III.C. CRYSTALS, SYMMETRY, AND DIFFRACTION

Biomacromolecules quite often occur naturally or in vitro as organized structures composed of subunits arranged in a symmetrical way. Such structures are readily studied by diffraction methods. The fundamental concepts concerning crystalline matter, symmetry relationships, and diffraction theory form a basic framework for understanding the principles and practice of image processing and interpretation of structural results. The main concepts, summarized here, are presented in more detailed, introductory form in excellent texts such as Eisenberg and Crothers (1979), Glusker and Trueblood (1972), Holmes and Blow (1965), and Wilson (1966). For additional references, see the reading list.

III.C.1. Definitions of Terms (compiled from Glusker & Trueblood, and Eisenberg & Crothers)

Asymmetric Unit: Part of the symmetric object from which the whole is built up by repeats. Thus, it is the smallest unit from which the object can be generated by the symmetry operations of its point group. Fig.III.3.

Bravais lattice: One of the 14 possible arrays of points repeated periodically in 3D space in such a way that the arrangement of points about any one of the points in the array is identical in every respect to that about any other point in the array. Fig.III.2.

Center of Symmetry (or Center of Inversion): A point through which an inversion operation is performed, converting an object into its mirror image. Fig.III.10.

Crystal: A solid having a regularly repeating internal arrangement of its atoms. Figs.III.5,6,7.

Crystal lattice: Crystals are composed of groups of atoms repeated at regular intervals in three dimensions with the same orientation. For certain purposes it is sufficient to regard each such group of atoms as replaced by a representative point; the collection of points so formed is the space lattice or lattice of the crystal. Each crystal lattice is a Bravais lattice. Figs.III.1,2,4,7.

Crystal Structure: The mutual arrangement of atoms, molecules or ions that are packed together in a lattice to form a crystal. Figs.III.5,6,7.

Crystal System: The seven crystal systems, best classified in terms of their symmetry, correspond to the seven fundamental shapes for unit cells consistent with the 14 Bravais lattices. Fig.III.2 and Table III.2.

Glide Plane: A symmetry element for which the symmetry operation is reflection across the plane combined with translation in a direction parallel to the plane. Figs.III.28 and III.29.

Inversion: A symmetry operation in which each point of an object is converted to an equivalent point by projecting through a common center (called center of inversion or center of symmetry) and extending an equal distance beyond this center. If the center of symmetry is at the origin of the coordinates, every point \(x, y, z\) becomes \(-x, -y, -z\). It converts an object or a structure into one of opposite "handedness", related to the first as is any object and its mirror image. Figs.III.10 and III.12.

Lattice: Rule for translation. Fig.III.1 and Tables III.1 and III.2.

Mirror Plane: A symmetry element for which the corresponding symmetry operation resembles reflection in a mirror coincident with the plane. It converts an object or a structure into one of opposite "handedness", related to the first as is any object and its mirror image. Figs.III.10,11.

Motif: Object that is translated. Figs.III.3,4,7.

Plane Group: Symmetry of a two-dimensional structure. There are 17 plane group symmetries possible (only 5 for biological structures). Figs.III.6,23-27.

Point Group: The collection of symmetry operations that describe the symmetry of an object about a point. Figs.III.8-17.

Reciprocal Lattice: The lattice with axes \(a^*, b^*, c^*\) related to the crystal lattice or direct lattice (with axes \(a, b, c\)) in such a way that \(a^*\) is perpendicular to \(b\) and \(c\); \(b^*\) is perpendicular to \(a\) and \(c\); and \(c^*\) is perpendicular to \(a\) and \(b\). Rows of points in the direct lattice are normal to nets of the reciprocal lattice, and vice versa. For a given crystal, the direct lattice and reciprocal lattice...
have the same symmetry. The repeat distance between points in a particular row of the reciprocal lattice is inversely proportional to the interplanar spacing between the nets of the crystal lattice that are normal to this row of points.

**Rotation Axis**: An axis of symmetry for an object, such as a crystal. When the object is rotated by \((360/n)°\) about an \(n\)-fold rotation axis, the new orientation is indistinguishable from the original one. Figs.III.9,12.

**Rotary-Inversion Axis**: An axis for which the corresponding symmetry operation is a rotation by \((360/n)°\) combined with inversion through a center of symmetry lying on the axis. Figs.III.12,13.

**Screw Axis**: An axis (designated \(n_m\)) for which the corresponding symmetry operation is a rotation about the axis by \((360/n)°\) followed by a translation parallel to the axis by \(m/n\) of the unit cell length in that direction. Figs.III.19-21.

**Space group**: A group or array of operations consistent with an infinitely extended regularly repeating pattern. It is the symmetry of a 3D structure. There are 230 space group symmetries possible (only 65 for biological structures).

**Symmetry**: An object is symmetric if some spatial manipulation of it results in an indistinguishable object. A symmetric object can be superimposed on itself by some operation.

**Symmetry Element**: Geometrical entity (such as a line, a point or a plane) about which a symmetry operation is carried out.

**Symmetry Operation**: The operation that leads to superimposition of an object on itself (i.e. results in moving the object to a position in which its appearance is indistinguishable from its initial appearance). Symmetry operations include rotation, inversion, reflection and translation.

**Translation**: A motion in which all points of an object move in the same direction, that is, along the same or parallel lines. Fig.III.18.

**Unit Cell**: The fundamental portion of a crystal structure that is repeated infinitely by translation in three dimensions. It is characterized by three vectors \(a, b, c\), not in one plane, which form the edges of a parallelepiped. Figs.III.1,2,7.

### III.C.2. Crystals

A crystal is a regular arrangement of atoms, ions, or molecules, and is conceptually built up by the continuing translational repetition of some structural pattern. This pattern, or **unit cell**, may contain one or more molecules or a complex assembly of molecules. In three dimensions (3D), the unit cell is defined by three edge lengths \(a, b, c\) and three interaxial angles \(\alpha, \beta, \gamma\). The different **3D crystal systems** (triclinic, monoclinic, orthorhombic, tetragonal, trigonal, hexagonal and cubic) arise from the seven fundamental unit cell shapes. The simplest, triclinic, is a parallelepiped, with no restrictions on cell lengths or angles. A cubic cell has equal \((a=b=c)\) and orthogonal \((\alpha=\beta=\gamma=90°)\) edges.

### III.C.3. Lattices

A lattice is a mathematical formalism that defines an infinite array of imaginary points: each point in the lattice is identical to every other point. That is, the view from each point is identical with the view in the same direction from any other point. This condition is not obeyed at the boundary of a finite, but otherwise perfect crystal. The crystal **structure** and the crystal **lattice** are NOT equivalent: the structure is an array of objects whereas the lattice is an array of imaginary, infinitely small points. A 2D lattice is defined by two translations, \(a, b\) and two axes at an angle \(\alpha\) to each other. A 3D lattice is defined by three translations, \(a, b, c\), and three axes at angles \(\alpha, \beta, \gamma\) to each other.

Crystal lattices (2D or 3D) may be **primitive**, with one lattice point per unit cell, or **centered**, containing two or four points per cell. There are **four 2D lattice systems** (Table III.1) which are subdivided into a total of five 2D lattices (Fig. III.1). There are **seven 3D crystal systems** corresponding to the seven basic space-filling shapes that unit cells can adopt.
These are subdivided into a total of fourteen, so-called Bravais lattices (Table III.2 and Fig. III.2). For example, the cubic crystal system includes three Bravais lattices: primitive \((P)\), body centered \((I)\), and face centered \((F)\).

<table>
<thead>
<tr>
<th>Lattice system</th>
<th>Lattice type</th>
<th>Conventional representation</th>
<th>Representative points</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oblique</td>
<td>(P)</td>
<td>(a = b) (\varphi &gt; 90^\circ)</td>
<td>((0, 0))</td>
</tr>
<tr>
<td>Square</td>
<td>(P)</td>
<td>(a = b) (\varphi = 90^\circ)</td>
<td>((0, 0))</td>
</tr>
<tr>
<td>Hexagonal</td>
<td>(P)</td>
<td>(a = b) (\varphi = 120^\circ)</td>
<td>((0, 0))</td>
</tr>
<tr>
<td>Rectangular</td>
<td>(P)</td>
<td>(a = \frac{b}{2}) (\varphi = 90^\circ)</td>
<td>((0, 0), \left(\frac{1}{2}, \frac{1}{2}\right))</td>
</tr>
</tbody>
</table>

Table III.1. Plane lattices. (From Sherwood, p.70).

<table>
<thead>
<tr>
<th>Crystal system</th>
<th>Lattice type</th>
<th>Conventional representation</th>
<th>Representative points</th>
</tr>
</thead>
<tbody>
<tr>
<td>Triclinic</td>
<td>(P)</td>
<td>(a = b = c) (\alpha \neq \beta \neq \gamma)</td>
<td>((0, 0, 0))</td>
</tr>
<tr>
<td>Monoclinic</td>
<td>(P)</td>
<td>(a = b \neq c) (\alpha = \gamma = 90^\circ \neq \beta)</td>
<td>((0, 0, 0))</td>
</tr>
<tr>
<td>Orthorhombic</td>
<td>(P)</td>
<td>(a = b \neq c) (\alpha = \beta = \gamma = 90^\circ)</td>
<td>((0, 0, 0), \left(\frac{1}{2}, \frac{1}{2}, \frac{1}{2}\right))</td>
</tr>
<tr>
<td>Tetragonal</td>
<td>(P)</td>
<td>(a = b \neq c) (\alpha = \beta = \gamma = 90^\circ)</td>
<td>((0, 0, 0))</td>
</tr>
<tr>
<td>Cubic</td>
<td>(P)</td>
<td>(a = b = c) (\alpha = \beta = \gamma = 90^\circ)</td>
<td>((0, 0, 0))</td>
</tr>
<tr>
<td>Trigonal</td>
<td>(P)</td>
<td>(a = b = c) (\alpha = \beta = \gamma &lt; 120^\circ \neq 90^\circ)</td>
<td>((0, 0, 0))</td>
</tr>
<tr>
<td>Hexagonal</td>
<td>(P)</td>
<td>(a = b \neq c) (\alpha = \beta = 90^\circ)</td>
<td>((0, 0, 0))</td>
</tr>
</tbody>
</table>

Table III.2. The fourteen Bravais lattices. (From Sherwood, p.73).
III.C.4. Crystal Structure

The crystal structure is built by placing a motif at every lattice point. The motif is the object that is translated, and may be asymmetric (e.g. a single polypeptide chain: Fig.III.3) or symmetric (i.e. containing two or more symmetrically arranged subunits). The crystal structure, crystal lattice, and motif are all restricted in the symmetries they can display, but biomacromolecular assemblies themselves are not restricted in the sense that they may display additional internal (non-crystallographic) symmetry. From this emerges the corollary that an asymmetric unit of the crystal structure may itself contain a symmetrical arrangement of identical, asymmetric molecules. The following introduction to symmetry will help clarify this.
Fig. III.4. Lattices and motifs. Each of the diagrams of Fig.III.4 may be reproduced by associating the motifs (a), (b), or (c) with the rectangular array of lattice points. (From Sherwood, p. 61)

Fig. III.6. The five two-dimensional plane groups for biological structures. The asymmetric unit is a triangle in every case; the motif is some group of triangles (1,2,3,4, or 6). (From Eisenberg and Crothers, p. 786)

Fig. III.7. (a) The generation of a 2D "crystal structure" from a lattice and a structural motif. The replacement of each lattice point by an apple leads to a 2D structure. There are many ways in which unit cells may be chosen in a repeating pattern. Various alternative choices are shown, each having the same area despite varying shape (note that the total content of any chosen unit cell is one apple). Infinite repetition in 2D of any one of these choices for unit cell will reproduce the entire pattern. (b) Perspective view of a triclinic lattice. To see the perspective, lines that define imaginary edges of a unit cell join the lattice points. One unit cell is shaded; it could have been chosen with a different shape (but the same volume), as in the 2D example in (a). (From Glusker and Trueblood, p. 10)
III.C.5. Symmetry

Biological objects may display symmetry about a point or along a line. An object is **symmetrical** if it is indistinguishable from its initial appearance when spatially manipulated. (Again, we ignore the loss of translational symmetry at the boundary of finite, crystalline objects.)

a. Symmetry operators

There are **four types of symmetry operations** that lead to superimposition of an object on itself: **rotation**, **translation**, **reflection** and **inversion**. A symmetry element is a geometrical entity such as a point, line, or plane about which a symmetry operation is performed. The symmetry of any object can be described by some combination of these symmetry operations. The symmetry of any aggregate or crystal of a biological molecule is only described by rotation and/or translation operations. This is because, for example, biological protein molecules mainly consist of L-amino acids, hence, reflection or inversion symmetries are not allowed.

b. Asymmetric unit

Symmetry operations give rise to other groups of atoms (the **asymmetric unit**) which are in equivalent, general positions. The number of equivalent general positions related by the symmetry operations equals the number of asymmetric units in the unit cell. The number of asymmetric units may be less than, equal, or greater than the number of molecules in the unit cell. If the number of asymmetric units is equal to or less than the number of molecules in the cell, than the molecule either contains no symmetry or it contains **non-crystallographic symmetry** (i.e. symmetry that is not contained within the allowed lattice symmetries). If the number of asymmetric units is greater than the number of molecules in the cell, than the molecules must occupy special positions and possess the appropriate symmetry element of the space group (see Sec. III.C.5.e).

c. Point groups (Symmetry about a single point)

1) Schoenflies and Herman-Mauguin point group notations

A **point group** is a collection of symmetry operations that define the symmetry about a point. Two systems of notations are used for point groups: i) the **S** or **Schoenflies notation** (capital letters; mainly used by spectroscopists) and ii) the **H-M** or **Hermann-Mauguin symbol** (an explicit list of the symmetry elements, commonly preferred by crystallographers).

2) Types of point group symmetry operators

The four types of symmetry about a point are rotational symmetry, mirror symmetry, inversion symmetry, and improper rotations. Each of these is described below with accompanying illustrations.

**Rotational symmetry** (**n**): the object appears identical if rotated about an axis by \( \alpha = \frac{360}{n} \) degrees = \( 2\pi/n \) radians. The only allowed \( n \)-fold axes for **crystal lattices** are \( n = 1,2,3,4, \) and 6 since lattices must be space filling. Figs.III.8,9,12.

![Fig.III.8. Two-, three-, four-, five-, and six-fold rotational symmetries. (From Bernal, pp.45,48,50,35, and 52)](image-url)
Fig. III.9. A four-fold rotation axis, parallel to \( c \) and through the origin of a tetragonal unit cell (a=b), moves a point at \( x, y, z \) to a point at \( (y, -x, z) \) by a rotation of 90 about the axis. The sketch on the right shows four equivalent points resulting from successive rotations. (From Glusker and Trueblood, p. 72)

**Mirror (reflection) symmetry \((m)\):** each point in the object is converted to an identical point by projecting through a mirror plane and extending an equal distance beyond this plane. Figs.III.10 and III.11

**Inversion symmetry \((i)\):** each point in the object is converted to an identical point by projecting through a common center and extending an equal distance beyond this center. Objects with \( i \) symmetry are said to be **centrosymmetric**. Fig.III.10.

Fig. III.10. (Left) A mirror symmetry operation. (Right) An inversion symmetry operation. (From Buerger, p. 8)
Improper rotations: rotations followed by $m$ or $i$. These include the rotoinversion ($n$ followed by $i$) and rotoreflection ($n$ followed by $m$). The only inversion axes for crystal lattices are $1, 2, 3, 4, 6$. Figs. III.12 and III.13.

Fig. III.11. Additional examples of mirror symmetry operations. (Right) $m_y$ and $m_x$. (From Bernal, pp. 26, 27, and 34)

Fig. III.12. Crystallographic rotation (a) and inversion-rotation (b) symmetry axes and their action on an asymmetric figure - a tetrahedron. (From Vainshtein, p. 67)
Fig. III.13. The operation $2$, a two-fold rotary-inversion axis parallel to $\mathbf{b}$ and through the origin, converts a point at $x, y, z$ to a point at $x, -y, z$. This is the result of, first, a two-fold rotation about an axis through the origin and parallel to $\mathbf{b}$ $(x, y, z$ to $-x, y, -z$) and then an inversion about the origin $(x, -y, z$ to $x, -y, z)$. This is the same as the effect of a mirror plane perpendicular to the $\mathbf{b}$ axis. Note that a right hand has been converted to a left hand. (From Glusker and Trueblood, p. 72)

3) Types of point groups

Cyclic, dihedral, and cubic (tetrahedral, octahedral and icosahedral) point groups define the collection of symmetry operations about a point. Klug (1969) and others (e.g. Wilson, 1966; Glusker and Trueblood, 1972; Eisenberg and Crothers, 1979; Bernal, Hamilton and Ricci, 1972) describe this and other aspects of symmetry in detail.

**Cyclic point groups**

These contain a single $n$-fold axis of rotation, where $n$ can be any positive integer and also may contain one or more planes of reflection (Fig.III.15, next page). Point groups that contain only an $n$-fold axis of rotation are given the symbol $n$ in the H-M system and $C_n$ in the S system ($C$ stands for cyclic). For example, the double-disk structure of tobacco mosaic virus (TMV) stacked disk aggregates contains 34 subunits (polypeptide chains) arranged with $C_{17}$ symmetry (S notation) (Fig.III.14). Non-biological molecules can also have mirror planes of symmetry either parallel (e.g. $nm$ or $nmm$ in the H-M notation or $C_{nv}$ in the S notation where $v$ stands for vertical) or perpendicular (e.g. $n/m$ in the H-M notation or $C_{nh}$ in the S notation where $h$ stands for horizontal) to the $n$-fold axis of symmetry.

![Double disk of protein from tobacco mosaic virus](image)

Fig.III.14. The TMV stacked disk structure with $C_{17}$ symmetry. (From Eisenberg and Crothers, p.757)
Fig. III.15. Examples of cyclic point groups. (From left to right and top to bottom) 2, 2mm, 3, 3m, 4, 4mm, 6, and 6mm (From Bernal, pp. 45, and 47-53)
Dihedral point groups

Dihedral point groups have axes of rotation at right angles to each other. These point groups consist of an \( n \)-fold axis perpendicular to \( n \) 2-fold axes. Most oligomeric enzymes display dihedral symmetry (Matthews and Bernhard, 1973). For example, the enzyme ribulose bisphosphate carboxylase/oxygenase (RuBisCO) has \( D_4 \) symmetry (422 in H-M notation). The number of asymmetric units in the point group \( D_n \) is \( 2n \), thus RuBisCO has eight asymmetric units. In this particular enzyme, each asymmetric unit contains two polypeptide chains: a large (\( \sim 55 \) kD), catalytic subunit and a small (\( \sim 15 \) kD) subunit whose function (regulatory?) is unknown.

Cubic point groups

The essential characteristic is four 3-fold axes arranged as the four body diagonals (lines connecting opposite corners) of a cube. The three cubic point groups that biological molecules can occupy are \( T \) (tetrahedral = 23 in H-M notation), \( O \) (octahedral = 432) and \( I \) (icosahedral = 532) (Fig.III.16). The tetrahedral point group contains 12 asymmetric units. Aspartate-ß-decarboxylase is presumed to display this point group symmetry (Eisenberg and Crothers, 1979). Dihydrolipoyl transsuccinyllase contains 24 asymmetric subunits arranged with octahedral (432) symmetry. There are a large number of plant, animal, and bacterial viruses, each containing 60 asymmetric units, which display icosahedral (532) symmetry (Fig.III.17). In most cases these spherical viruses contain a multiple of 60 copies of chemically identical or distinct protein or glycoprotein subunits. In those cases where the asymmetric unit contains more than one "subunit", not all subunits are equivalently arranged. Instead they occupy quasi-equivalent positions.

4) Lattice restrictions and non-crystallographic symmetry

The crystal structure and crystal lattice may only contain one-, two-, three-, four-, or six-fold rotational symmetry axes (because the crystal lattice must be space filling) though the motif can have additional symmetries. For example, the 34 subunits in the disc structure of TMV are arranged about a 17-fold axis of rotation (Fig.III.14). The TMV disc forms true 3D crystals and has been studied by X-ray crystallography. In the crystal, the disc occupies a general position in the unit cell, and therefore displays non-crystallographic symmetry. Many of the small, spherical viruses are icosahedral (cubic point group) and they contain symmetry elements compatible with allowed lattice symmetries, and crystallize and display crystallographic as well as non-crystallographic symmetry.
d. **Translational symmetry** (Symmetry along a line)

1) Repetition in one dimension

**Translation** is the symmetry operation of shifting an object a given distance, say \( t \), in a given direction, say the \( x \) direction, as illustrated in Fig.III.18 (one-dimensional crystal of right feet). The group of feet can be superimposed on itself if it is shifted in the \( x \) direction by the translation \( t \). In this one-dimensional crystal, each foot occupies one unit cell and the distance \( t \) is the **unit cell edge** of the crystal.

![Fig.III.18. A one-dimensional 'crystal' of right feet.](From Bernal, p.27)

2) **Screw axes**

A **screw axis** combines translation and rotation operations to produce a structure with helical symmetry (Figs. III.19-21). Screw axes are symmetry elements of crystals that are helices with an **integral** number of asymmetric units per turn of the helix. An \( n_m \) screw axis combines a rotation of \( 2\pi/n \) radians about an axis, followed by a translation of \( m/n \) of the repeat distance (unit cell edge). The screw axes found in crystals include \( 2_1, 3_1, 4_1, 4_2, 4_3, 6_1, 6_2, 6_3, 6_4, \) and \( 6_5 \). A crystal lattice only accommodates an integral number of asymmetric units per turn of the helix, although this need not apply to helices in general.

![Fig.III.19. A two-fold screw axis, 2_1, parallel to \( b \) and through the origin, which combines both a two-fold rotation \((x, y, z)\) to \((-x, y, -z)\) and a translation of \(b/2\) \((-x, y, -z)\) to \((-x, 1/2+y, -z)\). A second screw operation will convert the point \(-x, 1/2+y, -z\) to \(x, 1+y, z\), which is the equivalent of \(x, y, z\) in the next unit cell along \(b\). Note that the left hand is never converted to a right hand.](From Glusker and Trueblood, p. 73)
Fig. III.20. Some crystallographic four-fold screw axes showing two identity points for each. Note that the effect of 4₁ on a left hand is the mirror image of the effect of 4₃ on a right hand. The right hand has been moved slightly to make this relation obvious. (From Glusker and Trueblood, p. 74)

Fig. III.21. Crystallographic screw axes and their action on an asymmetric tetrahedron. (From Vainshtein, p. 69)
Aggregates such as actin thin filaments, microtubules, and the TMV particle are helical structures, all of which display symmetry along a line. For example, one turn of the basic helix of the TMV rod contains 16.33 protein subunits (Fig.III.22). The true repeat in the structure is, therefore, three turns of the basic helix containing 49 subunits. Most helical biological aggregates do not form 3D crystals suitable for diffraction studies, not because of strict symmetry constraints (they could occupy non-crystallographic positions), but rather as a consequence of their shape and large size.

![Fig. III.22. Drawing of part of the helical structure of tobacco mosaic virus. Each shoe-shaped protein subunit is bound to three RNA nucleotides. Part of the RNA chain is shown stripped of its protein subunits in a configuration if could not maintain without them. Each turn of 16.3 protein subunits is closely related to the disk structure of Fig.III.15. (From Eisenberg and Crothers, p. 782)](image)

e. Plane groups and space groups

The symmetry of a structure is described by a plane group if it is 2D or by a space group if it is 3D. All possible crystal symmetries are generated by combining all types of lattice symmetries with all types of motif symmetries. If the internal structure of the crystal is considered, additional symmetry exists due to the presence of screw axis and glide plane symmetries. This leads to 17 possible plane groups in two-dimensions and 230 space groups in three-dimensions. Thus, there are only 17 ways in which to generate a two-dimensional regular pattern from a motif associated with a two-dimensional lattice and 230 ways in which to generate a 3D regular pattern from a motif associated with a 3D lattice. Volume I of the International Tables for X-ray Crystallography (1969) provides descriptions and formulae for all the plane groups and space groups.

Only five plane groups and 65 space groups are compatible with the enantiomorphic biological structures. Figures III.6 and III.23 show some examples of 2D plane groups. The initial stage of most X-ray crystallographic structure analyses involves the space group determination. When this is known, the number of asymmetric units in the unit cell is also known, thus it is often possible to learn the packing and number of molecules within the unit cell, and prove if the molecules are symmetric. Symmetry operations in the unit cell give rise to systematic absences in the diffraction patterns and these absences often prove useful for determining the correct space group. The number of molecules per unit cell or per asymmetric unit is usually deduced from estimates of the molecular weight and measurements of the crystal density and unit cell volume. In many instances, image processing of electron micrographs of periodic biological specimens provides an objective means for determining or verifying these types of structural information.

Placing a motif at every point of a lattice generates a periodic structure. The lattice is a rule for translation and the motif is the object that is translated. Figures III.18 and III.24-27 illustrate examples of both one- and two-dimensional crystal structures. Notice that the motif does not always have to be an asymmetric object such as a right foot. In biological crystals, the motif often has the symmetry of one of the point groups or one of the screw axes. In such cases the periodic structure contains translational symmetry plus rotational (or reflection or inversion in some, mostly small biological molecules) symmetry. The periodic structure can be thought of as being built up in two steps. First a motif is generated from the asymmetric unit by the symmetry operations of the point group. Second, the structure is generated from the motif by the translational symmetry operations of the lattice:
Fig. III.23. The rotational symmetry elements (a) 1 (b) 2 (c) 4 (d) 3 and (e) 62. A single unit cell is outlined by a double set of lines. Other unit cells are shown with single lines. The asymmetric unit is shown shaded. (From Blundell and Johnson, p. 87)
Asymmetric unit | point-group symmetry | motif symmetry | lattice symmetry | structure

Glide plane symmetry (Figs. III.28-III.29) is produced by a translation followed by a mirror operation (or vice versa). Biological molecules do not, in general, display glide plane symmetries because they do not exist in enantiomorphous pairs. However, note that biological molecules (or crystals) when viewed in two-dimensions (i.e. in projection) can display mirror symmetry.
f. Examples of symmetrical biological molecules

1) Helical symmetry
- Actin filament
- Chromatin fibers
- Bacterial pili
- Tobacco mosaic virus
- T4 bacteriophage sheath (extended or contracted configuration)

2) Point group symmetry

<table>
<thead>
<tr>
<th>MOLECULE/AGGREGATE</th>
<th>S</th>
<th>H-M</th>
<th># ASU</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asymmetric aggregates: e.g. ribosome (monomer)</td>
<td>C1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Fibrous molecules: e.g. fibrinogen</td>
<td>C2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Enzymes:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>lactate dehydrogenase</td>
<td>D2</td>
<td>222</td>
<td>4</td>
</tr>
<tr>
<td>catalase</td>
<td>D2</td>
<td>222</td>
<td>4</td>
</tr>
<tr>
<td>aspartate transcarbamylase</td>
<td>D3</td>
<td>32</td>
<td>6</td>
</tr>
<tr>
<td>ribulose bisphosphate carboxylase/oxygenase</td>
<td>D4</td>
<td>422</td>
<td>8</td>
</tr>
<tr>
<td>glutamine synthetase</td>
<td>D6</td>
<td>622</td>
<td>12</td>
</tr>
<tr>
<td>aspartate-b-decarboxylase</td>
<td>T</td>
<td>23</td>
<td>12</td>
</tr>
<tr>
<td>dihydrolipoyl transsuccinylase</td>
<td>O</td>
<td>432</td>
<td>24</td>
</tr>
<tr>
<td>Spherical viruses: e.g. polyoma, polio, rhino, tomato bushy stunt, human wart, etc.</td>
<td>I</td>
<td>532</td>
<td>60</td>
</tr>
</tbody>
</table>
3) **Plane group symmetry** (two-dimensional crystals)
   - Bacterial cell walls (*e.g.* Bacillus brevis T layer)
   - Bladder luminal membrane
   - Gap junctions
   - Purple membrane

4) **Space group symmetry** (3D crystals)
   - Various intracellular inclusions
   - Various *in vitro* grown crystals suitable for x-ray crystallography

III.C.6. **Diffraction**

Diffraction methods, including x-ray, neutron, electron, and optical diffraction, provide a powerful way to study molecular structure. The ultimate goal is to understand the chemical properties of molecules by determining their atomic structure (*i.e.* the types of chemical bonds - ionic, covalent, or hydrogen-, their lengths and angles, Van der Waals radii, rotations about single bonds, etc.). Presently, only x-ray and neutron diffraction techniques are routinely capable of revealing the arrangement of atoms in molecular structures. In 1912 von Laue predicted that x-rays should diffract from crystals like light from a diffraction grating. Friedrich and Knipping later verified this prediction experimentally. W. L. Bragg developed the concept of diffraction from crystal planes (Sec. III.C.6.e) and that the diffraction pattern could be used to reveal atomic positions in crystals. The physical principles of x-ray diffraction form the fundamental basis of Fourier image processing techniques.

a. **Introduction to diffraction theory** (see also pp.7-9, Sec.I.A.3.c for review)

Diffraction is the **non-linear propagation of electromagnetic radiation** and occurs when an object scatters the incident radiation. The radiation scattered from different portions of the object interfere both constructively and destructively, producing a diffraction pattern that can be recorded on a photographic emulsion. In an electron microscope, electrons are scattered both by the electrons (inelastic scatter) and nuclei (elastic scatter) of the specimen atoms (Sec. I.C.1). X-rays are scattered by the electrons of atoms.

A characteristic of diffraction is that each point in the diffraction pattern arises from interference of rays scattered from all irradiated portions of the object. Structure determination by diffraction methods involves measuring or calculating the **structure factor** (*F*) at many or all points of the diffraction pattern. Two quantities, an **amplitude** and a **phase describe Each F. Amplitude** is the strength of interference at a particular point, and is proportional to the square root of the intensity in the recorded pattern (photographic film does not record the scattered amplitude, but rather the intensity which is proportional to the amplitude squared). **Phase** is the relative time of arrival of the scattered radiation (wave) at a particular point (*e.g.* photographic film), and this information is lost when the diffraction pattern is recorded. Phases cannot, therefore, be measured directly from x-ray diffraction photographs. A major concern of structure determination using x-ray crystallography is the regeneration of the lost phase information. Several techniques, including the heavy atom, isomorphous replacement and molecular replacement methods were devised to solve the so-called "phase problem" (see *e.g.* Eisenberg and Crothers, 1979; Glusker and Trueblood, 1972; Holmes and Blow, 1965; Wilson 1966).

X-ray phases could be obtained if it were possible to rediffract (focus) the scattered rays with a lens to form an image. Fortunately, we can **directly** visualize structure in electron and light microscopes because electrons and visible photons scattered by specimens can be focused with lenses to form images. In the absence of "noise", an image might be considered to contain structural information (amplitudes and phases) in directly interpretable form. A major advantage of image processing is that it provides an objective means to extract reliable structural information from
noisy images.

b. The Fourier transform

The concept of the Fourier transform is essential for understanding the principles of diffraction methods. The Fourier transform, named after the nineteenth century, French mathematician, Joseph Fourier, mathematically describes the distribution of amplitude and phase in different directions, for all possible directions of the beam incident on the object.

The Fourier transform of an object is a particular kind of weighted integral of the object. In one-dimension, the Fourier transform is mathematically expressed in the following way:

\[ F(X) = \int_{-\infty}^{\infty} \rho(x)e^{(2\pi ix)} \, dx \]  

(III.C.6.1)

\( F(X) \) is called the scattering function and \( \rho(x) \) is called the electron density function. The integration is over all values \( x \) in the structure. For the case in which there is a discrete summation over sampled points in the structure the above expression becomes:

\[ F(X) = \sum_{x} \rho(x)e^{(2\pi ix)} \]  

(III.C.6.2)

\( F \) is a shorthand notation for the Fourier transform of \( \rho \), i.e. \( F = T(\rho) \), where "\( T \)" represents the forward Fourier transformation operation.

A property of Fourier transforms is that the inverse relationship holds, namely:

\[ \rho(x) = \int_{-\infty}^{\infty} F(X)e^{(-2\pi ix)} \, dX \]  

(III.C.6.3)

Thus, \( \rho \) is the inverse transform of \( F \), i.e. \( \rho = T^{-1}(F) \), where "\( T^{-1} \)" signifies an inverse Fourier transformation operation.

Thus, \( \rho = T^{-1}(T(\rho)) \)  

(III.C.6.4)

This expression emphasizes an important property of the Fourier transform, called the inversion theorem: the Fourier transform of the Fourier transform of an object is the original object. This theorem is analogous to Abbe's treatment of image formation, which is considered to be a double-diffraction process (see Sec. III.C.6.d). The recorded diffraction pattern of an object is the square of the Fourier transform of that object.

c. Fourier synthesis

Joseph Fourier also showed that any periodic function may be mathematically represented by a summation of a series of sinusoidal waves. In one-dimension, the Fourier synthesis can be expressed in the following way:

\[ \rho(x) = \sum_{n=-\infty}^{\infty} A_n \cos(2\pi nx / a) \]  

(III.C.6.5)

where \( \rho(x) \) = one-dimensional density function  
\( x \) = coordinate of a point in the object  
\( a \) = repeat distance  
\( A_n \) = Fourier coefficient (amplitude term) for wave number \( n \)  
\( n \) = wave number (frequency) or cycles per repeat distance \( a \)  
\( (2\pi nx / a) \) = phase term (position of wave with respect to a fixed origin point in the repeating structure)
Fig. III.30 depicts a one-dimensional, periodic function and the relative amplitudes and phases of the six component waves. **Wave number** refers to the number of complete cycles of each wave per repeat distance. Mathematical combination of the waves to produce the periodic function is called **Fourier synthesis**, and the opposite process, decomposition of the function into its component waves, is called **Fourier analysis** (Figs. III.30-32). The latter is formally equivalent to analyzing the sound wave harmonics of a musical instrument. The analogy between music and structure can be represented as follows:

\[
tone = \sum \text{harmonics} \\
\text{structure} = \sum \text{structure factors}
\]

![Fig. III.30. Superposition of sinusoidal waves to yield a periodic function. The wave numbers and phases of the component waves are shown. (From Eisenberg and Crothers, p. 828)](image)

![Fig. III.31. (Left, upper) Representation of the electron density of a one-dimensional "crystal" by a superposition of waves. The crystal is formed by a periodic repetition of a diatomic molecule, as shown at the top of the right hand column. The component waves, each with proper phase and amplitude, are on the left. The curves on the right show the successive superposition of the five waves on the left. (Left, lower) Representation of another 1D crystal, this one containing a triatomic molecule. Note that this crystal is built up from the same waves as the crystal of (a); only the amplitudes and phases have been changed. (Top, right) The summation of two-dimensional waves to produce a two-dimensional "electron density". (From Eisenberg and Crothers, pp. 829-830)](image)
Fig.III.32. Each diffraction spot arises from an electron density wave in the crystal. If the amplitude and phase of each reciprocal lattice point are known, the crystal structure can be synthesized by adding together the appropriate electron density waves. A pictorial example of this is shown at the right. Note that the phase angle of the third wave added has considerable effect on the type of structure produced. (From Holmes and Blow, 7, p. 131)

Diffraction methods provide a direct way to display the decomposition of a function into component waves (frequencies), and for this reason, diffraction is often called spatial frequency spectrum analysis or harmonic analysis.

d. Image formation as a double-diffraction process

Image formation, according to Abbe's theory, is a two-stage, double-diffraction process (Figs.III.33-35). That is, an image is the diffraction pattern of the diffraction pattern of an object. With an "ideal" lens system, an image precisely depicts every detail present in the object. In the first stage of image formation, a collimated (parallel) beam of rays incident on the object is scattered and the interference pattern (Fraunhofer diffraction pattern) is brought to focus at the back focal plane of the lens. This stage is sometimes referred to as the forward Fourier transformation. The intensity distribution of the recorded diffraction pattern of an object is proportional to the square of the Fourier transform of the object. Although they are not equivalent, the terms, transform and diffraction pattern, are often used interchangeably.
Fig.III.33. Image formation in a lens can be described as a double-diffraction process.

Fig.III.34. Imaging of a specimen of periodic structure. (From Slayter, p.223)

Fig.III.35. Imaging of a specimen of irregular structure. (From Slayter, p.224)
A lens, essential for image formation, also acts to focus the diffraction pattern at a finite distance from the object (at the back focal plane of the lens if the object is illuminated by a coherent beam of radiation). If the lens is removed from behind the object, no image forms, but instead Fresnel diffraction patterns form at finite distances from the object (see Fig.I.7 on p.7 of Sec.I.A.3.c) and the Fraunhofer diffraction pattern forms at infinity (or large distance relative to the object size or wavelength of radiation used). In X-ray diffraction experiments, where there is no lens to focus the X-rays, a Fraunhofer diffraction pattern is generally photographed about 50 to 150 mm behind the specimen. This distance is large relative to the size of the diffracting objects (i.e. unit cells about 10-50 nm in dimension) and the wavelength of the incident radiation (usually 0.154 nm).

The second stage of image formation occurs when the scattered radiation passes beyond the back focal plane of the lens and interferes (recombines) to form an image (Figs.III.33-35). This is the back or inverse Fourier transformation stage. Note that the image cannot exactly represent the object because some scattered rays never enter the lens and cannot be focused at the image plane. Typical specimens for diffraction experiments are predominantly transparent to the radiation, that is, most rays pass straight through the object without being scattered.

Image formation is analogous to Fourier analysis in the first stage, and Fourier synthesis in the second stage. Fourier image analysis is a powerful method for analyzing a wide variety of periodic specimens because it separates the processing of electron micrograph images into two stages. The formation of the diffraction pattern in the first stage reveals structural information in a straightforward manner and conveniently and objectively separates most of the signal and noise components in the image. The transform may then be manipulated (Sec. III.D.2) and subsequently back-transformed in the second stage to produce a noise-filtered, reconstructed image.

e. Bragg diffraction

W. L. Bragg's simple description of the diffraction from crystals helps clarify our understanding of diffraction (Fig.III.36). Diffraction can be visualized as arising from the reflection of radiation from planes of electron density in the 3D crystal (or lines in a 2D crystal). These lattice planes are imaginary parallel planes within crystals. Each set of planes is identified by three Miller indices, hkl. These indices are the reciprocals of the intercepts in units of cell edge lengths that the plane makes with the axes of the unit cell (Fig.III.37). Diffraction from the hkl set of planes that are separated a distance \(d_{hkl}\) only occurs for certain orientations of the incident radiation according to the Bragg relation: (see also p. 95, Sec. I.F.1.a).

\[
n\lambda = 2d_{hkl}\sin\theta_{hkl}
\]

(III.C.6.6)

where

- \(n\) = integer
- \(\lambda\) = wavelength of incident radiation
- \(d_{hkl}\) = crystal lattice spacing between the [hkl] set of crystal planes
- \(\theta_{hkl}\) = angle of incidence and also of reflection

Fig.III.36. Interaction of two waves (1) and (2) with the same amplitude. (a) doubling of the amplitude when the waves are in phase, (b) mutual annihilation of the waves in counter-phase, (c) change of amplitude and phase in general case of a phase shift. (From Vainshtein, p.224)
The intensity of each hkl reflection is proportional to the distribution of electron density in the hkl planes. In other words, in some planes, the density may be evenly distributed and the corresponding reflection will be relatively weak. If in others, the density is concentrated in one region between the planes, the corresponding reflection will be strong.

Two Bragg-type planes are depicted in the two-dimensional crystal of hands (Fig.III.38, left). Density that lies between the dashed lines diffract at the reciprocal lattice point labeled [1,2] (and also its Friedel mate, [-1,-2], not shown). The spacing or perpendicular distance between the lines is inversely proportional to the distance of the [1,2] reciprocal lattice point from the origin. Also, relative to the transform origin (where \( \theta_{hkl} = 0^\circ \), which corresponds to the direction of unscattered radiation), the reciprocal lattice point appears in a direction normal to the set of lines. The dotted lines in Fig.III.38 (left) represent the [2,3] set of lattice planes.

For 2D, periodic structures, each Friedel pair of spots arises from a set of fringes (sinusoidal density waves) of particular spacing (frequency) and orientation in the crystal. The so-called Miller index of each spot corresponds to the two wave numbers \((h,k)\) which describe the number of wave cycles per repeat in the a and b directions. For example, the [2,3] set of waves (dotted lines in Fig.III.38) cross the a axis twice and the b axis three times in each unit cell.

In diffraction from 3D crystals, the Miller index of each spot is assigned three wave numbers \((h,k,l)\) corresponding to the number of wave cycles per repeat in the three unit cell directions \((a,b,c)\) (Fig.III.37c).

---

Fig.III.37. Miller indices of lattice planes in a crystal. (a) A lattice plane with intercepts \(a', b', \) and \(c'\) along the \(a, b,\) and \(c\) axes. (b) Lattice planes in a 2D lattice. (c) Lattice planes in a 3D lattice. (From Eisenberg and Crothers, p.811)
Each spot or reflection in the diffraction pattern may be mathematically represented as a plane wave whose amplitude is proportional to the square root of the spot intensity and whose phase is measured relative to a particular origin point in the crystal (e.g. the unit cell origin). The phase for a particular spot is the distance from the crest of the density wave to a chosen origin (Fig.III.30). When the amplitudes and phases (structure factors, $F_{hkl}$) of all spots in the 3D transform are known, the corresponding density waves can be mathematically summed (Fourier synthesis) to reconstruct the 3D object density (Figs.III.31 and III.32).

f. Structure factor

The structure factor describes the scattering of all atoms of the unit cell for a given Bragg reflection. Each diffracted ray, or reflection, is described by a structure factor, $F_{hkl}$. It is a complex number whose magnitude is proportional to the square root of the intensity of the $hkl$ reflection. Each structure factor may be regarded as a sum of the contributions of the radiation scattered in the same direction from all atoms within the unit cell. The mathematical formulation of the structure factor depends on whether the object is considered to be continuous or made of a discrete number of atoms.

**Objects consisting of discrete atoms**

For an object with $n$ atoms, the structure factor equation is:

$$F_{hkl} = \sum_{j=1}^{n} f_j \exp\left(2\pi i (hx_j + ky_j + lz_j)\right)$$

(III.C.6.7)

$f_j$ is the atomic scattering factor for atom $j$. This is the ratio of the amplitude scattered by the atom to the amplitude scattered by a single electron. At zero scattering angle, $f_j$ = atomic number, but, as the scattering angle increases, the value of $f_j$ decreases because of the finite volume over which the electrons in the atom extend. $hkl$ refers to the particular set of diffracting planes and $x_j, y_j, z_j$ are the fractional unit cell coordinates for each of the atoms in the unit cell.

Since $e^{i\theta} = \cos\theta + i\sin\theta$, equation III.C.6.7 can be rewritten:
\[ F_{\text{hkl}} = \sum_{j=1}^{n} f_j \left[ \cos \left(2\pi \left( h x_j + k y_j + l z_j \right) \right) + i \sin \left(2\pi \left( h x_j + k y_j + l z_j \right) \right) \right] \] (III.C.6.8)

\[ F_{\text{hkl}} = \sum_{j=1}^{n} f_j \cos \left(2\pi \left( h x_j + k y_j + l z_j \right) \right) + i \sum_{j=1}^{n} f_j \sin \left(2\pi \left( h x_j + k y_j + l z_j \right) \right) \] (III.C.6.9)

\[ = A_{\text{hkl}} + iB_{\text{hkl}} \] (III.C.6.10)

We see therefore, that the structure factor \( F_{\text{hkl}} \) is a complex quantity, with a real and an imaginary part represented by \( A_{\text{hkl}} \) and \( B_{\text{hkl}} \).

**Argand diagram**

\( F_{\text{hkl}} \) is conveniently depicted as a vector in an Argand diagram in which the horizontal axis represents the real axis and the vertical axis represents the imaginary axis (Fig.III.39). In this representation, the vector quantity \( F_{\text{hkl}} \) can be thought of as composed of the vector sum of \( A_{\text{hkl}} \), the real component, and \( B_{\text{hkl}} \), the imaginary component. The vector \( F_{\text{hkl}} \) makes an angle \( \alpha_{\text{hkl}} \) with respect to the real axis. The magnitudes of the vectors \( A_{\text{hkl}} \) and \( B_{\text{hkl}} \) are, respectively, \( |F_{\text{hkl}}| \cos(\alpha_{\text{hkl}}) \) and \( |F_{\text{hkl}}| \sin(\alpha_{\text{hkl}}) \).

**Fig.III.39.** Representation of structure factors by vectors in the complex plane. (a) The structure factor magnitude \( F(\text{hkl}) \) is represented by the length of a vector in the complex plane. The phase angle \( \alpha(\text{hkl}) \) is given by the angle, measured counterclockwise, between the positive real axis and the vector \( F \). (b) Complex numbers can be added by adding their real and complex components. (c) The structure factor for a reflection may be thought of as the vector sum of the X-ray scattering contributions from many atoms. Each of the \( j \) contributions may be represented as a vector in the complex plane, with amplitude \( f_j \) and phase \( \phi_j \). (From Eisenberg and Crothers, p.822)

The **structure factor amplitude** is defined as the modulus or magnitude of \( F_{\text{hkl}} \). That is:

\[
|F_{\text{hkl}}| = \left[ (A_{\text{hkl}})^2 + (B_{\text{hkl}})^2 \right]^{1/2}
\] (III.C.6.11)

The **structure factor phase** of \( F_{\text{hkl}} \) is equal to the angle \( \alpha_{\text{hkl}} \).
Since $F_{hkl} = A_{hkl} + iB_{hkl}$ \hfill (III.C.6.12)

\[ F_{hkl} = |F_{hkl}| \cos(\alpha_{hkl}) + i|F_{hkl}| \sin(\alpha_{hkl}) \] \hfill (III.C.6.13)

\[ F_{hkl} = |F_{hkl}| \exp(i\alpha_{hkl}) \] \hfill (III.C.6.14)

If $\theta_j$ is substituted for $2\pi(hx_j + ky_j + lz_j)$, the structure factor equation can be expressed in a simpler form:

\[ F_{hkl} = \sum_{j=1}^{n} f_j e^{i\phi_j} \] \hfill (III.C.6.15)

\[ F_{hkl} = \sum_{j=1}^{n} f_j \left( \cos \phi_j + i \sin \phi_j \right) \] \hfill (III.C.6.16)

**Structure with continuous density**

For a 3D structure with continuous density, $\rho(xyz)$, the structure factor equation becomes:

\[ F_{hkl} = \iiint \rho(xyz) e^{2\pi i(hx + ky + lz)} \, dx \, dy \, dz \] \hfill (III.C.6.17)

In this expression, the integration is over the entire unit cell volume, $V$. This reemphasizes a property we already know from the definition of the Fourier transform, namely, that every point in the object contributes to every point in the diffraction pattern.

g. Convolution and multiplication (sampling)

These concepts provide a fundamental basis for understanding diffraction from crystalline objects. Holmes and Blow (1965) give a general statement of the operation of convolution of two functions:

"Set down the origin of the first function in every possible position of the second, multiply the value of the first function in each position by the value of the second at that point and take the sum of all such possible operations."

This is expressed in mathematical terms as:

\[ c(u) = \int_{-\infty}^{\infty} f(x)g(u - x) \, dx \] \hfill (III.C.6.18)

This is known as the convolution of $f(x)$ and $g(x)$, and may be written as:

\[ c(u) = f(x) \ast g(x) \] \hfill (III.C.6.19)

Several examples of convolution are schematically depicted in Figs.III.40-42.

---

**Fig.III.40.** Convolutions with an array of $\delta$ functions. If $f(x)$ is an array of $\delta$ functions, and $g(x)$ the arbitrary function shown, then the result of the convolution $f(x) \ast g(x)$ is to associate the $g$ function with each $\delta$ function. This is always true on condition that $g(x)$ is narrower than the spacing of the $\delta$ functions so that $g(x)$ never overlaps two $\delta$ functions simultaneously. (From Sherwood, p.173)
The convolution theorem provides a precise way to describe the relationship between objects in real space and transforms in reciprocal space. It states that the Fourier transform of the convolution of two functions is the product of their Fourier transforms.

$$T(f \ast g) = F \times G$$  \hspace{1cm} (III.C.6.20)

The converse of the above also holds, namely that the Fourier transform of the product of two functions is equal to the convolution of the transforms of the individual functions.

$$T(f \times g) = F \ast G$$  \hspace{1cm} (III.C.6.21)

The symbols, $\ast$ and $\times$, correspond to the convolution and multiplication operations. $f$ and $g$ represent two functions and $F$ and $G$ are the respective Fourier transforms.

The convolution of a motif ($f_1$) with a finite lattice ($f_2$) to produce a crystal structure ($f_3$) is illustrated in Fig.III.41 and Fig.III.42. These examples are easy to conceptualize because, in each case, one of the functions ($f_2$) is a lattice. Fortunately, most periodic specimens we are considering obey this restriction. A crystal structure is thus equivalent to the convolution of a finite lattice with the contents of the unit cell. Mathematical computation of the convolution of two functions is usually required when neither function is a lattice.

All of the objects in Fig.III.43 can be described as convolutions of a motif and a 1D lattice. In these examples, a hand is convoluted with one, two, three or ten-point (i.e. finite), horizontal lattices. The diffraction patterns to the right of each "object" (Fig.III.43b,d,f,h) demonstrate the properties of transforms as formulated in the convolution theorem, namely:
\[ T(f_3) = T(f_1 * f_2) = F_3 = F_1 \times F_2 \quad (\text{III.C.6.22}) \]

\[ T^{-1}(F_3) = T^{-1}(F_1 \times F_2) = f_3 = f_1 * f_2 \quad (\text{III.C.6.23}) \]

Fig.III.43. Lattice sampling. (a) A single left hand (palm facing down) and (b) its diffraction pattern. (c) Two hands and (d) their diffraction pattern. Note that this is a sampled version of transform \( b \). (e) Three hands and (f) their diffraction pattern which is a more sharply sampled version of transform \( b \). (g) Row of ten hands and (h) its diffraction pattern. This transform is the most sharply sampled version of transform \( b \).

In other words, the transform of the crystal structure, \( F_3 \), is the transform of the unit cell contents (a single hand), \( F_1 \), multiplied (sampled) by the transform of the crystal lattice, \( F_2 \) (reciprocal lattice: the sampling interval is reciprocally related to the real space lattice repeat, \( i.e. \) the distance separating the hands). The transform of the contents of the unit cell is a continuous function (Fig.III.43b), whereas the transform from the crystal is discrete (Fig.III.43d,f,h). That is, the crystal transform is the transform of the single unit cell "sampled" at the reciprocal lattice points. Values of the Fourier transform at the reciprocal lattice points are called the structure factors, \( F_{hkl} \) (Sec. III.C.6.f).

The transform of Fig.III.43c, shown in Fig.III.43d, is the transform of a single hand (Fig.III.43b) multiplied by the transform of a two-point lattice (series of vertical lines). Note in Figs.III.43d,f,h that the thickness of the vertical "sampling" lines in each of these transforms is inversely proportional to the number of hands or points in the finite lattice (an infinite lattice would produce infinitely thin lines). In Figs.III.43d,f,h the transform of a single hand (Fig.III.43b) is sampled by the vertical fringes of the reciprocal lattice function. The hand transform (Fig.III.43b) is similarly described as the transform of a single hand sampled by an infinitely thick, vertical fringe (transform of a single point).

Also notice in Fig.III.43 that the 1D lattices give rise to transforms sampled in only one direction, whereas 2D lattices produce sampling on a 2D grid or reciprocal lattice. Thus, transforms of 2D crystals are the individual object transforms sampled at the reciprocal lattice points (Fig.III.44). If the phase and amplitude (structure factor) at each point \( hk \) in the reciprocal lattice can be obtained, the crystal and motif structures can be solved by mathematical Fourier synthesis (equivalent to an inverse Fourier transformation). Figs.III.45 and III.46 show additional examples of convolution and sampling as given in the text by Sherwood.
h. Other properties of Fourier transforms and diffraction patterns

Optical diffraction / "mathematical" transform analogy

An optical bench is an excellent device for demonstrating the properties of diffraction patterns and Fourier transforms and provides a clear analogy between X-ray and Fraunhofer diffraction. Taylor and Lipson (1964) and Harburn, Taylor and Welberry (1975) present hundreds of examples of objects and their optical diffraction patterns. These examples provide a solid foundation for understanding, at least within a visual framework, the principles of diffraction theory. In optical diffraction experiments (Sec. III.D.1), the diffraction grating (object) is a photographic transparency or an opaque material in which holes have been punched or etched out, and the incident radiation is a visible laser beam. The relationship between object structure (real space) and its transform (reciprocal space) will be illustrated in several figures in the following pages.
**3D RECONSTRUCTION OF MACROMOLECULES**

**Fig.III.45.** The diffraction pattern of N wide slits. This may be represented by the square of $T$ (one wide slit) $\cdot T$ (narrow slits) $\cdot T$ (shape function). The diffraction pattern of narrow slits and the shape function are shown in (a0 and (b) respectively, and the operation of their convolution gives rise to (c), the diffraction pattern of N narrow slits. On multiplying this by the pattern of one wide slit, (e), we derive (f), the diffraction pattern of N wide slits. (From Sherwood, p.254)

**Fig.III.46.** The diffraction pattern of N wide slits showing Fourier transform and convolution relationships. (From Sherwood, p.255)

**Asymmetric vs. symmetric structures and their transforms**

Simple, symmetric structures (Fig.III.47a-d and Fig.III.48 upper left) generally produce simple, symmetric transforms (Fig.III.47e-h and Fig.III.48 upper right), whereas asymmetric structures generally produce transforms that are more complex (Figs.III.48-50). Transforms are like *fingerprints* in the sense that specific object features often give rise to characteristic features in the transform. Each structure can be regenerated by back transformation **only** if the amplitudes and phases at all points of the transform are available. This may be accomplished for visible light (optical reconstruction, Sec. III.D.2) or for electrons (electron microscopy), but can only be achieved by mathematical computation for X-rays and neutrons where phases are indirectly measured.
Some shapes (Fig.III.47a-d) may be directly deduced from characteristic transforms (Fig.III.47e-h). Simple inspection of most transforms (Figs.III.48 bottom row and Figs.III.49 and 50) though, does not directly lead to a unique determination of structure.

Reciprocity

Dimensions in the object (REAL SPACE) are inversely related to dimensions in the transform (RECPROCAL SPACE). Thus, small spacings or features in real space are represented by features that are spaced far apart in reciprocal space. Diffraction rings become smaller and narrower as the diameter of the circular object increases (Fig.III.47a,b,c,e,f,g). The close, vertical edges of the rectangular object (Fig.III.d) give rise to the widely spaced fringes in the horizontal spike of the transform (Fig.III.h), and the wide, horizontal edges of the box produce the closely-spaced fringes in the vertical spike. Reciprocity is also illustrated by examples in Figs.III.38,49 and 50.
Resolution

Outer regions of the transform arise from fine (high resolution) details in the object. Coarse object features contribute near the central (low resolution) region of the transform. Unscattered rays (those scattered through an angle of zero degrees) appear at the exact center of the transform. For example, by blocking out portions of the hand transform (Fig.III.51a) and allowing the remaining rays to recombine to form an image (equivalent to optical filtering, Sec. III.D.2), we can demonstrate that the fingers are fine details (Fig.III.51b,c,e), whereas the hand as a whole is a coarse feature (Fig.III.51f,g). As high-resolution information is removed from the transform, fine structural details (fingers) are no longer distinguished in the reconstructions. This procedure is called low-pass filtering to denote that the low-resolution features (near the center of the transform) are allowed to interfere (resynthesize) at the image plane whereas the high-resolution features are removed. Removal of, or blocking out the low resolution Fourier components leads to accentuation of the high-resolution features (edges) (Fig.III.51d).

Fig.III.51. (a) Fourier transform of a hand. Rings denote the extent to which regions of the transform are allowed to recombine to form either low-pass (b,c,e,f,g) or high-pass (d) filtered image reconstructions.

Fig.III.52. Similar to Fig.III.50, showing the effect of taking only low angle diffraction to form the image of a duck object. A drawing of a duck is shown, together with its diffraction pattern. Also shown are the images formed (as the diffraction pattern of the diffraction pattern) when stops are used to cut progressively more of the high angle diffraction pattern. (From Holmes and Blow, p.120)
**Sharpness of diffraction spots**

Features in the diffraction pattern sharpen up as the number of diffracting objects or the distance between them increases (Figs.III.43, 53 and 54). As the horizontal, 1D 'crystals' increase in size by adding more units (Figs.III.43 and III.54) or by increased separation between units (Fig.III.53), the vertical fringes sharpen in the diffraction pattern. Sharpening reflects a situation of more complete, destructive interference away from the reciprocal lattice positions. The inverse relationship between sharpness and overall object size (extent) is another example of the reciprocity between object and reciprocal space.

![Fig.III.53. Effect of object separation on transform.](image)

(a) Single hand and its transform (b). (c,e,f) Pair of hands at increasing separation and the respective Fourier transforms (d,f,h).

**Geometry, intensity and symmetry**

The relative orientation and dimensions between real and reciprocal space lattices for a 2D crystal are illustrated in Figs.III.38, 44, 55, and 56. Parameters of the crystal lattice and reciprocal lattice are related by:

\[
d^* = K/d \sin \gamma^*
\]  

and \[\gamma^* = 180 - \gamma\]  

\(^{(III.C.6.24)}\) \(^{(III.C.6.25)}\)

\(K\) is the constant of diffraction (= \(\lambda L\), where \(\lambda\) is the wavelength of monochromatic radiation and \(L\) is the camera length, i.e. the distance from the specimen to the diffraction plane).

Notice that the reciprocal lattice edges, of dimensions \(a^*\) and \(b^*\), are respectively perpendicular to the cell edges \(b\) and \(a\). Only the position but NOT the relative intensity of diffraction at the lattice points is depicted in Fig.III.38. However, the relative intensities are depicted in Fig.III.44. Each spot is indexed according to its position in the reciprocal lattice, and is considered to arise by diffraction from a set of density planes in the crystal.
Fig. III.54. The significance of the diffraction pattern. The amplitude functions for one, three, N, and $\infty$ wide slits are depicted in (a), and the appropriate diffraction patterns in (b). Introducing a motif has the effect of altering the intensities of the main peaks, but the positions of these peaks remain the same, for the lattice determines this. Also, the shape of each main peak remains the same for this is a function of the overall shape of the entire molecule. (From Sherwood, p.249)
Asymmetric and symmetric objects produce different diffraction patterns. These examples illustrate the transform of the asymmetric unit (one object) and the symmetric unit (two objects) from the original grating to the diffraction pattern. The reciprocal relationships of a and b to the spacings of certain rows in the diffraction pattern are shown. The fact that the intensity is the same at all reciprocal lattice points in the diffraction pattern in (c) should NOT be thought to be general; it happens here because the scattering objects in the original "crystal lattice" are all particularly simple (isotropic holes) and are much smaller than the wavelength of the radiation used in this hypothetical experiment. Consequently, the intensities of the diffraction maxima show no variation in different directions and do not vary significantly with angle of scattering (which increases with increasing distance from the center of the pattern). (d) The relationships of a and b to a* and b* are shown. The reciprocal lattice which is, in this 2D example, the lattice of spacing a* and b*, is of great importance in diffraction experiments. For a particular diffraction pattern, the scale factor K depends upon the wavelength of the radiation used and upon the geometry of the experimental arrangement. (From Glusker and Trueblood, pp.24-25)

Comparison of Figs.III.43 and III.53, and Fig.III.49 demonstrate that the structure of the motif and NOT the spacings or geometry of the crystal lattice determine the intensity distribution in the transform.

Structural symmetry produces symmetrical intensity distributions in the transform (aside from Friedel symmetry, see below). For example, Fig.III.50 clearly illustrates that symmetric objects produce symmetric transforms. This property is one of the major reasons why optical diffraction (Sec. Ill.D.1) is a powerful method for diagnosing the presence of symmetry in biological specimens.

Objects with rotational symmetry (Fig.III.50a,c,e,g,i,k) give rise to transforms with the same n-fold symmetry if the n is an even-order number, or 2n-fold symmetry if n is an odd number. In these examples, the transform of the asymmetric unit (one hand, Fig.III.50a) is not sampled in the symmetric transforms because the motifs (Fig.III.50c,e,g,i,k) are not crystals. That is, the asymmetric units are not in identical orientations strictly related by translations.
**Screw-axis symmetry** in a crystal produces systematic absences in the transforms. The transforms of crystals with vertical or horizontal dyad screw-axes (2₁) appear with every other spot missing in the vertical or horizontal rows (shown with slides in class).

Not shown are examples of crystalline objects with symmetric motifs. One might envision a crystal with the motif of Fig.III.50g placed at the points of a square lattice. The transform of such a crystal is the transform of the motif (Fig.III.50h) sampled at the points of a square reciprocal lattice (\(a^*=b^*, \gamma^*=90^\circ\)). In this example, the asymmetric unit of the crystal is one-quarter of the motif (one hand) because the four-fold motif symmetry matches the four-fold lattice symmetry.
Projection theorem

The projection theorem states that the Fourier transform of the projected structure of a 3D object is equivalent to a 2D central section of the 3D Fourier transform of the object. The central section intersects the origin of the 3D transform and is perpendicular to the direction of projection. Optical transforms of projected specimen images are, therefore, 2D slices through the complete 3D transform of the specimen. This forms the basis of 3D reconstruction by Fourier methods, where several independent views of the projected structure are recorded and their transforms calculated to build up a complete 3D transform (Fig.III.58). The 3D structure is reconstructed from the 2D views by inverse Fourier transformation of the complete transform.

Fig.III.58. The mathematical principles of 3D reconstruction. (a) A 3D duck and (b) its 3D Fourier transform; (c) A projection of (a) and (d) the 2D Fourier transform of (c); (e) Another projection of (a) and (f) its 2D Fourier transform; (g) The 3D duck calculated from the 3D Fourier transform (h) which was reconstructed by sampling 3D Fourier space with the 2D transforms (d) and (f). (From Lake (Lipson), p.174)
**Friedel's Law**

Friedel's law states that there is an inversion center in the intensity distribution in the diffraction pattern from the projected structure of a real object. This means that the amplitude at any point in the pattern is identical at a point equidistant and opposite in direction from the transform origin. The phases at these two points are opposite. For periodic specimens with periodic patterns consisting of discrete spots (reflections), Friedel related spots are called Friedel pairs.

In X-ray diffraction, Friedel's law breaks down under conditions where there are atoms that scatter anomalously (refer to any basic text on X-ray diffraction methods for a detailed explanation). In optical diffraction experiments, Friedel's law generally breaks down (i.e. the pattern does not show perfect inversion symmetry in the intensity distribution) because the object is a photographic transparency, which causes phase shifts of the incident radiation (laser light) as it passes through the emulsion and backing of the film. It is possible (although quite messy!) to reduce these phase effects by using specially designed optical systems in which the transparency and diffraction lens are combined in a way in which the transparency can be immersed in an oil of refractive index closely matching the refractive index of both the transparency and lens.

Mathematically computed diffraction patterns should always show perfect Friedel symmetry (e.g. Figs.III.43, 44, 47, 49-53, and 58).