Symmetry, Chemistry, and Escher’s Tiles

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Abstract

When the first cave-woman pressed an olive or nut, she got an oil that wouldn’t mix with water. Why? One reason is because the molecules’ shapes have different symmetries. In brief, an asymmetrical molecule is better able to form a dipole, which permits dipole-dipole intermolecular attractions [1].

So, chemical reality is “somehow” related to mathematical symmetry. This by itself is exciting to those interested in chemistry, but others may need more visual examples. People who may feel intimidated by or bored with mathematical theories of groups will be drawn in to the world of symmetry in architectural tiles and in art using the ideas expressed in the art of M. C. Escher.

Symmetry and Chemistry

In general, highly symmetrical molecules are hydrophobic (“water-fearing”), while hydrophilic molecules (“water-loving”) have a unique direction, or polarity. For example, water (H–O–H or dihydrogen oxide) itself has a bent shape, so the electronic trends of its two O–H bonds can add together (figure 1a). In contrast, several non-polar, hydrophobic molecules have high symmetry. For example, O=C=O (carbon dioxide) is completely linear (figure 1b). The net electronic trend is also zero in C₆H₆ (benzene), with its six equal C–H sections in a hexagon (figure 1c). These trends all cancel exactly, because of the shape and symmetry of the molecule.

Figure 1. (a) The addition of electronic forces (thin arrow vectors) from the O–H bonds in water permits macroscopic physical effects (e.g., polarity, outline resultant vector), due to low symmetry. (b) High symmetry cancels any polarity in dry ice (CO₂) and (c) in benzene (C₆H₆).
Symmetry and Point Groups

Many of the drawings of M. C. Escher demonstrate certain symmetry relationships [2]. One technique he used follows the method of tiling or tessellation, which has been used for centuries in different cultures. This is the regular filling of a surface with identical (monohedral) or related objects, leaving neither gaps nor overlaps.

While molecules are three-dimensional objects, and tiling patterns are generally on a flat surface, it is possible to illustrate some symmetry relationships with tessellations. Artistic patterns can inspire and interest those who examine chemistry or mathematics, while less complex tilings with the same symmetries can clarify some relationships of molecules. For at least half a century, people have realized that tessellations demonstrate two-dimensional analogs of the three-dimensional symmetries of solid molecules, following the principles of crystallography [3,4,5].

The present work is not focused on the repetition of molecules in a crystal, but on the inherent symmetry of individual molecules, called point-group symmetry [6,7]. This molecular symmetry is often not used by crystals. Relationships are naturally indirect between non-repeating three-dimensional point-group symmetry and repetitive two-dimensional tessellation symmetry.

The elements of point-group symmetry are rotation, reflection, and inversion of all atoms in the molecule, in three dimensions about a fixed central point, with no net change to the molecule. In contrast, the elements of tessellation symmetry are rotation, reflection, glide reflection, and translation of a side of a polygon to one other side, to create a tile that can exactly cover the surface [8]. While these tessellation symmetries thus do not correspond to molecular symmetries, they can suggest them, without requiring three dimensions.

Symmetry Groups and Tessellation

M. C. Escher wrote, in regard to regular tessellation, that “it is the richest source of inspiration that I have ever tapped, and it has by no means dried up yet.” [12]

Escher twice visited the Alhambra in Spain, and made several sketches of the tile designs found there. About this he wrote that “The Moors were masters in the filling of a surface with congruent figures and left no gaps over.”

There can be no doubt about the great significance of this art form to Escher, when he wrote that “Periodic drawings are . . . not subjective; they are objective. . . . For once one has crossed over the threshold of the early stages, this activity takes on more worth than any other form of decorative art” [12].

For two-dimensional tilings of a surface, there are 17 different plane symmetry groups, cataloged by the Hungarian mathematician G. Pólya in 1924 [9]. These can be categorized by asking questions
as to the smallest symmetric rotation, and the presence of translation, reflections, and glide-reflec­
tions, and their relationships. These 17 symmetry groups yield 81 possible isohedral tiles [10]. Here,
a tiling is isohedral if, “for every choice of two tiles in the tiling, there is an element in the symme­
try group of the tiling that sends one of the tiles onto the other” [11].

The German mathematicians Heinrich Heesch and O. Kienzele later analyzed the types of
asymmetric tiles that can form a tessellation without reflections (Table 1) [13]. This leaves seven of
the 17 plane symmetry groups (called p1, p2, p3, p4, p6, pg, and pgg). From these, there are only
28 isohedral tile types.

<p>| Tafel 10. Die 28 Grundtypen des Flächenschlusses |</p>
<table>
<thead>
<tr>
<th>Netzecken</th>
<th>5</th>
<th>5</th>
<th>5</th>
<th>6</th>
<th>6</th>
<th>4</th>
<th>4</th>
<th>4</th>
<th>4</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Netze</td>
<td>333333</td>
<td>633333</td>
<td>434333</td>
<td>443333</td>
<td>6363</td>
<td>6434</td>
<td>4444</td>
<td>666</td>
<td>884</td>
<td>12,12,3</td>
</tr>
</tbody>
</table>

![Image] Tafel 10. Die 28 Grundtypen des Flächenschlusses

Table 1. Heesch’s table of the 28 types of asymmetric isohedral tiles [13].

Each tile has 3 to 6 edges. Each edge has a symmetry relationship within itself or with another
edge. Either one half of an edge coincides with its other half by a half-turn rotation (called C), or one
edge relates to a second edge, adjoining or across from it (called T, Cn, or G). For each tile, the shape
can be thought of as a basic figure (square, triangle, quadrilateral, pentagon, or hexagon), combined
with modifications or distortions of the edges, following the rules of its symmetry group.

The translation (T) operation means that, for example, a concave shape applied to one side of a
square requires the same shape applied to the opposite side, now in a convex sense. Thus, translation
conserves the area of the original rectangular or hexagonal figure (as do the other operations). A
rotation symmetry edge, \( C_n \), is the operation that copies a shape from one edge onto an adjoining edge, related by a pivoting on the common vertex by an angle of \( 2\pi/n \). A glide-reflection, \( G_n \), is also a translation to the opposite side of the figure, but one in which the shape of the edge is reflected across the line of translation. So, a bump out on the upper left becomes a bump in on the lower right.

The Heesch symbol that categorizes a tile “is obtained by traveling a circuit around its boundary and associating to each edge the appropriate letter. Edges with a center of half-turn symmetry \([C \text{ or } C_2]\) are related to themselves; other edges are related to an adjacent or to an opposite edge by a translation, glide-reflection, or rotation” [8].

Now that these symmetry categories have been established, visual examples are needed to illustrate them. These examples are chosen based on the symmetry of selected molecules (table 2). Each of the representative molecules in Table 2 is planar, and represents either a different point-group symmetry or chemical behavior, or both.

<table>
<thead>
<tr>
<th>Molecule</th>
<th>Schoenflies Point Group</th>
<th>International Symbol</th>
<th>Heesch Analog</th>
<th>International Symbol</th>
</tr>
</thead>
<tbody>
<tr>
<td>N=N=O</td>
<td>( C_{\infty v} )</td>
<td>( \infty mm )</td>
<td>none</td>
<td></td>
</tr>
<tr>
<td>O=C=O</td>
<td>( D_{oh} )</td>
<td>( \infty/mm )</td>
<td>( C )</td>
<td></td>
</tr>
<tr>
<td>BrF(_3)</td>
<td>( C_{2v} )</td>
<td>( mm )</td>
<td>( C_3C_3C_3C_6 )</td>
<td>( p6 )</td>
</tr>
<tr>
<td>HgCl(_3)</td>
<td>( D_{3h} )</td>
<td>( 6\cdot m2 )</td>
<td>( CCC )</td>
<td>( p2 )</td>
</tr>
<tr>
<td>C(_6)H(_6)</td>
<td>( D_{6h} )</td>
<td>( 6/mmmm )</td>
<td>( C_3C_3C_3C_3C_3 )</td>
<td>( p3 )</td>
</tr>
<tr>
<td>benzene</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Z-1,2-HFC=CHF</td>
<td>( C_{2v} )</td>
<td>( mm )</td>
<td>( TT )</td>
<td></td>
</tr>
<tr>
<td>difluoroethylene</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1,1-F(_2)C=CH(_2)</td>
<td>( C_{2v} )</td>
<td>( mm )</td>
<td>( TT )</td>
<td></td>
</tr>
<tr>
<td>difluoroethylene</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E-1,2-HFC=CHF</td>
<td>( C_{2h} )</td>
<td>( 2/m )</td>
<td>( G_1G_2G_1G_2 )</td>
<td>( pgg )</td>
</tr>
<tr>
<td>difluoroethylene</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>H(_2)C=CH(_2)</td>
<td>( D_{2h} )</td>
<td>( mmmm )</td>
<td>( CCCC ) or</td>
<td>( p2 ) or ( TTTT )</td>
</tr>
<tr>
<td>ethylene</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PtCl(_4)</td>
<td>( D_{4h} )</td>
<td>( 4/mmmm )</td>
<td>( C_3C_3C_3C_3 ) or</td>
<td>( p4 ) or ( G_1G_2G_1G_2 )</td>
</tr>
</tbody>
</table>

Table 2. Molecular and tessellation symmetries for selected planar molecule examples.
The table lists the Schoenflies three-dimensional point group symbol used by chemists, and a simplified three-dimensional international symbol used by crystallographers. The table also lists the two-dimensional international symbol of the tessellation group, and the specific tile symmetry under Heesch's system, chosen by its analogy to three-dimensional symmetry, while avoiding reflection operations.

These examples can now be illustrated, both with geometrical symbols, and with tessellation artwork. In this artwork, an individual shape has one or more sides modified according to Heesch's applicable symmetry rules. The shape is given patterns or colors, then is replicated by again applying these rules of symmetry. Escher also developed and applied patterns of shading or color symmetry on many of his tessellations, which we do here for artistic purposes only. Some tessellations can be conveniently created with the aid of appropriate computer software [14].

The planar-triangular HgCl$_3^-$ anion (point-group D$_{3h}$) has three mirror planes and a rotation symmetry element of $2\pi/3$ from each vertex (Cl) to the next. It also can be represented by a CCC tessellation figure (international symbol $p2$), where half of each side of an equilateral triangle has an arbitrary shape and can be rotated to the other half, under the $C$ operation (fig. 2).

![Planar-triangular HgCl$_3^-$ anion and related CCC tessellation](image)

**Figure 2.** Planar-triangular HgCl$_3^-$ anion, and a related CCC tessellation.
Benzene, $C_6H_6$, a discrete molecule with $D_{6h}$ point-group symmetry, has a 60° rotation center and 6 mirrors (3 shown, and their bisectors) in the benzene plane (fig. 3).

Figure 3. Benzene, $C_6H_6$. Three different but equivalent representations of one benzene molecule are shown.

Here, benzene is tessellated as a flower with a $C_3C_3C_3C_3C_3C_3$ form (international symbol $p3$), in figure 4.

Figure 4. Benzene is tessellated as a $C_3C_3C_3C_3C_3C_3$ flower.
Ethylene, \( \text{H}_2\text{C} = \text{CH}_2 \), is in point-group \( \text{D}_{2h} \) (in \( mmm \)), and has two mirror planes and a \( 180^\circ \) rotation axis. It can be tessellated as \( \text{CCCC} \), for example as a duck (fig. 5).

Tetrachloroplatinum, \( \text{PtCl}_4 \), is in point-group \( \text{D}_{4h} \) (in \( 4/mmm \)), and has three mirror planes and a \( \pi/2 \) rotation axis. It can be tessellated as \( \text{C}_4\text{C}_4\text{C}_4\text{C}_4 \), as was this image, a tribute to modern art (fig. 6 and 7).
Figure 7. A $C_4$, $C_4$, $C_4$, $C_4$ figure is tessellated to show the symmetry of the PtCl$_4$ molecule.

These results serve simultaneously to illustrate tessellation symmetries and three-dimensional molecular symmetries. Some analogies are more obvious than others, and further examples are possible. A future paper will discuss these in relation to the symmetries of frieze border patterns.
References

[1] Another reason is hydrogen-bonding, which is outside of the focus of this paper. Consult any introductory chemistry text.


[2b] See also figures 8 and 9 in our other paper in these proceedings, *Circles: A Paradigm for Paradox*.


[14] All tessellations herein were created by the authors or by Jill Etheridge with the program *Tesslemania!*™, version 1.1 for Macintosh, Kevin D. Lee, Sandpiper Software Inc., © MECC, Minnesota, 1994.