were performed as early as the 1970s using MCS(71) and MSCAT(72) to understand the effects of multiple scattering on the measured spectra (73). In the 1990s, with increasing needs of simulating neutron instruments for the purpose of instrument design and optimization, several MC neutron ray tracing packages emerged, including McStas(74; 75), Vitess(76), Ideas(77), and NISP (78). Simulations with these packages were intended primarily for instrument design, such as calculat-
ing energy resolution under different operating conditions (see, for example, (79; 80; 81; 82; 83; 84; 85; 86; 87; 88)). Unlike MCS and MSCAT, most of these newer MC software packages (with a notable exception in NISP) treat a neutron instrument as a linear chain of neutron optical components. Each component performs one modification of neutron beam characteristics such as spatial divergence and energy distribution. The approach based on direct acyclic graphs (Sect. ??) greatly simplifies the coding of instrument simulations. Such an approach is adequate when neutrons are rarely scattered back to the upstream components. These software packages, especially McStas and Vitess, have made important contributions to neutron instrument design.

A Monte Carlo neutron ray tracing simulation follows the neutron from the moderator through the guides and choppers in the primary flight path, to the sample with various possible complexities, and then to a detector or get-lost pipe. There are substantial differences in these components, and natural differences in the structure of the software programs to simulate them.

Detector system

The detector systems in modern neutron scattering instrumentation have a modular and sometimes hierarchical organization, facilitating their construction, maintenance and validation. For example, the four direct geometry time-of-flight spectrometers at Spallation Neutron Source (SNS) (89), ARCS (90), SEQUOIA (91; 92), CNCS(93), and HYSPEC(94) have detector arrays that use use the same so-called “8-packs” (95), each of which is a detector module with eight $^3$He linear position sensitive detector tubes (LPSD). The 8-packs are arranged in a vertically-oriented cylindrical geometry around the sample position, forming a hierarchical organization of pixels, tubes, packs, and detector rows. A simplified illustration of the detector hierarchy can be found in Figure 12.2(c).

Sample Assembly

Samples, sample holders, and sample environments comprise a “sample assembly.” The sample assembly is an aggregation of neutron scatterers near the sample position, as illustrated in Figure 12.2(b). A challenges in simulations of a sample assembly is that neutrons can scatter back and forth among its constituents. Even within a sample of some thickness, multiple scattering is possible. Section 12.3.2 shows simulated and measured effects of multiple scattering in the MICAS furnace, which contains heating elements, heat shields, in addition to the sample (96).

12.2.2 General capabilities of MCViNE

MCViNE(97) is a general purpose neutron ray tracing package that combines the two different approaches taken by MSCAT-like packages and McStas-like packages. At the instrument level, it allows users to create a simulation application as a linear chain of neutron components, each fully configurable by
its type and corresponding geometrical and physical properties using either command line options or an xml-based configuration file. At the component level, two general components exist with support of hierarchical representation for the sample assembly and the detector system. A detector system can be specified with an xml file describing its hierarchy, while a sample assembly is specified by a collection of xml files, one for the geometrical organization of the constituent scatterers, and others describing the scattering mechanisms for them. Multiple-scattering among scatterers in a sample assembly can be turned on and off by a command line option.

12.2.3 Software Architecture of MCViNE

MCViNE is implemented in C++ and Python. The two C++ layers contain fundamental mathematics, the neutron ray tracing mechanism, the mechanism to include components from other MC ray tracing packages, and support of multiple scattering and composite scattering kernels. The neutron components from McStas are included in the C++ layer so they can be used individually rather than as a whole compiled instrument. For the sample, the combination of the object oriented features of C++ and the speed of the compiled code make C++ appropriate for implementing the composite kernel and scattering kernels. The “composite scatterer,” of MCViNE will be explained in detail in Sect. 12.2.4.

A Python layer on top of the two C++ layers allows construction of a component chain similar to McStas-like neutron ray tracing software packages, provides the interface to the C++ components to include in that chain, and allows for introduction of components that are completely written in Python. This last feature makes it extremely easy to create simple neutron components, and to create prototypes of more sophisticated neutron components.

12.2.4 Composite Scatterer

This section starts with basic concepts of the “composite scatterer,” followed by an introduction to the essential object-oriented software designs centered around it, and finishes with examples of ray-tracing algorithms in sample assemblies and detector systems enabled by these designs.

Concepts

In MCViNE, a “composite neutron scatterer” represents a group of physical objects, for example, a powder sample in an aluminum can, a single crystal sample surrounded by a furnace, or (perhaps surprisingly) a detector system. An “elemental scatterer” is a scatterer without constituent scatterers. A “homogeneous scatterer” is one kind of elemental neutron scatterer, whose scattering function is homogeneous within its volume. The scattering properties are modeled using one “scattering kernel” or a combination of several “scattering kernels.”
Figure 12.2: Concepts in scattering composites: composite scatterer, homogeneous scatterer, shape, scattering kernels. (a) an abstract hierarchy of an arbitrary scattering composite. (b) and (c) are concrete examples of such hierarchies. (b) top view of a sample assembly consisting of an aluminum can and a copper plate. (c) a detector system – how it is constructed from an elemental scatterer (detector tube) in a three-level hierarchy.
In the spirit of Sect. 1.1, each scattering kernel allows one scattering mechanism, such as incoherent one-phonon nuclear scattering or coherent magnetic scattering.

Figure 12.2(a) shows an abstract hierarchy of a composite scatterer in MCViNE. Figure 12.2(b) depicts a sample assembly that is a composite of two-level hierarchy, in which the bottom level consists of two homogeneous elemental scatterers: one aluminum can and one copper plate sample. Figure 12.2(c) represents a detector system consisting of $^3$He eight-packs that form roughly a cylindrical arrangement around the sample position. The detector system is represented in MCViNE in a three-level hierarchy: at the bottom level is the $^3$He detector tube; at the middle level, 8 such tubes construct an 8-pack; at the top-level, the detector system consists of a collection of 8-packs. Such hierarchical representations allow MCViNE to model the closely the physical reality in a way that parallels the experimental work.

Object-Oriented Designs

Design patterns for object-oriented programming are developed in the classic text by Gamma, et al., (98). Two important object-oriented design patterns used in MCViNE are

- The “composite pattern,” which describes the part-whole relationship and to represent a hierarchy. It allows clients to treat composites and their constituents in a uniform way.

- The “visitor pattern”, which allows separation of operations from the objects to be operated on, so new operations can be added without touching these objects.

Using these design patterns for scattering composites unifies the programming interfaces to the operations on both the composites and on the individual elemental objects. Composite and visitor patterns are used in three major aspects of the MCViNE neutron scattering model: neutron scatterers, geometric shapes of scatterers, and scattering kernels, as described next.

**Neutron scatterers.** By using the composite pattern, algorithms for multiple scattering are consolidated in one implementation. Scattering from a composite neutron scatterer starts with a determination of which constituent intersects the incident neutron ray, and then delegates the scattering assessment to that particular constituent. The constituent could be a composite itself that requires another delegation for scattering. The hierarchical representation of neutron scatterers and this recursive algorithm work for both samples and detector systems, and can improve computing efficiency and code maintenance.

**Geometric shapes** Using constructive solid geometry (CSG) (see for example (99)), composite shapes are constructed from basic shapes such as cylinders and blocks, and composites by using operations such as union, intersection and difference. Ray-tracing through shapes is therefore simplified as visitor methods of the primitive shapes and the binary shape operations.
12.2. SIMULATION OF NEUTRON SCATTERING EXPERIMENTS WITH MCViNE

Scattering kernels
Composite scattering kernels make it easy for the Monte Carlo algorithm to sample a total $S(Q, \omega)$ consisting of both slowly-varying regions and regions containing sharp features, such as diffraction and coherent phonon scattering, by allowing users to combine different kernels such as incoherent and coherent kernels in a kernel composite. In addition, users can organize scattering kernels into groups; this makes it easy to apply importance sampling (assign different weights to different kernels or kernel groups) in simulations.

A “scattering kernel” in MCViNE is an abstraction of the scattering mechanisms such as diffraction, nuclear scattering by phonons, or magnetic scattering by spin waves. It does not include the sample geometry, but only the scattering physics. By separating the implementation of “scatterer,” “shape,” and “scattering kernel,” a sample in MCViNE generally consists of a combination of scattering kernels that can be added or deleted easily. Furthermore, the scattering kernel library in the MCViNE framework can be extended without affecting the logic of geometric ray tracing, which is implemented in “shape” and “scatterer.” For example, simple scattering kernels taking a histogram form of $S(Q, \omega)$ can be used, or more sophisticated phonon scattering kernels that take phonon energies and polarizations as inputs (these were used in the examples of vanadium and aluminum in Sect. 1.1). Many others are possible.

12.2.5 Algorithms

Ray Tracing
In components such as sample assemblies or detector systems, neutron scatterers are represented by a hierarchy of objects with shapes and scattering mechanisms. Ray tracing of a neutron happens by first determining which neutron scatterer at the top level of the composite hierarchy intercepts the neutron. This is done by computing the intersections of the forward ray of the neutron and all the shapes of the top-level constituents. A random selection might be necessary if multiple top-level objects intercept the neutron. After the top-level neutron scatterer is identified, the neutron is propagated to the front surface of the scatterer if necessary (not necessary if the neutron is already inside the scatterer) with appropriate attenuation, and the ray tracing algorithm then recurses into itself if the scatterer is a composite. Otherwise a point in the forward path of the neutron inside the scatterer is picked at random, and the neutron is propagated to that point with attenuation. At this point a scattering (or an absorption) mechanism of the scatterer is randomly picked, and the neutron is either scattered with its probability adjusted, or is absorbed.

Multiple Scattering
Multiple-scattering (MS) is naturally supported in MCViNE scattering composites, implemented with a recursive algorithm. Here we differentiate between two types of multiple scattering.
Figure 12.3: An example of multiple scattering within one scatterer. The incident neutron was scattered three times by three different scattering kernels. At each scattering point, the original neutron is also propagated out of the scatterer with proper attenuation. Red arrows are paths of neutron propagation. Circles highlight the location of scattering. Different scattering events are coded using different colors.

- single-scatterer multiple scattering (SSMS) to describe the multiple scattering within one neutron scatterer
- multiple-scatterer multiple scattering (MSMS) to describe multiple scattering among neutron scatterers.

Figure 12.3 shows an SSMS in which a neutron is scattered three times before exiting a scatterer. Each time a neutron is scattered inside a homogeneous scatterer, the original incident neutron packet is split into two neutron packets for computational efficiency. One neutron packet is propagated through the scatterer with its probability lowered by attenuation, while the other is scattered by one of the scattering kernels, chosen by a random selection, at a point also randomly selected along the forward path. This splitting process repeats for the scattered neutron several times, as illustrated in Figure 12.3, until either the neutron probability is lower than a pre-selected limit, or the maximum order of multiple scattering is reached. When either termination condition is met, any neutron that is still inside the scatterer is allowed to propagate out, with its probability attenuated appropriately.

Figure 12.4 shows an example of multiple scattering in a sample assembly consisting of a “concentric” arrangement of a sample and a sample environment. Only five possible multiple scattering paths are illustrated, and the splitting processes are not shown. Sample components in some linear-chain-based MC ray tracing packages can support SSMS, but they do not have abstractions similar to composite scatterer or scattering kernel. As a result, the SSMS algorithm must be duplicated in these sample components as explained in Sect. ?? In MCViNE, implementations of new scattering kernels can be added without reimplementing the multiple scattering algorithm. McStas supports MSMS partially for a concentric sample assembly by, for example, adding a second outer cylinder into the simulation component chain (100). Path (e) of Figure 12.4 is included this way, but path (c) of Figure 12.4 is supported only with more work on the user’s part. The multiple scattering algorithm of MCViNE is more
12.2. SIMULATION OF NEUTRON SCATTERING EXPERIMENTS WITH MCViNE

Figure 12.4: An example of multiple scattering in a concentric sample assembly. Five (out of infinite) possible multiple scattering paths are illustrated (see text).

comparable to that of MSCAT(72) in which the sample and the sample environment are treated together in the multiple scattering loop, offering complete treatment of SSMS and MSMS. However, MCViNE allows a straightforward increase in complexity of the sample and sample environment interactions by using the recursive MS algorithm enabled by OOP.

Ray tracing in a detector system

For detector systems, MCViNE takes advantage of the hierarchical representation for neutron scatterers, only in this case the elemental homogeneous neutron scatterer is the \( ^3 \text{He} \) detector tube that intercepts neutrons and records them. MCViNE reuses the code for ray tracing in a composite scatterer for simulating \( ^3 \text{He} \) detector systems, and the new code needed is a scattering kernel for the \( ^3 \text{He} \) that takes into account gas absorption. When a neutron is sent to a detector system shown in Figure 12.2(c), for example, the generic ray tracing algorithm for composite scatterers first checks whether the top level composite scatterer is penetrated by the neutron. If so, all constituents of the composite scatterer, i.e. the detector packs, are examined to determine which of them intercepts the neutron. Unless a neutron traverses a gap between detector packs, the detector pack is identified and then its constituents, the 8 detector tubes, are examined for neutron detection. The path of a neutron through the detector tube is then computed by ray tracing of the neutron through a cylinder, and a MC sampling picks a point in the path for the neutron to be absorbed. (The ray tracing through a cylinder takes care of the parallax effect of the detector tube.) Additionally, the appropriate weighting multiplier for neutron detection probability (computed from absorption probability depending on the \( ^3 \text{He} \) pressure and the length of the neutron path through the tube) and the time-of-flight are computed for the neutron to be recorded as a detector event in a “virtual detector electronics device.” This hierarchical approach to detectors allows for adding more physical features, including details of the charge cloud around the anode wire of the detector.
12.3 Example of a Neutron Simulation

12.3.1 Relationship to Data Reduction

Inelastic neutron scattering data contains a complex overlap of signals with several origins. Multiple scattering and multi-excitation scattering are obvious sources of signal that are usually unwanted in final data analysis. These contributions will vary depending on factors such as sample temperature, sample thickness, additional scattering components in the beam, such as sample containers and sample environments, and the complex geometrical arrangement of all these components. In addition, the instrument has a complex resolution function. Attempts to describe this resolution function analytically separate contributions to the resolution and include uncertainty in timing from the source and differences in path length due to the finite chopper opening (\(\theta\)). The data reduction process attempts to strip the measured data of unwanted contributions to the measured signal, leaving only single-excitation inelastic and elastic scattering from the sample. Because the individual signal contributions are not known, this process is done with a series of approximations that are imperfect, and are also incapable of capturing the complex overlap of signals.

As instruments improve, more detailed information is available from experimental measurements. Some trends and phenomena can be ignored when counting statistics are poor, but become more evident in data of higher quality. There is a growing need for more accurate and reproducible approaches to data reduction. Simulations of neutron experiments offer an alternative way to identify and isolate the individual and mixed contributions to measured intensities.

Section 12.2 described some technical features of the Monte Carlo Virtual Neutron Experiment (MCViNE) software package. In short, MCViNE is a Monte Carlo neutron ray tracing package developed during the commissioning of the ARCS instrument at the SNS (97; 101; 102). This package has several improvements over previous Monte Carlo neutron ray tracing packages, including the use of a recursive function to inherently handle multiple scattering and a flexible architecture for re-arrangement of components.

12.3.2 Scattering from a Sample in a Furnace

A high-temperature furnace used in the ARCS spectrometer is shown in Fig. 12.10. It has many layers of shielding that cause scattering that appears as an unwanted experimental background. In experimental practice, this background can be measured and then subtracted from data acquired from the sample in the furnace. There are two subtleties about this practice that bring ambiguities to this experimental practice:

- When the sample is in the furnace, it removes some of the incident beam, perhaps 10%, by scattering or absorption. The downstream part of the
furnace does not see the full incident beam that was seen for the empty furnace measurement. The typical practice is to assume half of the furnace scattering is from the downstream side, so the full background of an empty furnace is not subtracted from the measured data with the sample. If the sample scatters 10% of the incident neutrons, perhaps 95% of the background from the empty furnace should be removed by subtraction.

- Unfortunately, the parts of the furnace upstream and downstream from the sample are not in the same position, and they give different background signals. Sometimes it is possible to make a third measurement (in addition to the empty furnace, furnace plus sample) with a strongly absorbing material in place of the sample. In this case, the downstream part of the furnace will receive less of the incident beam, and perhaps the scattering contributions from the upstream and downstream parts of the furnace can be isolated by comparing these measurements of a furnace with a black absorber to the measurements of the empty furnace. This can be challenging when the sample and absorber are smaller than the full width of the incident beam, but in principle it can work well.

- A much more difficult problem occurs when there is a first scattering from the furnace to the sample and a second scattering from the sample to the detector (or vice-versa). The intensity from these neutrons that interact with both the sample and the furnace can only be observed when the sample is in the furnace, and an empty furnace measurement is not helpful. Perhaps samples of different thickness can be used to help analyze these multiple scattering effects, but this is not a popular method.

These three problems present an opportunity for simulations. Accurate simulations of the empty furnace, furnace containing the sample, and the sample by itself, can elucidate the contributions of background scattering from the furnace, and between the furnace and sample. These results could show how much of the empty furnace measurement to subtract from the data with the sample, and then show how much multiple scattering between the sample and furnace needs to be removed as well.

### 12.3.3 Steps of Simulations with MCViNE

A “sample assembly” is constructed from several scatterers, arranged in a specific geometrical configuration to model the physical layout of all components in the path of the beam. Construction of the sample assembly is a primary task before starting the simulation, much in the same way as an experimentalist modifies the sample and its environment, rather than the instrument itself. The instrument configuration of the various guides and choppers in the primary flight path upstream of the sample has been previously developed, and neutron beams on the sample for many conditions have already been simulated. The detector configuration is static, and also does not require user modification. The simulation sequence follows four steps illustrated in Fig. 12.5.
Figure 12.5: The simulation proceeds in four steps, as shown for a schematic of the ARCS instrument. First, the neutrons travel from the moderator to the sample (yellow path). Second, the neutrons are incident on the sample and scatter from the sample (pink path). Third, the neutrons are intercepted by the detector array (green path). Fourth, the event-mode NeXus file is reduced using Mantid.

1. First, the beam is simulated as the neutrons travel from the moderator to the sample. The instrument components before the sample are modeled in a linear chain, and scattering probabilities are propagated and updated by each component in the chain until the neutrons reach the sample. This beam simulation is archived, and does not need to be repeated unless the user selects a new configuration of operating parameters.

2. Second, the neutrons are scattered from the sample assembly. The sample assembly includes the sample and furnace as described below.

3. Third, the scattered neutrons are intercepted by the detector array. This assigns a detector pixel ID and time-of-flight to each neutron that reaches the detector. The collected neutrons are then processed into an event-mode NeXus file. This approach to data collection is the same as is currently in use at the SNS.

4. The final step in the simulation sequence is the reduction of the NeXus file using Mantid. This reduction step uses the routines and procedures that are used for reduction of experimental data.

It is also possible to introduce a radial collimator between the second and third steps. The neutrons are forced to travel in straight lines through the collimator after scattering from the sample and before entering the detectors. The collimator triangulates the neutron paths to converge on the sample, and
Figure 12.6: Photograph of the inner edge of collimator septa around the sample region of the ARCS spectrometer. The perspective of the photo is from a camera slightly above the position of the sample. The detector array lies approximately 2 m beyond the collimator towards the top of the photograph.

eliminates much of the scattering from the furnace components that are at different radii. The radial collimator gave a remarkable improvement in the background from the ARCS instrument. Figure 12.6 shows the actual radial collimator in position around the sample space. For the simulation, the radial septa are configured with the same geometrical positioning and spacing as the real collimator, but are modeled as “ideal” in that they are infinitely thin and have 100% absorption.

12.3.4 Furnace Simulation Template

12.3.5 Furnace Geometry

The MICAS furnace, shown in Fig. 12.7 is capable of heating samples to 1900 K (or more practically, 1500 K). In use, the entire furnace is lowered with a crane into the sample region. The bottom-most portion of the furnace, the many-layered region in the schematic and the crinkled-foil region in the photo, is the part of the furnace in the path of the neutron beam. This is this region that must be accurately described in any Monte Carlo simulation.

The critical region of the furnace in the neutron beam contains the sample at
Figure 12.7: Schematic of the MICAS furnace (left) and the furnace itself (right), supported on a yellow cart.
12.3. EXAMPLE OF A NEUTRON SIMULATION

Table 12.1: Outer diameters of the concentric cylinders of heating elements, shielding, and the outer vacuum tank containment for the MICAS furnace. All of these components are made from high-purity Nb foil with a thickness of 0.002 in.

<table>
<thead>
<tr>
<th>Component</th>
<th>Diameter (in)</th>
</tr>
</thead>
<tbody>
<tr>
<td>inner heating element</td>
<td>3.13</td>
</tr>
<tr>
<td>outer heating element</td>
<td>3.65</td>
</tr>
<tr>
<td>heat shield 1</td>
<td>5.10</td>
</tr>
<tr>
<td>heat shield 2</td>
<td>5.52</td>
</tr>
<tr>
<td>heat shield 3</td>
<td>5.93</td>
</tr>
<tr>
<td>heat shield 4</td>
<td>6.35</td>
</tr>
<tr>
<td>heat shield 5</td>
<td>6.77</td>
</tr>
<tr>
<td>heat shield 6</td>
<td>7.18</td>
</tr>
<tr>
<td>heat shield 7</td>
<td>7.60</td>
</tr>
<tr>
<td>heat shield 8</td>
<td>8.01</td>
</tr>
<tr>
<td>outer tank</td>
<td>11.54</td>
</tr>
</tbody>
</table>

the center, which is attached to the bottom of a stick and inserted into the furnace. Concentric cylinders around the sample providing heating and thermal radiation shielding. Their dimensions are given in Table 12.1. Figure 12.8 depicts these concentric cylinders. The two inner-most cylinders are the heating element. They are surrounded by between 1 to 8 layers of shielding for thermal radiation that suppress heat loss from the furnace. (Dissipation of heat from the heating element into the sample region of the instrument tank is an issue that must be observed closely, ideally by monitoring the temperature on the outer tank layer. In recent modifications of the MICAS furnace design, this problem is mitigated by water cooling above and below the sample region of the furnace.) It was discovered that the precise diameters of these heating elements and heat shields is crucial for accurately modeling their scattering. The values given in Table 12.1 were obtained from an actual measurement of the heating element and heat shields with a micrometer when the furnace was deconstructed and not in use. Initially, all of the concentric layers were simulated as individual scatterers. This was cumbersome and proved to be unnecessary, however.

Details of Furnace Model

The furnace assembly was modeled as two scatterers: (1) the outer vacuum container, and (2) the two heating elements and eight heat shields. The outer vacuum container is modeled as a hollow cylinder with radius 5.77 in., thickness 0.1 mm, and height 15 in. As is obvious from the photo in Fig. 12.7, this outer layer is not a smooth cylinder because of some mechanical abuse. It also serves as the outer vacuum containment for the furnace, so any pressure imbalance
Figure 12.8: Drawing (not to scale) of the heating element and heat shield region of the MICAS furnace. The inner and outer heating elements and the outer tank are fixed. Heat shields can be removed, beginning with the outermost, depending on the maximum temperature of the experiment.

that is created during pump down and venting of the furnace and sample area can cause this thin layer to crinkle. So instead of modeling this region as a perfectly dense cylinder of thickness 0.002 in. (0.05 mm), it is modeled with twice this thickness, and a reduced “packing factor” of 0.5. In Fig. 12.8, it is depicted as the blue cylinder.

The scatterer describing the remaining components also has a hollow cylinder shape with inner radius 1.56 in., outer radius 4 in., and height 15 in. This hollow cylinder has a reduced packing factor of 0.008 to capture the considerable amount of ‘empty space’ present in this cylinder. In Fig. 12.8, this is depicted by the volume encompassed by the two red cylinders. Figure 12.9 is an example of the furnace template in the sample assembly file. In practice, the sample assembly is an xml file modified by the user to add or remove components of the sample to the simulation. In this example file, the furnace template is in use as indicated by the blue labels. The shape, composition, and geometric conditions described for these components are input variables. A separate “scatterer” file for each of the components provides details including the packing factor, scattering kernels, and scattering probabilities.

Computational Results

Neutron scattering was both measured and computed for an “empty” furnace at room temperature, both with and without the radial collimator. Figure 12.10 compares experimental results (a, b) with simulations (c-f). For these test cases, the empty furnace is not truly empty. The experiment contained an empty Nb
Figure 12.9: The furnace template contributes two components to the sample assembly. The blue labels indicate which component is being described. The “outer most” and “Nb heating elements etc. 2+8” make up the furnace.
sample sachet fixed in a BN absorbing frame at 45 degrees to the incident beam. The simulation was simpler, however, containing two pieces of Nb foil at the sample position, also at 45 degrees to the incident beam.

The experimental data in (a) and (b) are plotted on the same intensity scale, demonstrating the dramatic effect of the radial collimator in reducing unwanted scatter from the sample environment. Prior to the introduction of the radial collimator, the result in (a) shows the considerable background scattering that needed to be removed from the experimental data. Most notable is the high intensity scattering along the elastic line, including a peculiar split in the elastic intensity, especially towards higher $Q$. Inelastic scattering intensity is also visible, and it is even possible to see some phonon dispersions from Nb.

The broad Bragg diffractions, which split into two at higher $Q$, originate from two distinct regions of scatterers in the furnace. There is a high concentration of Nb near the sample position, including the Nb foil at the sample position and 10 concentric layers of Nb foil within 4 in. of the beam center. This produces a set of elastic scattering peaks with an intensity somewhat broadened along $E$. The second set of elastic scattering peaks results from the outermost Nb foil layer, which is at a considerable distance of nearly 6 in. from the sample center. A gap of nearly 2 in. exists between the sample and shielding, and this outer cylinder, causing its own set of diffraction peaks from scattering. In the experimental measurement with the collimator introduced (b), this effect is entirely eliminated. This is as expected because the radial geometry of the collimator is highly effective in removing scattering that occurs at angles not radiating directly from the sample position. The collimator is less effective at low $Q$, and the lowest-order diffractions from Nb have moderate intensity with the collimator in place.

The simulations in Fig. 12.10 show the furnace without the collimator (c, e) and with the collimator (d, f), and provide a comparison with (e, f) and without (c, d) multiple scattering. The intensity scales are matched in all of the simulated results to demonstrate the high efficiency of the collimator in reducing this unwanted background scattering (so it is easier to analyze the results for the simulation without the collimator). Panel (c) shows the empty furnace with all scattering kernels in use, but without multiple scattering. This reproduces the experiment data in (a) reasonably well, although intensity along the elastic line, especially at low $Q$, is noticeably absent. This is also evident in comparison of (b) and (d), the experiment with the collimator in place and simulation without multiple scattering. In (e), multiple scattering is introduced, which provides the missing intensity near $E=0$ and also smears out the scattering along $Q$. This appears to also be the case in (f), though it is less obvious. Multiple scattering often involves two elastic scattering events, but these tend to be incoherent and contribute intensity between the Bragg peaks from the Nb. A similarly broad elastic line is seen in Fig. 1.1e for multiple scattering from fcc Al, for which the individual scatterings were assumed coherent.

These empty furnace measurements and simulations demonstrate the viability of simulating background intensities in inelastic neutron scattering experiments. Not only are the results interpretable in terms of individual scattering
12.3. EXAMPLE OF A NEUTRON SIMULATION

Figure 12.10: The empty furnace provides significant background, as is visible from the experiment without the collimator (a), but the background is considerably reduced with the collimator (b). The simulation without the collimator is shown without multiple scattering (c) and with multiple scattering (e). The simulation with the collimator is also shown without multiple scattering (d) and with multiple scattering (f).
processes, but they are quantitatively accurate. Nevertheless, this approach is not fully proven. It still seems prudent to make experimental measurements of the main contributions to the background, such as the furnace itself, and use the simulations to correct for more difficult effects such as multiple scattering between the sample and the furnace.

12.4 The Future of Computational Scattering Science

The opportunities of computational scattering science, and the effort to achieve them, are not new ideas. The authors had understood many of them around the end of the 1900's. It is somewhat disappointing that they are not more widespread today. Some of this is understandable, since it a project in computational scattering science requires success in both experiment and in computation. Obviously it is easier to address only one of these, and this has been the historical norm. However, “Without deviation from the norm, progress is not possible,” as pointed out by Frank Zappa. What follows are a number of observations about the opportunities and the paths needed to achieve them.

12.4.1 Some General Thoughts

Today it requires less effort to use sophisticated tools for computational materials science than it did about 10 years ago. Today a graduate student may routinely use more than a million CPU hours per year, and this will increase steadily. Yet there remains a large learning curve for applying these tools to Scattering Science.

The bigger challenge is to incorporate these computational tools into workflows for doing new science. This is not moving forward very rapidly, and the status quo of an individual scientist/developer is an inefficient cottage industry. The scattering community needs help to make the new computational tools easier to use. There is some need to develop new codes for computational scattering science, but the bigger need is to use the existing codes for calculating the properties of materials, and integrate them into workflows to interpret the results of experimental scattering data. These workflows could be archived and shared, allowing them to be adapted quickly for new scientific endeavors, supporting scattering scientists who need them.

It is of course paramount to quantify the accuracy of quantitative science. When sophisticated features are extracted from a workflow that combines computing and experiment, or when different types of experimental data are combined to develop an underlying materials model, we have little experience with the reliability of the results. Uncertainty quantification needs to be considered when building new scientific workflows. Although underutilized in scattering science, Bayesian methods can be incorporated naturally into efforts that combine computation with experiment, using results from one as prior information
for the other, for example. These methods are challenged when the models for obtaining conditional probabilities are not well-known, however.

Nearly all scattering science research with X-ray and neutron facilities involves studies of materials, but the experiments cover an enormous range of different materials and phenomena. Nevertheless, some computational materials science methods are important for broad classes of materials. All atoms obey quantum mechanics, for example. Modern methods for calculating electronic structure, molecular dynamics, and tools for modeling atomic structure and dynamics are well established. Software that simulates materials at the atomic level using quantum mechanics and statistical mechanics has become both commonplace and essential for understanding a large range of phenomena. It is time to develop specific scientific workflows that use these modern tools of computational materials science to assist in interpreting specific scattering experiments.

12.4.2 Some Specific Workflows

Direct experiment simulations have not reached their full potential in 2016. These simulate neutrons sent into a beamline, how the beams are monochromated and focused, and how they are scattered from the sample into detectors. The quirks of the instrument are included in these simulations, and in many cases it is more reliable to compare simulated data to real data, rather than trying to correct experimental data for characteristics of the instrument. Central to these simulation workflows is the scattering from the sample. We know how a neutron is deflected by different interactions with the atoms in the sample, and we now have the tools to calculate the positions of atoms, their vibrational dynamics, their magnetic moments, and how local magnetic moments respond to temperature and magnetic field. These ab-initio codes are a triumph of quantum mechanics in the past two decades. We should be using them to calculate the relevant correlation functions and cross-sections for scattering events, and using these results in the simulations of experimental data.

There is broad agreement that density functional theory (DFT) codes are the standard workhorses for computational materials science today, and there should be no major issues incorporating them into workflows that calculate diffraction patterns and phonon dispersions of regular crystalline material, for example. To date, many studies have already benefited from DFT workflows. Classical molecular dynamics codes could also be made available this way, but using them for accurate simulations of real materials requires expertise in “training” their force fields. This detailed work may be beyond the patience of computational scattering scientists. Although ab-initio molecular dynamics demands much more of the computer, it demands less from the scientist. Finally, there is widespread interest, both in theory and in experiment, in understanding materials when their structures are not in equilibrium. Codes to calculate the structures and dynamics of materials with excited electrons are emerging, and are currently limited to much smaller systems than are typical DFT calculations due to the less favorable scaling with system size of these methods.
Uncertainty quantification is hardly used by the scattering science community today, even though it is almost always desirable to estimate errors in experimentally-derived quantities. When these quantities are not obtained by direct measurement, but instead the measurements are used to optimize parameters in an underlying atomic model, for example, it is challenging to estimate the uncertainties. There are some families of workflows where classical Bayesian statistics could be adapted to estimate uncertainty. These methods are not implemented in core packages of computational materials science. Software workflows need to give more attention to uncertainty quantification, especially those that combine multiple sets of data and computation. It is unfortunate that there is so little reward for the effort of doing this, but this work is especially important when the models themselves are uncertain or controversial. In such cases the traditional Bayesian approach may not be appropriate, but there are new efforts in the applied mathematics of uncertainty quantification that could help over the next years.

Adapting computational tools to new scientific workflows is most efficient when the software packages are modular, and have consistent interfaces for I/O, for example. The rules of good object-oriented programming go a long way towards satisfying this goal, although compatibility must be designed. Also necessary will be workflow documentation, discussion forum management, and some brokerage of computing resources. Software development with professionalism, and with steady input from the scattering community, is essential for building useful and sustainable software for computational scattering science. We are not doing this so well today, but the promise is there. It seems too obvious to go unrecognized for much longer.

Further Reading

The contents of the following are described in the Bibliography.
Appendix A

Appendix 3: Selected Derivations and Physical Constants

A.1 Convolutions and Correlations

A.1.1 Convolution Theorem

It is easiest to explain convolutions in terms of a broadening of a sharp peak caused by making a measurement with a blurry instrument. The instrumental broadening function is \( f(k) \). We seek the true specimen diffraction profile \( g(k) \). What we actually measure with our diffractometer is the convolution of \( f(k) \) and \( g(k) \), denoted \( h(k) \) (where \( K \) is the shift of the detector across the diffraction intensity). Deconvolution will require the Fourier transforms of \( f(k) \), \( g(k) \), \( h(K) \):

\[
\begin{align*}
  f(k) & = \sum_n F(n) e^{i2\pi nk/l} \quad \text{equipment,} \quad (A.1) \\
  g(k) & = \sum_n G(n') e^{i2\pi n'k/l} \quad \text{specimen,} \quad (A.2) \\
  h(K) & = \sum_n H(n'') e^{i2\pi n''K/l} \quad \text{measurement.} \quad (A.3)
\end{align*}
\]

Note that \( l \) has units of inverse distance, so \( n/l \) is a real space variable. The range in \( k \) of the Fourier series is the interval \(-l/2\) to \(+l/2\), which includes all

\footnote{Measurements are typically in scattering angle, which is interpretable as a \( k \)-space variable.}
features of a diffraction peak. The convolution of \( f \) and \( g \) is defined as:

\[
h(K) = \int_{-\infty}^{\infty} f(K - k) g(k) \, dk.
\] (A.4)

We must choose an interval so that that \( f \) and \( g \) vanish outside the range \( \pm \frac{l}{2} \), so we can change the limits of integration from \( \pm \infty \) to \( \pm \frac{l}{2} \). Substitute (A.1) and (A.2) into (A.4):

\[
h(K) = \frac{l}{2} \sum_{n} \sum_{n'} G(n') F(n) e^{i2\pi nK/l} e^{i2\pi n'k/l} \, dk.
\] (A.5)

We rearrange summations over the independent variables \( n \) and \( n' \), and remove from the integral all factors independent of \( k \):

\[
h(K) = \sum_{n} \sum_{n'} G(n') F(n) e^{i2\pi nK/l} \int_{-\frac{l}{2}}^{\frac{l}{2}} e^{i2\pi (n' - n)k/l} \, dk.
\] (A.6)

Now we employ the orthogonality condition:

\[
\int_{-\frac{l}{2}}^{\frac{l}{2}} e^{i2\pi (n' - n)k/l} \, dk = \left\{ \begin{array}{ll} 1 & \text{if } n' = n \\ 0 & \text{if } n' \neq n \end{array} \right\}.
\] (A.7)

With the orthogonality condition of (A.7), the double sum in (A.6) is reduced to a single sum:

\[
h(K) = l \sum_{n} G(n) F(n) e^{i2\pi nK/l}.
\] (A.8)

Compare (A.8) to the definition for \( h(K) \) in (A.3). We see that that the Fourier coefficients \( H(n') \) are proportional to the product of \( G(n) \) and \( F(n) \):

\[
l G(n) F(n) = H(n).
\] (A.9)

By comparing (A.4) and (A.9), we see that a convolution in \( k \)-space is equivalent to a multiplication in real space (with variable \( n/l \)). The converse is also true; a convolution in real space is equivalent to a multiplication in \( k \)-space. This important result is the convolution theorem.

---

2We don’t care about \( f(k) \) and \( g(k) \) outside this interval, but with (A.1)–(A.3) these Fourier transforms repeat themselves with a period of \( l \). We confine ourselves to one period, and require that \( f \) and \( g \) vanish at its ends.

3Verified by writing the exponential as \( \cos(2\pi (n' - n)k/l) + i \sin(2\pi (n' - n)k/l) \). The sine integration vanishes by symmetry. The cosine integration gives \([2\pi (n' - n)]^{-1} \sin(\pi (n' - n)) \sin(\pi (n' - n))\), which = 0 when \( n' - n \neq 0 \). In the case when \( n' - n = 0 \), the integrand in (A.7) equals 1, so the integration gives \( l \).
A.2. FOURIER TRANSFORM OF SCREENED COULOMB POTENTIAL

A.1.2 Deconvolutions

Equation (A.9) shows how to perform the deconvolution of \( f(k) \) from \( h(K) \); perform a division in \( n \)-space. Specifically, when we have the full sets of Fourier coefficients \( \{F(n)\} \) and \( \{H(n)\} \), we perform a division in \( n \)-space for each Fourier coefficient:

\[
G(n) = \frac{1}{I} \frac{H(n)}{F(n)}. \tag{A.10}
\]

We obtain each \( F(n') \) by multiplying both sides of (A.1) by \( \exp(-i2\pi n'k/l) \) and integrating over \( k \):

\[
\frac{1}{l} \int_{-l/2}^{l/2} f(k) e^{-i2\pi n'k/l} \, dk = \sum_n F(n) \int_{-l/2}^{l/2} e^{i2\pi (n-n')k/l} \, dk. \tag{A.11}
\]

The orthogonality relationship of (A.7) causes the right-hand-side of (A.11) to equal zero unless \( n = n' \). Equation (A.11) therefore becomes:

\[
\frac{1}{l} \int_{-l/2}^{l/2} f(k) e^{-i2\pi n'k/l} \, dk = F(n'). \tag{A.12}
\]

The Fourier coefficients \( H(n) \) are obtained the same way. The simple division of Fourier coefficients in (A.10) then provides the set of Fourier coefficients for the true specimen profile, \( \{G(n)\} \). If we then use (A.2) to take the Fourier transform of the \( \{G(n)\} \) from (A.10), we obtain \( g(k) \), the true specimen diffraction profile.

A.2 Fourier Transform of Screened Coulomb Potential

In this subsection we calculate the Fourier transform of a “screened Coulomb” potential, a result that is useful in calculations of form factors of atoms for example. This screened Coulomb potential, \( V(\vec{r}) \), is:

\[
V(\vec{r}) = -\frac{Ze^2}{r} e^{-r/r_0}. \tag{A.13}
\]

The exponential factor accounts for the screening of the nuclear charge by the atomic electrons, and \( r_0 \) is an effective Bohr radius for the atom. Interestingly, the exponential decay also facilitates the mathematics of working with a potential that is otherwise strong at very large distances.

We now use the first Born approximation, (1.75), to calculate the atomic scattering factor, \( f(\Delta \vec{k}) \), as the Fourier transform of \( V(\vec{r}) \):

\[
f_{el}(\Delta \hat{k}) = -\frac{m}{2\pi \hbar^2} \int_{\text{all space}} e^{-i\Delta \hat{k} \cdot \vec{r}} V(\vec{r}) \, d^3 \vec{r}. \tag{A.14}
\]
Substituting the potential (A.13) into (A.14):

\[ f_{el}(\vec{\Delta k}) = \frac{mZ\varepsilon^2}{2\pi\hbar^2} \int \frac{e^{-i\vec{\Delta k} \cdot \vec{r}}}{r} d^3\vec{r}. \]  

(A.15)

The integral, \( I(\vec{\Delta k}, r_0) \), in (A.15) occurs in other contexts, so we pause to solve it.

\[ I(\vec{\Delta k}, r_0) \equiv \int \frac{e^{-i\vec{\Delta k} \cdot \vec{r}}}{r} d^3\vec{r}, \]  

(A.16)

which is the 3-dimensional Fourier transform of the screened Coulomb potential (A.13). It is natural to use spherical coordinates:

\[ I(\vec{\Delta k}, r_0) = \int_0^\infty \int_0^\pi \int_0^{2\pi} e^{-i\vec{\Delta k} \cdot \vec{r}} \frac{e^{-r/r_0}}{r^2} r^2 \sin\theta \, d\theta \, d\phi \, dr. \]  

(A.17)

The trick for working with the exponential in (A.17), \( e^{-i\vec{\Delta k} \cdot \vec{r}} \), is to align the vector \( \vec{\Delta k} \) along the z-axis so that \( \vec{\Delta k} \cdot \vec{r} = \Delta kz \). Also, since \( z = r\cos\theta \):

\[ dz = -r\sin\theta \, d\theta. \]  

(A.18)

The limits of integration are changed as:

\[ \theta = 0 \Rightarrow z = r, \]  

(A.19)

\[ \theta = \pi \Rightarrow z = -r. \]  

(A.20)

With the substitution of (A.18)–(A.20) into (A.17):

\[ I(\vec{\Delta k}, r_0) = \int_0^\infty \int_0^\pi \int_0^{2\pi} e^{-i\vec{\Delta k} \cdot \vec{r}} e^{-r/r_0} d\phi (-dz) dr, \]  

(A.21)

\[ I(\vec{\Delta k}, r_0) = 2\pi \int_0^\infty \int_{r_0}^r e^{-i\vec{\Delta k} \cdot \vec{r}} e^{-r/r_0} dz \, dr. \]  

(A.22)

Writing the exponential as \( e^{-i\Delta k z} = \cos(\Delta k z) - i \sin(\Delta k z) \), the z-integration of the sine function vanishes by symmetry in the interval \(-r\) to \(+r\), and the cosine integral is:

\[ \int_{-r}^{r} \cos(\Delta k z) \, dz = \frac{+2}{\Delta k} \sin(\Delta k r), \]  

(A.23)
which does not depend on the direction $\Delta \vec{k}$. Using (A.23) for the $z$-integration in (A.22), we obtain:

$$I(\Delta k, r_0) = \frac{4\pi}{\Delta k} \int_{r=0}^{\infty} \sin(\Delta kr) e^{-r/r_0} dr . \quad (A.24)$$

Equation (A.24) is the Fourier transform of a decaying exponential. This integral can be solved by twice integrating by parts. The result is a Lorentzian function:

$$\int_{r=0}^{\infty} \sin (\Delta kr) e^{-r/r_0} dr = \frac{\Delta k}{\Delta k^2 + \frac{1}{r_0}} . \quad (A.25)$$

We substitute the result (A.25) into (A.24), completing the evaluation of (A.16):

$$I(\Delta k, r_0) = \int_{\text{all space}} e^{-i\Delta k \cdot r} e^{-r/r_0} \frac{1}{r} d^3r = \frac{4\pi}{\Delta k^2 + \frac{1}{r_0}} . \quad (A.26)$$

For later convenience, we now obtain a related result. The use of an exponential screening factor to perform a Fourier transform of the Coulomb potential is a useful mathematical trick. By letting $r_0 \to \infty$, we suppress the screening of the Coulomb potential, so $e^{-r/r_0} = 1$ in (A.13). The Fourier transform of this bare Coulomb potential, with its mathematical form of $1/r$, is obtained easily from (A.26):

$$\int_{\text{all space}} e^{-i\Delta k \cdot r} \frac{1}{r} d^3r = \frac{4\pi}{\Delta k^2} . \quad (A.27)$$

---

4Defining $U \equiv e^{-r/r_0}$ and $dV \equiv \sin(\Delta kr) dr$, we integrate by parts: $\int U dV = UV - \int V dU$. The integral on the right hand side is evaluated as: $(\Delta kr_0)^{-1} \int_{r=0}^{\infty} \cos(\Delta kr) e^{-r/r_0} dr$, which we integrate by parts again to obtain: $-(\Delta kr_0)^{-2} \int_{r=0}^{\infty} \sin(\Delta kr) e^{-r/r_0} dr$. This result can be added to the $\int U dV$ on the left hand side to obtain (A.25).
### A.3 Fundamental and Derived Constants

#### Fundamental Constants

- \( \hbar = 1.0546 \times 10^{-27} \text{ erg-sec} = 6.5821 \times 10^{-16} \text{ eV-sec} \)
- \( k_B = 1.3807 \times 10^{-23} \text{ J/(atom-K)} = 8.6174 \times 10^{-5} \text{ eV/(atom-K)} \)
- \( R = 0.00198 \text{ kcal/(mole-K)} = 8.3145 \text{ J/(mole-K)} \) (gas constant)
- \( c = 2.998 \times 10^{10} \text{ cm/sec} \) (speed of light in vacuum)
- \( m_e = 0.91094 \times 10^{-27} \text{ g} = 0.5110 \text{ MeV/c}^2 \) (electron mass)
- \( m_n = 1.6749 \times 10^{-24} \text{ g} = 939.55 \text{ MeV/c}^2 \) (neutron mass)
- \( N_A = 6.02214 \times 10^{23} \text{ atoms/mole} \) (Avogadro constant)
- \( e = 4.80 \times 10^{-10} \text{ esu} = 1.6022 \times 10^{-19} \text{ coulomb} \)
- \( \mu_0 = 1.26 \times 10^{-6} \text{ henry/m} \)
- \( \epsilon_0 = 8.85 \times 10^{-12} \text{ farad/m} \)

- \( a_0 = h^2/(m_e c^2) = 5.292 \times 10^{-9} \text{ cm} \) (Bohr radius)
- \( \epsilon^2/(m_e c^2) = 2.81794 \times 10^{-13} \text{ cm} \) (classical electron radius)
- \( \epsilon^2/(2a_0) = R \) (Rydberg) = 13.606 eV (K-shell energy of hydrogen)
- \( \epsilon \hbar/(2m_e c) = 0.9274 \times 10^{-20} \text{ erg/oersted} \) (Bohr magneton)
- \( \hbar^2/(2m_e) = 3.813 \times 10^{-16} \text{ eV s cm}^{-2} \)

#### Definitions

- 1 becquerel (B) = 1 disintegration/second
- 1 Curie = 3.7 \times 10^{10} \text{ disintegrations/second}

**radiation dose:**
- 1 roentgen (R) = 0.000258 coulomb/kilogram
- Gray (Gy) = 1 J/kg

Sievert (Sv) is a unit of “radiation dose equivalent” (meaning that doses of radiation with equal numbers of Sieverts have similar biological effects, even when the types of radiation are different). It includes a dimensionless quality factor, Q (Q \approx 1 for x-rays, 10 for neutrons, and 20 for α-particles), and energy distribution factor, N. The dose in Sv for an energy deposition of D in Grays [J/kg] is:

\[
\text{Sv} = Q \times N \times D \text{ [J/kg]}
\]

Rad equivalent man (rem) is a unit of radiation dose equivalent approximately equal to 0.01 Sv for hard x-rays.

- 1 joule = 1 J = 1 W·s = 1 N·m = 1 kg·m²·s⁻²
- 1 joule = 10⁷ erg
- 1 newton = 1 N = 1 kg·m·s⁻²
- 1 dyne = 1 g·cm·s⁻² = 10⁻⁵ N
- 1 erg = 1 dyne-cm = 1 g·cm²·s⁻²
- 1 Pascal = 1 Pa = 1 N·m⁻²
- 1 coulomb = 1 C = 1 A·s
- 1 ampere = 1 A = 1 C/s
### A.3. FUNDAMENTAL AND DERIVED CONSTANTS

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<thead>
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<th>Element</th>
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<th>$K\alpha_2$</th>
<th>$K\beta_1$</th>
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<tbody>
<tr>
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<td>1.54052</td>
<td>1.54433</td>
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<td>Mo</td>
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<td>Ag</td>
<td>0.56083</td>
<td>0.55936</td>
<td>0.56377</td>
<td>0.49701</td>
</tr>
</tbody>
</table>

1 volt = 1 V = 1 W·A$^{-1}$ = 1 m$^2$·kg·A$^{-1}$·s$^{-3}$
1 ohm = 1 Ω = 1 V·A$^{-1}$ = 1 m$^2$·kg·A$^{-2}$·s$^{-3}$
1 farad = 1 F = 1 C·V$^{-1}$ = 1 m$^{-2}$·kg$^{-1}$·A$^2$·s$^4$
1 henry = 1 H = 1 Wb·A$^{-1}$ = 1 m$^2$·kg·A$^{-2}$·s$^{-2}$
1 tesla = 1 T = 10,000 gauss = 1 Wb·m$^{-2}$ = 1 V·s·m$^{-2}$ = 1 kg·s$^{-2}$·A$^{-1}$

### Conversion Factors

<table>
<thead>
<tr>
<th>Conversion</th>
<th>Value</th>
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<tbody>
<tr>
<td>1 Å = 0.1 nm = 10$^{-4}$ µm = 10$^{-10}$ m</td>
<td></td>
</tr>
<tr>
<td>1 b (barn) = 10$^{-24}$ cm$^2$</td>
<td></td>
</tr>
<tr>
<td>1 eV = 1.6045 × 10$^{-12}$ erg</td>
<td></td>
</tr>
<tr>
<td>1 eV/atom = 23.0605 kcal/mole = 96.4853 kJ/mole</td>
<td></td>
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<tr>
<td>1 cal = 4.1840 J</td>
<td></td>
</tr>
<tr>
<td>1 bar = 10$^5$ Pa</td>
<td></td>
</tr>
<tr>
<td>1 torr = 1 T = 133 Pa</td>
<td></td>
</tr>
<tr>
<td>1 kG = 5.6096 × 10$^{29}$ MeV·c$^{-2}$</td>
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</tr>
</tbody>
</table>

### Useful Facts

- energy of 1 Å photon = 12.3984 keV
- $\hbar$ for $10^{12}$ Hz = 4.13567 meV
- 1 meV = 8.0655 cm$^{-1}$
- temperature associated with 1 eV = 11,600 K
- lattice parameter of Si (in vacuum at 22.5°C) = 5.431021 Å

### Neutron Wavelengths, Energies, Velocities

$E_n = 81.81 \lambda^{-2}$ (energy-wavelength relation for neutrons [meV, Å])
$\lambda_n = 3955.4/v_n$ (wavelength-velocity relation for neutrons [Å, m/s])
$E_n = 5.2276 \times 10^{-6} v_n^2$ (energy-velocity relation for neutrons [meV, m/s])

### Relativistic Electron Wavelengths

For an electron of energy $E\text{[keV]}$ and wavelength $\lambda\text{[Å]}$:

\[
\lambda = \frac{\hbar}{2m_eE\left(1 + \frac{E}{2m_e c^2}\right)^{1/2}} = \frac{0.3877}{E^{1/2} \left(1 + 0.9788 \times 10^{-3}E\right)^{1/2}}
\]
Table A.1: Parameters of high-energy electrons

<table>
<thead>
<tr>
<th>$E$ [keV]</th>
<th>$\lambda$ [Å]</th>
<th>$\gamma$</th>
<th>$v$ [c]</th>
<th>$T$ [keV]</th>
</tr>
</thead>
<tbody>
<tr>
<td>100</td>
<td>0.03700</td>
<td>1.1957</td>
<td>0.5482</td>
<td>76.79</td>
</tr>
<tr>
<td>120</td>
<td>0.03348</td>
<td>1.2348</td>
<td>0.5867</td>
<td>87.94</td>
</tr>
<tr>
<td>150</td>
<td>0.02956</td>
<td>1.2935</td>
<td>0.6343</td>
<td>102.8</td>
</tr>
<tr>
<td>200</td>
<td>0.02507</td>
<td>1.3914</td>
<td>0.6953</td>
<td>123.6</td>
</tr>
<tr>
<td>300</td>
<td>0.01968</td>
<td>1.587</td>
<td>0.7765</td>
<td>154.1</td>
</tr>
<tr>
<td>400</td>
<td>0.01643</td>
<td>1.7827</td>
<td>0.8279</td>
<td>175.1</td>
</tr>
<tr>
<td>500</td>
<td>0.01421</td>
<td>1.9785</td>
<td>0.8628</td>
<td>190.2</td>
</tr>
<tr>
<td>1000</td>
<td>0.008715</td>
<td>2.957</td>
<td>0.9411</td>
<td>226.3</td>
</tr>
</tbody>
</table>

kinetic energy \( \equiv T = \frac{1}{2} m_e v^2 = \frac{1}{2} E^{1+\gamma} \frac{1}{\gamma} \)
Appendix A

Appendix 2: Software Design

A.1 Extending *DANSE*: Writing C++ Extensions to Python

Important sources of information on this topic are available elsewhere: the Python extension and embedding manual, and the Python-C API reference manual. These documents are available at the Python website: http://python.org, but better yet, just look in the ‘ext’ and ‘api’ subdirectories of your Python distribution’s Doc directory for HTML versions.

This section was written to supplement those documents, first by giving an example of dynamically allocating C++ objects and keeping track of them, and second by introducing some of the API functions for working with aggregate types like tuples and lists, for which the easiest Python-C conversion tools don’t work. Also, I’ve concocted some simple, try-this-at-home examples to illustrate the process. Finally, I’ve added a more real-life example.

Note: In the following, C++ is used to mean both C and C++. The Python-C API is written in C, but for the most part, but that can be called seamlessly from C++. There’s only one item that must have C linkage, the init function, described below.

A.1.1 Overview

Why write C++ extensions to Python? To reuse existing code, and to gain better performance. A great deal of software has already been written in C and C++ (not to mention FORTRAN), and, at least at the present, nothing beats compiled languages for performance in the numerically intensive codes that *DANSE* supports. Extending Python allows us to turn all that code into building blocks for solutions. Libraries of extensions package those building blocks into kits that users can adapt to solve their problems.

Of course you can do all that without Python, so why use it at all? Python has an easier learning curve than C++, and it’s more flexible and immediate,
making it available to a wider set of users. Once a library of extensions is available on a platform, users can call out to that library, mixing and matching components and testing combinations, without the overhead of compiling and linking. (This immediacy should not be underestimated.) And with all its standard library packages, Python can be manipulated to do some pretty amazing things that would be more difficult to realize in C++, like parsing XML documents, setting up computations, etc.

Writing C++ extensions for Python can be learned in an afternoon, especially if there are some examples to follow. The idea is this: you have a C++ class or function, and you’d like to make it callable from the Python interpreter. You’ll need to (I) write some wrapper code in C++, and (II) compile it into a C++ library that the Python interpreter can dynamically load and use. You then typically (III.) write a Python module that mediates between the Python user and this library. This makes life plush for those users who don’t want to be bothered with the details of finding out what’s in the library and how to call it. More importantly, it gives us the chance to check inputs as soon as possible for errors. When a bad pointer is sent to the C++ level, the results are disastrous (a core dump on a good day), so our software can be made more robust if we handle the pointers ourselves.

It takes more care to create a library that is complete: catches exceptions, checks that preconditions and postconditions have been met, and so on. These are more advanced topics, covered in other parts of the book (SOMEDAY NAME A FEW??). Low-level DANSE programmers will be expected to incorporate these techniques into their code, but first things first.

A.1.2 A Little More Detail

Here are those three easy steps again, in slightly more detail:

- Write the bindings. There are three essential components:
  
  1. Wrapper function(s). (One for each function you want callable from Python). The wrapper typically calls a function or a class method, or it creates a heap object. Once you learn how to write one wrapper, you know how to do it, because all wrappers do the same three things.

  2. Method table. (One entry in the table for each function you want callable from Python). The method table tells the Python interpreter

---

1If it’s so easy, why hasn’t someone written a program to write the wrappers automatically? They have! Packages like the Simple Wrapper Interface Generator (SWIG, http://www.swig.org) will do nearly all of the work for you. There are arguments for and against automatic code-generators like SWIG, and I’ve worked both ways. At the moment, writing the bindings is such an easy task, and I do so relatively little of it, that I prefer to do it myself. Others no doubt feel differently, and I have no interest in changing their minds. But even if you’re going to use SWIG, or a similar library, there’s merit in putting in some time writing your own wrappers to learn how and why things get done. Then you can judge well for yourself which approach suits your situation.
A.1. EXTENDING DANSE: WRITING C++ EXTENSIONS TO PYTHON

which C++ functions it can call from the library.

3. init function. (One per module) This is the interpreter's entry point into the C++ library.

These three steps are accomplished with generous aid from the Python-C extension API.

• Compile it. This is slightly platform dependent (Michael Aivazis's system for processing source code removes this platform dependence for UNIX flavors, including cygwin; Windows is in the works). The goal is to compile into a shared object library (unix) or a dynamically linked library (the beloved Windows dll).

• Call it from Python. One typically writes a Python module that acts as a layer between the user and the C++ library. By doing things like providing Python classes that mirror the C++ classes, one can make the experience quite similar. Or dissimilar. The choice is yours.

A.1.3 A Lot More Detail: Wrappers

Every wrapper function does three things:

a) Converts a Python object with the arguments to the C++ function into C++ objects,

b) calls the C++ function,

c) converts the output to a Python object with the result and return it.

Let's first run through these steps with numbers and strings, for which there's an immediate connection between Python and C++ types; we can use a function, PyArg_ParseTuple(), which is built in to the API. Later examples look at tasks like working with C++ class instances and using aggregate Python types, such as dictionaries and lists.

Simple Example: PyArg_ParseTuple, Py_BuildValue
a) Convert arguments from Python to C++

To convert the Python arguments into C++ objects, first define variables of the appropriate type, one for each C++ argument. Then pass the addresses of these variables into PyArg_ParseTuple(). This function takes the args tuple, pulls PyObject's out of it, and converts them to C++ types according to a format string.

Here are some examples:

//One integer:
int a;
int ok = PyArg_ParseTuple(args, "i", &a);
if(!ok) return 0;
//Two integers
int a, b;
int ok = PyArg_ParseTuple(args, "i", &a, &b);
if(!ok) return 0;

//One integer, a string, two doubles
int anint;
char * astring;
double dub1, dub2;
int ok = PyArg_ParseTuple(args, "isdd", &anint, &astring,
                           &dub1, &dub2);
if(!ok) return 0;

A complete list of what can go into the format string is given in the extension
documentation (look in the ext subdirectory of the doc directory in your Python
distribution, or look online at http://python.org/doc/current/ext/ext.html). In the cur-
rent (Oct. ’02) documentation, you want section 1.7, “Extracting Parameters in
Extension Functions”.

`PyArg_ParseTuple()` checks the types of the objects in the tuple `args` against
the types given in the format string. If there’s a discrepancy, it sets the exception
context and returns 0 to our wrapper. If our wrapper detects that, it returns 0 to
the Python interpreter, which understands that to mean failure, and raises an
exception. So if you’re using `PyArg_ParseTuple()`, most of the error checking
is done for you! It is not idiot-proof, but it is smart-friendly.

Once `PyArg_ParseTuple` has successfully returned, do any additional pro-
cessing or checking of the input data that’s appropriate. For instance, Python
does not have an unsigned integer type. With a C++ function that takes an
unsigned int, you’ll need to pass an int to `PyArg_ParseTuple`, check that the int
is greater than -1, and then convert it to an unsigned int.

b) Call your code

It’s your function, call it.

c) Convert output to a Python object, and return it

The API gives a function called `Py_BuildValue`. It returns a pointer to a `PyObject`;
it takes a format string and variables. The format strings are the same as
those used in

```c
PyObject *py_result = Py_BuildValue("i", result);
```
or—

```c
PyObject *py_result = Py_BuildValue("s", astring);
```

etc. What’s returned by `Py_BuildValue` is what the wrapper will return. One
note: don’t return 0; the interpreter will take this as sign of failure. You could,
of course, return a Python integer with value zero: `Py_BuildValue("i", 0)`.
You can specify more than one item to return, in which case `Py_BuildValue` will
place the items in a tuple.

Here’s a complete example of wrapping a function, “bogus”, that takes a
doouble, a string, and an int (in that order) and returns an int. We expect that
the arguments will come from Python in the order string, int, double.
static PyObject *wrap_bogus(PyObject *, PyObject * args){
    //First, get the arguments from Python
    int anint = 0;
    double adub = 0;
    char * astring = 0;
    int ok = PyArg_ParseTuple(args,"sid", &astring, &anint, &adub);
    if(!ok) return 0;
    //do any checking of arguments here
    //Second, make the function call
    int result = bogus(adub, astring, anint);
    //do any extra stuff you want with the return result here
    //Third, build a Python object to return
    return PyBuildValue("i",result);
}

Wrapping classes: PyCObject_FromVoidPtr, PyCObject_AsVoidPtr

a) Creating C++ objects

Wrapping functions is well and good, but what about C++ classes? Python can work with a C++ object by dynamically allocating it and holding onto a pointer. That pointer can be passed back to subsequent wrappers that can then invoke class methods on the object. The pointer is handled in Python by a type PyCObject. To convert a pointer to a PyCObject, use PyCObject_FromVoidPtr(). This API function takes two arguments: the void pointer, and a pointer to a function that takes a void pointer and no return. The purpose of the function is to delete the C++ object when nothing in the Python session is paying attention to it any more.

We get the arguments to the constructor from the Python API, create the object using new, and return a pointer to that object to the interpreter. Use the API function PyCObject_FromVoidPtr() to create the Python object to return to Python. Here’s an example with a real (if dull) class called Numbers:

```cpp
class Numbers
{
public:
  Numbers(int first, double second) : m_first(first), m_second(second){};
  ~Numbers(void){};
  double NumMemberMult(void){return (double)m_first*m_second;}
private:
  int m_first;
  double m_second;
};
```

Here’s a wrapper that creates a new instance of Numbers:

```cpp
PyObject *wrap_new_Numbers(PyObject *, PyObject* args){
    //First, extract the arguments from a Python tuple
    int arg1;
    double arg2;
    int ok = PyArg_ParseTuple(args,"id", &arg1, &arg2);
    if(!ok) return 0;
    //Second, dynamically allocate a new object
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Numbers *newnum = new Numbers(arg1, arg2);
//Third, build a Python object to return
PyObject * py_newnum = PyCObject_FromVoidPtr( static_cast<void *>(newnum),
&DelNumbers);
return py_newnum;
}

Look familiar? This wrapper has essentially the same form as wrap_bogus().
That's because ALL wrappers have essentially this form.

The pointer to the dynamically allocated object, newnum, goes out of scope
as soon as wrap_new_Numbers() returns. The only thing keeping this from being
the mega-classic memory leak is that the Python interpreter has an object that
will keep track of the address of the new object. The interpreter keeps track of
that object for us, and when we lose interest in it (when its reference count goes
to zero), the interpreter will call a C++ function to delete the C++ object. So, the
second slot in PyCObject_AsVoidPtr() is a pointer to a function with return
type void and one void argument. The signature of PyCObject_AsVoidPtr() is:
PyObject * PyCObject_FromVoidPtr( void *, void (*DeleteFunction)(void*));

You must supply the function pointed to (in this example called DelNumbers).
It has the delete corresponding to the new above. Here's it is:

static void DelNumbers(void *ptr)
{
    Numbers * oldnum = static_cast<Numbers *>(ptr);
    delete oldnum;
    return;
}

This strategy can be used for any dynamically allocated resource, such as file
handles or arrays.

Using the object

The user can't actually do anything with the pointer in the Python layer, except
send it back to the C++ layer to do something else: call a C++ class method on
it, or give it as an argument to another function. Here's an example of wrapping
a class method.

#include <Python.h>
PyObject *wrap_Numbers_MemberMult(PyObject *, PyObject* args)
{
    // First, extract the PyCObject that has the
    // Python version of the address
    // from the args tuple
    PyObject *pynum = 0;
    int ok = PyArg_ParseTuple(args, "O", &pynum);
    //"O" is for Object
    if(!ok) return NULL;
    //Convert the PyCObject to a void *
    void * temp = PyCObject_AsVoidPtr(pynum);
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//cast void pointer to Numbers pointer
Numbers * thisnum = static_cast<Numbers *>(temp);
//Can combine the two lines into one:
//Numbers *thisNum = static_cast<Numbers *>(
//  PyCObject_AsVoidPtr(pynum));
//Second, make the function call
double result = thisnum->NumMemberMult();
//Third, build a Python object with the return value
return Py_BuildValue("d",result);
}

All you have to do is fish the PyCObject out of the tuple, extract the void pointer to a C++ variable, cast it to the appropriate type, and use it; then bundle up the result and send it back to the interpreter.

Working with composite types

What if you want to pass a Python list of numbers to a C++ function? There’s no format code to pass to PyArg_ParseTuple for lists. The solution is to extract the list from the args tuple as a PyObject (format code: “O”). Then use the Python-C API functions for working with lists to load the Python list, item-by-item, into a C++ array. Suppose our target function has the signature

    double sum_some_numbers(double *numbers, int array_length)

Here’s some code that could wrap this function. Note that we have to take responsibility for error checking. We can set the exception using PyErr_SetString(). We can verify that Python objects are the type what we think they are by Pytype_Check().

#include <Python.h>
#include <valarray>  // This example uses the std::valarray class.
PyObject *py_sum_some_numbers(PyObject *, PyObject* args)
{
    PyObject *pyList;
    int ok = PyArg_ParseTuple(args, "O", &pyList);
    if(!ok) return 0;
    //Did the user send a Python list?
    int isList = PyList_Check(pyList);
    if(!isList)
    {
        //If not, complain to the Python user and raise an exception:
        PyErr_SetString(PyExc_TypeError, "You fool! That's not a list!");
        return 0;
    }
    //How many items are in the list?
    int numNums = PyList_Size(pyList);
    //Maybe you want to do something here if the size of the list is 0.
    // Now transfer the contents of the list to an array
    // valarray: this C++ standard library class is a great
    // way to avoid memory leaks, and much more
    std::valarray<double> nums(0.0, PyList_Size);
    for(int i=0; i<numNums; i++)
    {
        //Extract the next object in the list:
        //
PyObject *temp = PyList_GetItem(pyList, i);
   // Was the list item a Python float? If not, quit.
   // Note that we don't need to worry about cleaning
   // up the memory: nums will be automatically destroyed
   // when execution exits the scope of nums.
   if( !PyFloat_Check(temp))
   {
      // Just what was in that list?
      PyErr_SetString(PyExc_TypeError, "You fool! That's not a float");
      return 0;
   }
   // Now convert the Python float to a double, and load
   nums[i] = PyFloat_AsDouble(temp);
 }

/*Step 2: Call the function*/
double sum = sum_some_numbers(&nums[0], nums.size());
/*Step 3: return result.*/
return Py_BuildValue("d", sum);

Sometimes you'll want to return a Python object, such as a list. In this
 case, you'll have created a PyObject pointer at some point. You can return
 that pointer directly. If you need to return several Python objects, you can use
 Py_BuildValue() with the "O" format code.

The Python-C api is very complete and well-documented. Similar func-
tions exist for inserting objects into lists, and working with dictionaries, tuples,
modules, and so on. Consult Chapter 7 of the api reference. Hopefully, these
examples have given you the flavor for wrapping C++ functions.

A.1.4 A Lot More Detail: Method Table

The method table is sort of like a table of contents for the Python interpreter.
When it loads the library, it reads the method table to find pointers to the
functions in the library.

static PyMethodDef numbersMethods[] = {
   {"PyNumbers", wrap_new_Numbers, METH_VARARGS,
   "create new Numbers object"},
   {"PyNumbers_MembMult", wrap_Numbers_MemberMult, METH_VARARGS,
   "Multiply Numbers object’s members"},
   {NULL, NULL}
};

The name of the table must match the second argument in the init function.
Each function in the library gets an entry in the table, and each entry has four
components.

1. The string ("PyNumbers" or "PyNumbers_MemberMult") is what you’ll
call from the interpreter.

2. The name wrap_whatever is the name of the corresponding C++ function.
3. METH_VARARGS indicates that one is using the “tuple named args” approach.

4. The final string will appear as the docstring for this function in the Python layer.

The {NULL, NULL} marks the end of the table for the interpreter.

A.1.5 A Lot More Detail: Init Function

The final component in the bindings is the init function. This function has to be named initname_of_library(). If the filename of the library is numbers.dll or numbers.so, this function must be named initnumbers; if it’s _numbers.dll, then this function is named init_numbers. For this example, let’s call the extension library_numbers. The init function’s return type is void, and it takes no arguments. It calls Py_InitModule(), which takes two arguments: a string literal with the name of the library, and the name of the Methods table. It must match the name of the Method table. Also, the function must have C linkage, not C++, meaning the function must be declared extern “C” if you’re using a C++ compiler. Also, on Windows, the function must be exported by the dll, hence the __declspec(dllexport). To keep some platform independence, wrap this in a pre-processor conditional. This function is executed when the library is loaded, so if there’s other initialization steps you need to take, this is the place.

```c
extern "C"
#ifdef WIN32 || _WIN32
__declspec(dllexport)
#endif
void init_numbers(void)
{
    (void) Py_InitModule("_numbers", numbersMethods);
}
```

A.1.6 More Detail: Compile

Compiling the Numbers example under Linux

Here’s a way to compile the Numbers example under Linux. You’ll need files with the Numbers source code (Numbers.cpp and Numbers.h), and the wrapper, Numbers_bindings.cpp. I assume you’re using gcc; if not, you’ll need to modify the compiler flags appropriately.

To compile, compile each source file:

```bash
gcc -I/usr/include/python2.2 -I. -c -fpic Numbers.cpp
gcc -I/usr/include/python2.2 -I. -c -fpic Numbers_bindings.cpp
```

and link them into a shared library:

```bash
gcc Numbers.o Numbers_bindings.o -lm -lc -fpic -shared -o _numbers.so
```
In the compile lines, you’ll of course need to make sure that you’ve pointed to the directory where your Python.h file lives. The “-fpic” specifies position independent code, “-shared” a shared library that can be dynamically linked.

Once you’ve compiled _numbers.so, move it into a place on your system’s PYTHON_PATH and go to town.

Compiling the Numbers example under Windows

Well, of course we want all our ARCS modules to run under Linux/Unix, but for all those times Windows needs a helping hand, here’s how to do it:

For working in Windows, it may be best to use MS Visual C++. Here’s what you’d do to create a project and so on in VC7.

1. From the Start page select New Project.

2. From the Project Types, pick Visual C++ Projects, from the Templates, choose Win32 project. Fill in the name for your library.

3. On the next window, pick Application Settings, and set Application Type to DLL.

4. In the Solution Explorer,
   - get rid of stdafx.cpp
   - right-click on the name of the project, and choose Properties
   - Under the C/C++/General folder, add the additional include directory in which your distribution’s Python.h file resides.
   - Under the Linker/General tab, set the output file to _your_project_name.dll. Note that everything up the “.dll” must be the same as what follows “init” in the init_your_project_name function in the bindings. In the Numbers example, we called that function init_numbers (it’s the last function in the file). So the dll has to be called _numbers.dll. If the name of the file and the init function don’t agree, the interpreter will get lost.

5. In Solution Explorer, look in the file your_project_name.cpp – you’ll need to get rid of that crap about APIENTRY DllMain. Better yet, just get rid of the automatically generated code.

Now fill add the source files Numbers.cpp, Numbers_bindings.cpp, etc. Build, and when the _your_project_name.dll appears in the output directory, move it into your PYTHON_PATH, and enjoy.

A.1.7 More Detail: Call it from Python

In keeping with the Numbers example used above, here’s a Python class called Numbers. It’s a “shadow class” for the C++ Numbers class. Pretty much everything you can do with the C++ class can also be done with the Python.
import _numbers

class Numbers:
    def __init__(self, an_int, a_float):
        # Check an_int
        if type(an_int) != type(1):
            raise TypeError, "Fool! an_int must be an integer"
        # Check a_float
        if type(a_float) != type(3.14159):
            raise TypeError, "Fool! a_float must be a float"
        self.this = numbers.PyNumbers(an_int, a_float)

    def MemberMult(self):
        return numbers.PyNumbers_MembMult(self.this)

If this were saved in a module named numbers.py (stored somewhere on
your PYTHON_PATH), then an interpreter command line session might look like

>>> import numbers
>>> n = numbers.Numbers(2,3.14)
New Numbers object created
>>> n.MemberMult()
6.280000000002
>>> n = 1

The variable n is the only thing keeping track of the Numbers object. When we
reassign n, the interpreter calls our bit of code from the C++ library that deletes
the object, preventing the resource leak.

So now the Python user has something like “interpreted C++”. Pretty cool,
right?

A.1.8 More realistic example

Let’s wrap a function from the NeXus API, NXopen, with all the bells and
whistles. This wrapper is part of a larger library that wraps the entire NeXus C
API. The latter, of course, is a simplified interface to the HDF libraries that the
NeXus standard currently uses.

Since this is one of several dozen libraries, we split the bindings up into
several files. The wrappers live in pairs of files, one pair for each major func-
tional group: file.h/file.cc for file level operations, group.h/group.cc for group
level, etc. Only two file level operations from the original NeXus C API are in
the Python NeXus API: NXopen and NXflush. Therefore, file.h looks like the
following:

#ifndef NeXus_file_h
#define NeXus_file_h

// Python bindings for file level operations:
// NXopen
extern char pyNeXus_NXopen__name__[];
extern char pyNeXus_NXopen__doc__[];
extern "C" PyObject * pyNeXus_NXopen(PyObject *, PyObject *args);
// NXflush
extern char pyNeXus_NXflush__name__[];
extern char pyNeXus_NXflush__doc__[];
extern "C" PyObject * pyNeXus_NXflush(PyObject *, PyObject *args);
#endif

This file will be included into the file that contains the methods table. Note that in addition to the actual wrapper functions, we declare two char arrays for the name and docstring. Defined in file.cc, these variables keep the methods table neat (the docstrings in particular may get lengthy).

How do we implement the wrapper for NXopen? Begin with the signature of NXopen, located in napi.h in the NeXus source distribution:

```c
NX_EXTERNAL NXstatus CALLING_STYLE NXopen(CONSTCHAR * filename, NXaccess access_method, NXhandle* pHandle);
```

The various types NXstatus, CONSTCHAR, NXaccess, and NXhandle are defined in the NeXus C API header files; we need to track them down so we can know what the Python user will have to give us in order to satisfy the function call.

Searching through napi.h, we learn that NXstatus is a typedef for int, CONSTCHAR is a typedef for char,\(^2\) NXaccess is an enumeration with members like NXACC_READ, and NXhandle is a typedef for void *. We can’t map the NXaccess enumeration directly into Python types, so we’ll expect a string from the user; by comparing values of the string we’ll assign the proper value to an NXaccess variable. As for the CONSTCHAR * filename, we can derive that directly from a Python string.

What about the NXhandle * pHandle? This is interesting. Let’s open up the NeXus C API source code and find out what exactly is done with that void pointer. Reading through napi.c, we discover that NXopen dynamically allocates a structure of type NexusFunction, and that the NXhandle we pass to NXopen becomes a handle to that object. In other words, pHandle is an output of NXopen, not an input. We don’t need to trouble the Python user with giving us a NXhandle object; instead, we’ll give them one.

Expecting two inputs, both Python strings, we write the first few lines of the wrapper as follows:

```c
PyObject * pyNeXus_NXopen(PyObject *, PyObject *args) {
    char *filename = 0;
    char *acc_method = 0;
    int ok = PyArg_ParseTuple(args, "ss", &filename, &acc_method);
    if(!ok) return 0;
```

and we’ve got what we needed from the Python user. Or do we? We owe it to ourselves to check the inputs from the user. This could be done either here or in the Python layer. Checking that the inputs are Python strings has already been performed by PyArg_ParseTuple. There are two questions about the filename:

\(^2\)Prefering to not mislead the readers of our code, we might have chosen CONSTCHAR as typedef const char.
A.1. EXTENDING DANSE: WRITING C++ EXTENSIONS TO PYTHON

is it the name of an actual file, and is it the name of an appropriate HDF file?
The first question is easily answered in a platform independent way in Python.
The second can only be answered by essentially doing what we do anyway in
NXopen. We need to inspect the value of acc_method anyway to convert it to
the appropriate member of the NXaccess enumeration. We can do that easily
using the C++ standard library class string:

```cpp
// Check access_method:
std::string methstring(acc_method);
NXaccess mode;
if (methstring == "r") mode = NXACC_READ;
else if(methstring == "rw") mode = NXACC_RDWR;
else if(methstring == "c") mode = NXACC_CREATE;
else if(methstring == "c4") mode = NXACC_CREATE4;
else if(methstring == "c5") mode = NXACC_CREATE5;
else {
    std::string errstr("NeXus_bindings.cc pyNeXus_NXopen(): ");
    errstr += "unrecognized access_method string.");
    PyErr_SetString(PyExc_ValueError, errstr.c_str() );
    return 0;
}
```

Here we’ve used PyErr_SetString to set the (Python) exception context if
we don’t recognize what the Python user wants, and then forced the interpreter
to raise the exception by returning 0.

At this point, we’re nearly ready to call NXopen—we only need to declare a
variable of type NXhandle:

```cpp
NXhandle handle;
NXstatus status = NXopen(filename, mode, &handle);
if(status != NX_OK)
{
    std::string errstr("NeXus_bindings.cc pyNeXus_NXopen(): ");
    errstr += "NXopen failed.");
    PyErr_SetString(PyExc_IOError, errstr.c_str() );
    return 0;
}
```

If we get to this stage, we’re almost ready to return. Since we are not content
with any NXstatus but NX_OK, all we need to do is to return the NXhandle ini-
tialized by NXopen. But there’s another issue here: NXopen allocates a resource,
so we must release that resource when we’re finished. As with the discussion
of allocating C++ objects (§ A.1.3), we can use PyCObject_FromVoidPtr and a
helper function to release the resource when the Python user is finished:

```cpp
return PyCObject_FromVoidPtr(handle, pyNeXus_NXclose);
}
```

The helper function (declared static to avoid ) wraps NXclose, which in turn
disposes of the resources allocated in NXopen:
Finally, we define the name and docstring for this function. The name should be something sensible, while the docstring is an opportunity to incorporate a little documentation.

```c
char pyNeXus_NXopen__name__[] = "nxopen";
char pyNeXus_NXopen__doc__[] = "Open a nexus file\n" "2 Arguments: filename, access_method\n" "Input: \n" " filename (Python string)\n" " access_method (Python string)\n" " allowed values: r (read only)\n" " rw (read/write)\n" " c (create)\n" " c4 (create HDF 4)\n" " c5 (create HDF 5)\n" "Output: (return)\n" " PyCObject holding pointer to NXhandle\n" "Exceptions: ValueError, IOError\n";
```
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