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The topology of fullerenes

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Fullerenes are carbon molecules that form polyhedral cages. Their bond structures are exactly the planar cubic graphs that have only pentagon and hexagon faces. Strikingly, a number of chemical properties of a fullerene can be derived from its graph structure. A rich mathematics of cubic planar graphs and fullerene graphs has grown since they were studied by Goldberg, Coxeter, and others in the early 20th century, and many mathematical properties of fullerenes have found simple and beautiful solutions. Yet many interesting chemical and mathematical problems in the field remain open. In this paper, we present a general overview of recent topological and graph theoretical developments in fullerene research over the past two decades, describing both solved and open problems. © 2014 The Authors. WIREs Computational Molecular Science published by John Wiley & Sons, Ltd.

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INTRODUCTION

Buckminsterfullerene \( \text{C}_{60} \cdot I_h \) is a convex and 'spherical' molecule (in the sense that the atoms lie on the surface of a sphere) with a highly symmetric icosahedral structure. It was originally conjectured independently by Ōsawa in 1970\(^1\) and Stankevich, Bochvar, and Galpern in 1973\(^3,4\) discovered by mass spectrometry in 1985 by Kroto et al. through laser evaporation of graphite,\(^5\)–\(^8\) and synthesized in larger amounts by Kritschmer et al.\(^9\) in 1990 (the first mass-spectroscopic studies on small carbon cluster were performed by Mattauch et al.\(^10\) in 1943). Fullerenes are the polyhedral analogs to the two-dimensional graphene sheets.\(^6\) They occur in nature and have been detected in interstellar space, albeit in minute amounts.\(^11\)–\(^14\) In general, (classical) fullerenes are cage-like, hollow molecules of pseudo-spherical symmetry consisting of pentagons and hexagons only, resulting in a trivalent (and in the most ideal case) convex polyhedron with exactly three edges (bonds) joining every vertex occupied by carbon, idealized as sp\(^2\) hybridized atoms.\(^15\) In graph theoretical terms, fullerenes belong to the class of cubic, planar, three-connected, and simple graphs, consisting of 12 pentagons only (the 12 Pentagon Theorem) and \( F_6 \) hexagons. This gives the general formula of \( \text{C}_{20+2F_6} \) (\( F_6 \geq 0 \) and \( F_6 \neq 1 \)) for carbon fullerenes, with the number of hexagonal faces \( F_6 = 20 \) for \( \text{C}_{60} \). The smallest possible fullerene is \( \text{C}_{20} \) (\( F_6 = 0 \)), a dodecahedron consisting of 12 connected pentagons, and the only Platonic solid in the family of fullerene polyhedra.\(^16\) The truncated icosahedron \( \text{C}_{60} \cdot I_h \) is the minimal arrangement such that no two pentagons share an edge (isolated pentagons): The edges of each pentagon join only hexagons, and the edges of each hexagon alternately join pentagons and hexagons. Thus, it is the smallest member in the class of fullerenes fulfilling the so-called isolated pentagon rule (IPR), which brings thermodynamic stability to a fullerene cage.\(^18\)
There are infinitely many fullerenes, the number of isomers growing as $O(N^6)$ for $N$ carbon atoms as shown in Figure 1.19 The reason for the $N^6$ growth comes from algebraic geometry, and is discussed below. These isomers come in many different shapes depending on the distribution of the pentagons as shown in Figure 2. As experimental techniques advance to synthesize new fullerene isomers with many interesting chemical and physical applications, see for example the recent bottom-up synthesis of $C_{60}$ by Scott et al. and Kabdulov et al., one naturally wants to know more about their thermodynamic stability and electronic properties. As the isomer space rapidly grows into the millions and beyond, this becomes a daunting task for larger fullerenes. Even when the search is restricted to fullerenes that have isolated pentagons, known to yield the best stability, the number of candidates for most thermodynamically stable isomers is huge, many orders of magnitude too large for quantum chemical methods to be a viable means of analyzing stability. But, remarkably, the fullerene graphs contain all the information we need, and we are able to sort through the millions of isomers, finding a few candidates for the most stable, by way of simple, easily computed topological indicators, such as the neighbor indices for pentagons or hexagons, or the incremental assignment of heat of formations to certain face patches.

There has been an intense activity in the field of topological and graph theoretical descriptions of molecules such as fullerenes over the past 20 years, to the extent that it has become a major sub-discipline within mathematical chemistry. It comes perhaps as a surprise that graph theory applied to molecular structures has not yet made it into main-stream university teaching. Furthermore, since the publication of the standard reference work in this area, An Atlas of Fullerenes by Fowler and Manolopoulos, there has not been a comprehensive review on this subject highlighting the many activities in recent times. In contrast, there are excellent reviews and books available on the chemistry and physics of fullerenes or on their electronic structure calculations. In this overview we aim to close this gap and report on the many new and exciting developments in the topological and graph theoretical treatment of fullerenes, which have taken place over the past decade or so.

### TOPOLOGICAL PROPERTIES OF FULLERENES

**A Short Introduction to Graph Theory and Embeddings**

A graph is a pair $G = (V, E)$, where $V$ defines a set of vertices, and $E$ is a set of edges, each an unordered pair connecting two vertices. If a graph can be drawn in the two-dimensional plane without edge crossings, then it is called planar, and in this case it can also be drawn using only straight lines. A planar embedding is a map $V \rightarrow \mathbb{R}^2$ assigning a 2D-coordinate to each vertex such that straight-line drawings of the edges do not cross. For planar graphs in general, planar embeddings are not unique. But whenever at least three vertices must be removed before the graph separates into two disconnected components, the graph is called three-connected, and there is essentially only one way to embed it in the plane. Because of this, a three-connected planar graph has a well-defined set of faces, $F$. We can therefore represent a three-connected planar graph as $G = (V, E, F)$ without ambiguity, and will do so where convenient.

Whenever a graph is planar, it can also be embedded onto a surface of a sphere without edge crossings, and vice versa. Once again, this embedding is unique (in the same sense as for a planar embedding) if and only if the graph is three-connected, and we obtain the same faces. In fact, the graphs describing three-dimensional polyhedra are exactly the three-connected planar graphs, and are therefore called polyhedral graphs.

The number of neighbors to a vertex $v$, that is, the number of edges incident to $v$, is called its degree, abbreviated as $\text{deg}(v)$. If every vertex of a graph $G$ has the same degree $k$, then $G$ is called $k$-valent or $k$-regular. The special case of three-valent graphs is called cubic, and has a host of attractive mathematical

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**FIGURE 1** Number of distinct (non-isomorphic) fullerene isomers $C_60$ (with and without fulfilling the IPR) with increasing number of carbon atoms $N$ up to $N = 400$ (double logarithmic scale). (Data taken from the House of Graphs Ref 34)
properties, not least of which is their deep connections to algebraic geometry.19

Fullerenes have the neat property that the graphs formed by their bond structure are both cubic, planar, and three-connected, for which all faces are either pentagons or hexagons. Because of this, the mathematics describing them is in many cases both rich, simple, and elegant. We are able to derive many properties about their topologies, spatial shapes, surface, as well as indicators of their chemical behaviors, directly from their graphs.

Planar connected graphs fulfil Euler's polyhedron formula,

$$N - E + F = 2$$

with $N = |V|$ being the number of vertices (called the order of the graph), $E = |E|$ the number of edges, and $F = |F|$ the number of faces (for fullerenes these are...
hexagons and pentagons). The hand-shaking lemma in graph theory tells us that

$$\sum_{i=1}^{N} \text{deg}(v_i) = 2E$$

Since for fullerenes \(\text{deg}(v_i) = 3\) for all vertices, we obtain

$$E = \frac{3}{2} N, \quad F = \frac{1}{2} N + 2 = \frac{1}{3} E + 2, \quad \text{and} \quad E = 3F - 6$$

Hence, for fullerenes and fulleroids one has an even number of vertices. The total number of faces is \(F = F_5 + F_6\), where \(F_5\) and \(F_6\) are the number of pentagons and hexagons respectively. From this we derive from Eq (3) \(E = 3F_5 + 3F_6 - 6\) and \(N = 2F_5 + 2F_6 - 4\). On the other hand, each pentagon (hexagon) has five (six) edges, which gives \(E = (5/2)F_5 + 3F_6\). From these equations together with Eq (1) we obtain \(F_5 = 12\) for regular fullerenes (the ‘12 Pentagon Theorem’ for fullerenes), which gives the number of hexagons as \(F_6 = (N - 20)/2\) with \(N \geq 20\), and the general formula \(C_{20+2F_5}\) for fullerenes. Hence, from the fact that we only allow for cubic planar graphs consisting of pentagons and hexagons, we restrict the number of pentagons to exactly 12 (no more and no less!), and every addition of a hexagon adds two new vertices. Moreover, \(N = 22\) with just 1 hexagon and 12 pentagons is not valid, but every number of hexagons two or greater yields at least one fullerene.\(^{51}\) The most stable \(C_{22}\) molecule is, however, a cage-like fulleroid with a four-membered ring.\(^{52}\)

In this article, we will often work with the dual graph. The dual graph \(G^*\) of a planar connected graph \(G\) has a vertex corresponding to each face of \(G\), and an edge joining two neighboring faces in \(G\). By this process, the old vertices become the new faces. The dual of a planar graph is itself a planar graph, and for fullerenes it is a triangulation with 12 vertices of degree 5 and the remaining of degree 6. For planar graphs in general, the dual is not unique, but as discussed above, it is unique for three-connected graphs such as fullerenes, and hence the dual operation is well defined.\(^{48}\) The dual operation is an involution, that is, \((G^*)^* = G\), so we can think of the dual as just another representation of the same graph. Figure 3 shows the duals of two representative fullerenes with their corresponding 2D graph representation discussed in the next section.

A useful representation of a graph is the adjacency matrix \(A_{ij}\) well known to chemists, as it is widely used in Hückel theory of conjugated organic
molecules. The adjacency matrix is a symmetric matrix defined by \( A_{ij} = 1 \) if the graph \( G \) contains an edge \( v_i - v_j \), and \( A_{ij} = 0 \) otherwise. It is equivalent to the edge-set representation, and is much easier to work with in many settings. As the number of edges for fullerenes is \( E = 3N/2 \), the adjacency matrix is very sparse. If two graphs \( G_1 \) and \( G_2 \) are isomorphic, the corresponding adjacency matrices \( A_1 \) and \( A_2 \) are related by vertex permutation, and thus share the same set of eigenvalues and eigenvectors. Matrices play a special role in chemical graph theory, and the reader is referred to Ref 53 for more details.

### Drawing Fullerene Graphs: Methods for Planar Embedding

Planar embeddings can be drawn on paper and give us a way to visualize the structures of planar graphs. However, not all choices of embeddings are equally informative. We want the drawings to be simple, not too cluttered, and to expose as much as we can of the graph’s structure and symmetry. In general the full symmetry group can not be realized in two dimensions, but we can find embeddings that show as much of it as possible. Figure 4 shows planar drawings of three icosahedral fullerenes, \( C_{20} \cdot I_b \), \( C_{60} \cdot I_h \), and \( C_{540} \cdot I_h \), exposing 10-fold \( D_{5} \)-symmetry out of the full \( I_h \) group. In this section, we will discuss methods for constructing good planar embeddings.

While a small graph can easily be drawn by hand, larger graphs such as \( C_{540} \) shown in Figure 4(c) require a computer algorithm and a drawing program. The first method we will discuss is called the Schlegel projection, and requires first an embedding of the graph as a three-dimensional polyhedron. In 1886, Victor Schlegel showed that a planar graph can be obtained from a three-dimensional polyhedron by projection in the following way: A point \( x_0 \) is placed slightly above one of the faces, and a plane \( P \) is chosen to lie below the polyhedron. Lines are then drawn from \( x_0 \) through each of the polyhedron’s vertices. The intersection of each line \( x_0 - x_i \) with \( P \) defines the 2D coordinate of the vertex \( v_i \). If \( x_0 \) is chosen sufficiently close to the face, and if the points of the polyhedron lie on a sphere, the resulting 2D embedding will be planar. While Schlegel only proved that the projection produces a planar layout when the vertices lie on a sphere, it still works in many cases where this is not the case, and the method is widely used to the extent that planar graph embeddings sometimes (incorrectly) are called Schlegel diagrams. However, Schlegel diagrams are specifically the embeddings that result from the Schlegel projection. The Schlegel projection is illustrated in Figure 5(a).

A problem with the Schlegel projection is that it often fails, and produces crossing edges, when used with many non-spherical polyhedra or when the parameters are not well chosen. The cone-projection method, shown in Figure 5(b), which projects the vertices out on an enveloping cone of the fullerene polyhedron and then down onto a plane, seems to be more robust.\(^{35}\)

Both the cone-projection and the Schlegel projection require first a polyhedral embedding of the graph, and on top of that often necessitate bit of hand-tuning to produce a planar embedding. The third method we mention is called the Tutte embedding,\(^{54}\) and is guaranteed to always yield a planar embedding for three-connected planar graphs. In the Tutte embedding, one fixes the coordinates of a single, outer face, and solves a sparse linear system

\[
\sum_j T_{ij} x_j = \overline{y}_i
\]

such that in the result, every vertex is placed in the barycenter of its neighbors. For fullerenes, \( T_{ij} = \delta_{ij} - A_{ij}/3 \) and \( \overline{y}_i = 0 \) for every \( i \) where \( v_i \) is not a fixed vertex, and for the vertices \( v_i \) on the outer face, \( T_{ii} = \delta_{ii} \) and \( \overline{y}_i \) is the fixed coordinate. While the Tutte embedding is guaranteed to be planar, the solution to the linear system results in exponential crowding of the vertices when embedding large graphs, making the result difficult to interpret. Hence, it is useful to follow the Tutte embedding by an optimization to make the face sizes more even.\(^{35}\)
FIGURE 6 | (a) Canonical face spiral for $C_{60}-I_h$. (b) The first fullerene ($C_{28}-D_2$) with a failing face spiral.

We note that there are many different graph drawing algorithms available, each with advantages and disadvantages, but none that are guaranteed to always yield good results.\textsuperscript{54–57}

The Large Isomer Space

How does the number of $C_N$-isomers grow as we increase the number of vertices $N$? Naively one might assume a hypothetical sphere of ‘connected’ hexagons (which is of course impossible because of Euler’s theorem) and substitute 12 hexagons by pentagons to obtain a set containing all possible fullerene isomers fulfilling Euler’s theorem (just like playing Lotto), we obtain an upper bound for the number of isomers of the order $\mathcal{O}(N^{12})$. This is, however, three orders of magnitude too large, but the proof is nontrivial.

In 1998, William Thurston showed that all the triangulations of the sphere with vertex degree at most 6 form a lattice of integer points in a complex hyperbolic space $\mathbb{C}(1,9)$ through a parameterization as described in Ref 19. The triangulations with $N$ vertices or less correspond to the lattice points inside the intersection between a certain cone and a ball of radius $\sqrt{N}$ in this space. Hence their number is roughly proportional to $N^{10}$, and the number of triangulations with exactly $N$ vertices is roughly proportional to

$$\Delta(N^{10}) = N^{10} - (N - 1)^{10} = 10N^9 - 45N^8 + 120N^7 + \cdots$$

or, in short, $\mathcal{O}(N^9)$. Because they are the largest class of the finitely many triangulation classes with vertex degree six or less, the fullerenes have the same asymptotic count. Hence, the number of $C_N$ isomers grows as $\mathcal{O}(N^9)$. We note in passing that the connection of fullerenes to combinatorial manifolds is a deep one and gives rise to many of the beautiful mathematical properties exhibited by fullerenes.

The high order polynomial growth of fullerene isomer counts makes it difficult to establish a complete database for fullerene graphs up to high vertex numbers.\textsuperscript{34} Moreover, the proportion of IPR isomers to the total number of isomers grows with increasing $N$, gradually approaching 1 with increasing $N$, as is shown on Figure 4.

Face Spiral Representations of Fullerene Graphs

One of the first methods for encoding fullerene graphs was the face spiral algorithm by Manolopoulos et al.,\textsuperscript{58} which unwinds all faces of a fullerene ‘like an orange peel’ (the Manolopoulos face spiral algorithm). More specifically, starting with a sequence of three mutually adjacent faces, new faces are added to the string such that the next face is adjacent to the previous one and the one that was added to the string earliest, and that has neighboring faces left which are not part of the spiral string (yet). As the result, one obtains a string of length $F$ of 12 fives and $F - 12$ sixes, corresponding to the sizes of faces.

There are $6N$ possible spiral starts resulting in up to $6N$ spirals per fullerene graph. In most fullerenes, some spirals cannot be completed because the two rules for choosing the next face cannot be both fulfilled. Since a fullerene spiral always consists of 12 fives and $N/2 - 10$ sixes, this spiral string can be abbreviated as a list of the 12 pentagon positions, called the face spiral pentagon indices (FSPI) $\{S_n|n = 1, \ldots, 12\}$. The lexicographically smallest of all successful spirals is the canonical spiral representation of the graph.
For example, for $C_{60}$ we have the canonical FSPI of \{1, 7, 9, 11, 13, 15, 18, 20, 22, 24, 26, 32\} as shown in Figure 6.\(^{31}\)

While it was initially conjectured that every fullerene can be unwound into a face spiral\(^{58}\) (for good reason: the first counterexample comes after over $10^{12}$ isomers) there is an exceedingly small proportion of fullerene graphs\(^{59}\) (but infinitely many in total) that do not admit a face spiral,\(^{60}\) that is, where all $6N$ spiral starts fail. See Figure 6 for a case where a face spiral fails. The smallest of these unspiralable fullerenes is $C_{380-T}$\(^{61}\). Manolopoulos et al.,\(^{60}\) Fowler et al.,\(^{62}\) Yoshida et al.,\(^{63}\) and Brinkmann et al.\(^{64}\) compiled lists of non-spiral fullerenes with up to 1000 atoms and selected point groups (all $T_+$, all isolated pentagon triple (IPT) fullerenes with $D_2$ and $T$ symmetry, all $D_{3+}$, and all truncated IPT fullerenes respectively). Two of those are shown in Figure 7.

In order to encode any fullerene graph, it is necessary to extend the face spiral algorithm to a general version in which the spiral may ‘jump’ from one face to a non-adjacent face. Both Brinkmann\(^{65}\) and Fowler et al.\(^{66}\) published general face spiral algorithms for cubic polyhedral graphs, including fullerene graphs. Fowler’s algorithm extends the two above mentioned rules such that the subgraph that has not been added to the spiral yet never gets disconnected, and as a result the spiral never gets stuck. Brinkmann’s algorithm, in contrast, adds faces as defined by the non-general version, but defines how to proceed if all neighbors of the current face have been added to the spiral already. Both algorithms yield the same spiral string as the non-general algorithm if no jump is required.

For the non-general as well as both general algorithms for finding the canonical face spiral of a graph, there is an inverse algorithm that efficiently regenerates the graph from the (general) spiral string representation. With the extension to general spirals, all fullerene isomers with any vertex number can be generated. In addition, all the information of the fullerene graph is contained in a general face spiral consisting of 12+ numbers (12 for most fullerenes). This makes it rather convenient for database storage as the face spiral uniquely defines the fullerene.

In a similar way to face-spirals we can introduce vertex spirals (or Hamilton spirals) useful for identifying locations of substituents (such as hydrogen) or reaction centers\(^{67}\) (for details see Ref 68). For fullerenes with no vertex spirals the scheme can be extended to general spirals with jumps similar to general face spirals.

**Generation of Fullerene Graphs**

In order to explore chemical, physical, or graph theoretical properties for a wide range of fullerenes, it is important to have access to a list of (stored) fullerene graphs. For this, one requires an exhaustive and efficient generator for all fullerene isomers of a given vertex number $N$. The general face spiral algorithm is well suited for compactly storing and recreating specific fullerene graphs. It also allows for sorting all isomers of $C_N$ for a given vertex number $N$. The general face spiral algorithm is well suited for compactly storing and recreating specific fullerene graphs. It also allows for sorting all isomers of $C_N$ for a given vertex number $N$ according to their FSPIs including information about jumps if required. It is, however, extremely inefficient to generate all isomers $C_N$ directly through a face spiral algorithm. In the non-general case one attempts to create graphs from $\binom{F}{12}$ spiral strings while the number of non-isomorphic graphs only grows $\sim N^9$.

A large and quickly growing portion of all potential spiral strings are either non-canonical spirals, duplicates or are not successful for coding a fullerene. Including general spirals increases the overhead by two orders of magnitude, as jumps could be inserted at any step and with arbitrary lengths $l < N$.

A different approach to generating fullerene graphs is by adding faces to an existing graph, while...
considering different sites for addition at each step. Liu et al.\textsuperscript{69} introduced a method which grows a graph starting with a single face and thus only reaches a fullerene graph in the last step. Brinkmann et al.\textsuperscript{70} use patches, rather than single faces, as building blocks. These patches can be obtained by subdividing existing fullerene graphs.

Further development in efficient graph generation came from transforming an existing fullerene graph into a new, larger one by adding faces. Using \( C_{24} \) as a ‘seed’, Brinkmann et al.\textsuperscript{71} were able to generate almost all fullerene graphs up to \( C_{200} \). However, three graphs were not accessible by the applied set of transformations. To cure this shortcoming, Hasheminezhad et al.\textsuperscript{72} defined a set of patch replacements (or growth operations) which, starting from either \( C_{20} \) or \( C_{28} - T_{d} \), yield every fullerene in a systematic way. The set consists of one single operation that elongates a nanotube with minimal caps, one class of linear generalized Stone-Wales transformations\textsuperscript{73,74} that depend on one parameter, and one class of bent generalized Stone-Wales transformations that depend on two parameters. Brinkmann et al.\textsuperscript{75} used this set of growth operations to define a fast and complete algorithm that recursively generates all fullerene isomers up to a given maximal \( N \). The algorithm prunes the recursion tree in such a way that the only one representative of each isomorphism class is ever considered. Without such a scheme, exhaustively generating all fullerene graphs would succumb to combinatorial explosion.\textsuperscript{76} The result is an incredibly efficient algorithm which has been used to generate an exhaustive database up to \( C_{400} \), which is available at the House-of-Graphs website.\textsuperscript{34} Sometimes one is not interested in generating exhaustive isomer lists (which are huge for large \( N \)), but rather to generate in a directed fashion fullerenes that have specific properties. For this, one may start with one specific isomer \( C_{N} \) and derive others by certain transformations. This will be discussed in the following section.

Transformation of Fullerene Graphs

Transformations of fullerene graphs can be divided into local transformations that leave all but a certain region of the graph unchanged, and global transformations such as the Goldberg-Coxeter transformation.\textsuperscript{77,78} A planar graph can be transformed locally by replacing one fullerene patch\textsuperscript{74,79–84} by either a different patch or the same patch in a different orientation. A patch is a set of faces that is bounded by a simple cycle,\textsuperscript{82,85} that is, a cycle that traverses no vertex or edge twice. Some authors relax this condition and allow cycles that are non-selfintersecting, but are permitted to traverse any edge at most twice. This definition includes patches that have bridges, and is used in the complete characterization of fullerenes by Hasheminezhad et al.\textsuperscript{72}

Two patches can be exchanged if they share the same boundary code. The boundary code of a patch is the sequence of free valencies of the vertices that lie on the boundary (in case of cubic graphs, a sequence of zeros and ones). A few patch replacements are shown in Figure 8.

Replacing patches of equal size can be understood as a formal isomerization, while replacement by larger or smaller patches, referred to as vertex insertion or deletion, or growth operations, formally derives a molecular graph of a different size. Two fullerene graph patches with the same boundary code contain the same number of pentagons, as can be seen from Euler’s polyhedron formula. The most prominent example for formal isomerizations is the Stone-Wales (or pyracylene) transformation (SW),\textsuperscript{86} which can be generalized\textsuperscript{73,87} by allowing a variable distance between the two pentagons. Astakhova et al.\textsuperscript{88} extended this rearrangement mechanism even further to higher numbers of pentagons. Brinkmann et al.\textsuperscript{84} generated a catalog of isomerization pairs with up to five pentagons.

Isomerization operations can be reformulated as sequences of vertex-pair switches, called 2-switches.\textsuperscript{89} According to Berge’s switching theorem,\textsuperscript{90} the whole isomer space of a given \( C_{N} \) is accessible by repeated application of 2-switches. However, intermediate structures of this procedure might have faces different from pentagons and hexagons, or result even in non-planar graphs.
Growth operations can be classified according to the number of pentagons and the number of vertices that are added. There are no growth operations for fullerene graphs that involve no or only one pentagon. The smallest example is the Endo-Kroto transformation (EK), which involves two pentagons and adds two vertices. Additional noteworthy examples are the addition of four and six vertices at a patch that contains three pentagons and has $C_3$ symmetry. Brinkmann et al. compiled an extensive list of growth pairs. The three classes of patch replacements defined by Hasheminezhad et al. form the basis of the currently fastest fullerene graph generator, as discussed in a previous section.

With respect to the polyhedral representation of a fullerene graph, growth operations can be understood as the capping of a domain by additional vertices. As a result the pentagons in that domain move toward each other. Patches in which all pentagons are fused cannot be capped. Conversely, the inverse of a growth operation corresponds to the truncation of a domain of high curvature: The distances between the affected pentagons increase.

While patch replacements are useful from a graph theoretical point of view to obtain new fullerene isomers, the EK $C_2$ insertion and the SW transformation have also been suggested to resemble viable reaction pathways. As Stone and Wales pointed out in their initial article, a concerted mechanism for the SW transformation ([2+2]) is Woodward-Hoffmann forbidden in the electronic ground state. Other mechanisms that involve the breaking of one bond or the catalysis by additional carbon atoms are therefore more likely. For a discussion of different mechanisms see Ref. 93. For every suggested pathway, however, the activation barrier is so high that SW transformations are only feasible at very high temperatures. The SW transformation is believed to be one of the main mechanisms by which fullerene cages equilibrate during formation to form the most stable isomer. Starting at $C_{60} - I_h$, 1709 out of the 1811 other isomers are accessible by consecutive SW transformations.

Endo and Kroto proposed a concerted mechanism for what has since been known as the EK $C_2$ addition. However, more recent experimental and theoretical investigations show, that the concerted reaction leads to an unstable adduct and is followed by $C_2$ ejection. For a summary of alternative mechanisms of the $C_2$ addition to fullerene cages see Ref. 95.

Given any fullerene, it is possible to construct an infinite series of larger ones with (essentially) the same three-dimensional shape. This is possible through the Goldberg-Coxeter transformation, named after an infinite series of icosahedral-symmetry polyhedra described by Goldberg in 1937. By superimposing a hexagonal mesh on the surface of the dodecahedron, as shown in Figure 9(a), a new polyhedron is obtained with the same number of pentagons and an increased number of hexagons. The number of vertices in the new polyhedron is $k^2 + kl + P$, where $k$ and $l$ are integers describing the scale and orientation of the mesh, as shown for the dual in Figure 9(b).

While originally only defined for $C_{20}$, yielding exactly all the fullerenes of icosahedral symmetry, Dutour and Deza have shown that it is well defined for all cubic planar graphs. However, it is not

**FIGURE 9** Illustration of GC-transform. (a) Original figure of a hexagonal sheet from Goldberg’s paper. (b) The GC-transform acting on a face in the fullerene dual for various values of $k$ and $l$. 
trivial to determine how to automatically perform the general transform, the main difficulty lying in how to ‘glue’ the graph back together when the new edges cross the transformed triangles in complex ways. The general construction by Dutour and Deza is quite algebraically heavy handed, and is not easily understood, nor lends itself easily to implementation. In the case of fullerenes, however, it is quite easy to give a procedural description of the transformation in terms of the fullerene dual: Because a fullerene dual is a triangulation of the sphere with no vertices of degree more than 6, it can be unfolded onto the plane of equilateral triangles, called the Eisenstein plane. The unfolded surface forms a polygon in the Eisenstein plane with every degree 5 vertex on the polygon periphery, and with each edge on the periphery appearing twice, once in each direction.

The vertices in the Eisenstein triangulation is a sub-ring of the complex plane, and we can write

\[(a, b) = a + bω \text{ where } ω = e^{i\frac{2\pi}{3}}\]  \hspace{1cm} (6)

Because it is a sub-ring, multiplication by Eisenstein integers \((a, b)\) are exactly the operations that bring \((1,0)\) into every other vertex point. Because \(ω^2 = ω - 1\), we get the multiplication rule

\[\begin{align*}
(a, b) \cdot (c, d) &= ac + (ad + bc) ω + bdω^2 \\
&= (ac - bd, bc + (a + b) d)
\end{align*}\]  \hspace{1cm} (7)

and the magnitude of \((a, b)\) is \((a + bω)(a + bω^{-1}) = a^2 + ab + b^2\). Hence, if \(T\) is an equilateral triangle with unit area, the area of \((k, l)T\) is \(k^2 + kl + l^2\). This is no coincidence: multiplication in the Eisenstein ring exactly corresponds to the Goldberg-Coxeter transformation in the dual formulation. This yields a simple and practical formulation of the Goldberg-Coxeter transform of any triangulation of the sphere with degree \(\text{deg}(v) \leq 6:\)

\[\text{GC}_{k,l}(G^*) = \text{fold} \left( (k + lω) \cdot \text{unfold} (G^*) \right)\]  \hspace{1cm} (8)

The procedure is illustrated in Figure 10. Details for how to efficiently perform the fold and unfold operations are given in Avery (unpublished manuscript). Note that two successive GC-transformations on a graph \(G^*\) can be expressed as a single GC-transformation by multiplying the two Eisenstein numbers together. Similarly, a GC-transformation can be inverted by dividing instead of multiplying. For any particular fullerene, we can even use Euclid’s algorithm to find out whether it is a GC-transform of a smaller fullerene, and to find the smallest such ‘parent’ fullerene.\(^{59}\)

Notice, that if one were to cut out the two diagrams in Figure 10, gluing together the edges so that the numbers on the vertices match, one obtains the three dimensional structure of the given \(C_{12}\) and \(C_{224}\) fullerenes. The reader is invited to do so.

There are two special cases of the Goldberg-Coxeter transformation that warrant closer study: the case \(l = 0\), named the halma transform (or \(k\)-inflation) for the construction’s similarity to the game board of halma, or ‘Chinese Checkers’; and the case \(k = l = 1\), called the leapfrog transform. Both of these types of Goldberg-Coxeter transformations are simple to understand: every triangle is subdivided and reconnected the same way, as illustrated on Figure 11. While the GC-transform in general can introduce, reverse, or remove chirality (but otherwise preserves symmetry), both the halma and leapfrog transformation always preserve symmetry.

### Geometry of Fullerenes

Many of the beautiful properties of fullerenes derive from their relation to algebraic and differential geometry. This relation is mostly out of scope for this review, but in this section we will touch on the subject informally. The subject is treated in great depth by Thurston\(^{19}\) and others.

An important quantity for understanding the geometry and shapes of fullerenes is the Gaussian curvature. The Gaussian curvature is the product of the two principal curvatures, which for each point on the surface are the maximal and minimal curvatures in any direction through that point. By the Bertrand-Diquet-Puiseux theorem, the Gaussian curvature in a point is the same as the difference between \(2\pi\) and the angle required to make a circle in a neighborhood around this point. Figure 12 illustrates three categories of surfaces with zero, positive, and negative Gaussian curvature respectively. If the Gaussian curvature is zero in a point, the surface only bends in one direction around that point. A surface that has zero Gaussian curvature everywhere is characterized by being ‘flat’ in the sense that it can be unwrapped onto a plane without tearing. Positive Gaussian curvature bends the same way in all directions. A positive curvature surface can be cut open and unwrapped onto a plane. Finally, negative curvature around a point constitutes a saddle point, yielding a wobbly pringle-like surface. Negative curvature surfaces cannot be unwrapped onto a plane, because there is no room: around a negative curvature point, the angle of a circle is greater than \(2\pi\). If one were to make a cut in a negative curvature surface, it could only be flattened out by allowing parts of it to overlap itself.
The Gaussian curvature only depends on the topology of the surface, and is independent of how it is isometrically embedded in space. In the same way, the surface metric (or Riemann metric), which determines the geodesics and distances between points along the surface, can be derived directly from the graph (Avery et al., unpublished manuscript), and is also independent of the spatial embedding. However, while we do not require a 3D embedding to analyze these surface properties, they themselves do determine the possible ways that the surface can embed isometrically into space, and what is its 'natural' three-dimensional shape.

Fullerenes are all closed surfaces with non-negative Gaussian curvature everywhere. Surprisingly, this is simply due to them having faces no larger than hexagons! In fact, the slightly larger class of three-connected cubic planar graphs with largest face size 6, and their dual triangulations, are in a sense all the positive curvature genus 0 surfaces.

It is most easy to understand the geometry of fullerenes when considering their duals. These are equilateral triangulations of a closed surface. The equilateral triangle plane, also called the Eisenstein plane, is the dual of a hexagonal mesh. Hexagon planes (like graphene) and their dual degree-six triangulations are flat in the sense that they have Gaussian curvature 0 everywhere: any surface of equilateral triangles with vertex degree only six can be unwrapped into the plane without stretching or tearing. What happens now if we set the degree to 5 of a single vertex in the plane? To do this, we must cut out an infinite triangular wedge of angle $2\pi/6$. Gluing together the edges forms an infinite cone with Gaussian curvature zero everywhere except the degree 5 vertex at the cone’s apex, which introduces the positive curvature $2\pi/6$. 

**FIGURE 10** | The steps in the GC$_{2,1}$ Goldberg-Coxeter transformation from C$_{32}$-D$_{3h}(5)$ to C$_{224}$-D$_{3h}$. The transform and the diagrams of the unfolded fullerenes were automatically generated using the program Fullerene$^{35}$. 

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FIGURE 11 | Subdivision of a triangle in the dual graph; (a) for the leapfrog transformation ($k = l = 1$), and (b) for the lowest few halmas ($l = 0$).

Every time we introduce a vertex of degree 5 (or a pentagon in the hexagonal plane), it introduces a surface cut that is glued together to locally introduce the positive Gaussian curvature $2\pi/6$ at the vertex. After 12 pentagons we reach $4\pi$, the Gaussian curvature of the sphere, closing the surface. Unfolding again along the 12 cuts would result in a polygon similar to the diagrams shown in Figure 10. Had we instead introduced, for example, degree 4 vertices, each would contribute with curvature $2\pi/3$, requiring only 6 to close the surface. The positions at which we placed the 12 pentagons determine where and how strongly the surface bends, and through that the natural shape of the fullerene, giving rise to the many interesting polyhedral shapes shown in Figure 2.

Because fullerenes always have exactly 12 curvature-$2\pi/6$ bends, with the remaining surface of Gaussian curvature zero, they adopt polyhedral shapes rather than being spherical in nature (the latter defined as atoms lying on a sphere), and the most symmetric shapes that they can achieve have icosahedral symmetry. The non-spherical nature of fullerenes was pointed out already by Bakowies and Thiel in 1995.104,105

The curvature of cubic graph surfaces also provides an intuitive understanding of why fullerenes, having only hexagon and pentagon faces, would be more prevalent in nature than other polyhedral carbon structures: smaller face sizes introduce much sharper bends, reducing stability. Heptagons by themselves introduce no sharper bends than pentagons, but each heptagon necessitates an extra pentagon to balance out the negative Gaussian curvature and reach a closed surface, increasing the total number of bends by two. However, with the right distribution of heptagons and pentagons, it is possible to construct quite smoothly curving near-spherical fulleroid surfaces that could potentially be stable in nature. We discuss fulleroids, a generalization of fullerenes that allow arbitrary face sizes, in the final section.

Generating Accurate 3D Geometries

The Cartesian coordinates for the ideal molecular geometry of small highly symmetric fullerenes such as $C_{20} - I_h$ or $C_{60} - I_h$ can be calculated directly using basic geometry.106 This is not the case for fullerenes in general: They come in many different shapes, most of which do not have a simple mathematical formula. However, we do know that the ideal geometry (in which the faces of the dual polyhedron are equilateral triangles) can be derived directly from the fullerene graph. This is because of their special properties discussed above. However, while we know that a fullerene graph determines the ideal 3D structure as a conformal, isometric embedding of the surface into space, it is not yet known how to compute it directly. However, we can use numerical optimization methods to obtain good 3D structures—both efficiently and with results that are very close to the physical molecular geometry.

Generating Initial Structures

Before optimizing a fullerene structure by for example a force-field method, we need a reasonable initial structure. Specifically, we need an embedding of the
graph as a polyhedron (i.e., no crossing edges) such that its faces are the same ones as in the final polyhedron, and such that the magnitude of the force experienced by the vertices does not vary too greatly, causing slow convergence or numerical instability. We here outline two methods for obtaining initial geometries that are suitable for input to the fullerene force-field optimization.

The Fowler-Manolopoulos 3D embedding algorithm (also called Adjacency Matrix Eigenvector or AME algorithm) diagonalizes the \(N \times N\) adjacency matrix \(A_{ij}\) of the fullerene graph to obtain eigenvectors \(\mathbf{x}_i\) (\(i = 1, \ldots, N\)), of which three \('P\)-type eigenvectors' with one nodal plane are chosen to construct the Cartesian coordinates (termed topological coordinates) by Manolopoulus and Fowler for the initial structure.\(^{31,107}\) This often yields suitable coordinates because fullerenes can be modeled as spherical aromatic systems fulfilling approximately the Laplace differential equation with real spherical harmonics \(Y_{lm}(\theta, \phi)\) as eigenfunctions, of which the three \(Y_{lm}(l = 1)\) are associated with the coordinates \((x_i, y_i, z_i)\).\(^{31,108}\) Cartesian coordinates constructed in this way already reflect the space symmetry of the fullerene. The \('P\)-type eigenvectors can easily be found for near-spherical fullerenes like \(C_{20}\) or \(C_{60} - I_h\), because the sequence of eigenvalues mimics the spherical harmonics. However, for distorted fullerenes, or for very large fullerenes where the spectrum of the adjacency matrix becomes very dense with quasi-degeneracies, it becomes difficult to find the appropriate eigenvectors, and the assumption that the eigenvectors resemble spherical harmonics becomes decreasingly valid. For example, for large fullerene nano-tubes aligned around the \(z\)-axis, such as the one shown in Figure 2, the eigenvalues belonging to the \('P_x\) and \('P_y\) type eigenvectors do not lie close to the \('P_z\) eigenvalue. Hence, one must search for the appropriate set of eigenvectors, and identifying them is not always easy.\(^{109-111}\) Second, the bond lengths tend to vary substantially in the constructed fullerene, leading to unreasonable large bond distances once the minimum bond distance is set to \(\approx 1.4\ \text{Å}\), and a scaling procedure for the eigenvectors may only solve part of this problem.\(^{31}\) Finally, because it is often necessary to search for the right eigenvectors and hence calculate a large number of them, the algorithm scales up to \(O(N^3)\), which quickly becomes a heavy operation as \(N\) grows.

We have found it useful to sacrifice some accuracy in the initial geometry for a method that is more stable, and which scales approximately as \(O(N)\). The fullerene force-field optimization is insensitive enough to the starting geometry that the difference is not noticeable. The method starts with any planar embedding of the fullerene graph. We choose the Tutte-embedding, which is guaranteed to be planar, and can be computed in near-linear time by solving two sparse linear systems of equations.\(^{54}\) The planar embedding is then mapped onto the surface of a sphere (or e.g., ellipsoid, if the fullerene is a priori known to be elongated) in the following way: For every vertex \(v\), let the topological distance from the outer face (the 'depth' of the vertex) be \(d_v\), and its coordinate in the Tutte-embedding be \(\mathbf{x}_v\). Let \(D\) be the maximum depth, and \(\mathbf{x}_c\) be the barycenter in the embedding. We assign one angle \(\phi(v) = \langle d_v, +0/2\pi/D + 1\rangle\), placing vertex layers equidistantly along the \(\phi\)-angle according to their depth. The second angle \(\theta(v)\) is the angle of \(\mathbf{x}_v\) around \(\mathbf{x}_c\). Then assigning the angles \(v \mapsto (\theta(v), \phi(v))\) yields an embedding on the sphere with no crossings, since the Tutte-layout is guaranteed to be planar, but which avoids the exponential crowding of the planar Tutte-embedding, such that the vertices are distributed in a reasonable manner across the surface of the sphere.\(^{b}\) It remains, however, a challenge to embed the fullerene graph on a more realistic surface.

### Fullerene Force-Field

This section describes how to obtain good results for the molecular geometry numerically by way of specially tailored force-field optimization methods. This yields geometries that are very close to the ones obtained by long and computationally heavy quantum chemical calculations, as we shall see.

It is common for molecular force-field optimizations to distinguish between single and double bonds, even introducing different types of single and double bonds depending on atoms and groups in close proximity. For each bond, that is, edge in the graph, one sums over all the Kekulé structures and assigns a fraction of bond character to each edge. This strategy is not practical for fullerenes, even for small cages like \(C_{60}\) due to exponential growth in perfect matchings and hence Kekulé structures, as discussed below. Force fields for fullerenes should therefore be designed in a way that avoids explicit treatment of double bonds. As a good approximation it is assumed that the properties of bonds, angles, and dihedral angles in a fullerene are determined by the size of their neighboring faces alone. The force fields discussed below are specifically designed for fullerenes based on this assumption.

The first force field tailored to fullerenes by Wu et al.\(^{114}\) was designed for \(C_{60} - I_h\) and uses harmonic force field terms. It distinguishes between two bond types: bonds adjacent to two hexagons, and bonds
adjacent to a pentagon and a hexagon, which are optimized for bond lengths $r_0$ with values of 1.54 Å and 1.41 Å, respectively. Angles are either part of a pentagon ($\theta_0 = 3\pi/5$) or a hexagon ($\theta_0 = 2\pi/3$). The force constants $k_1$ through $k_4$ are given (in $10^3$ dyn/cm) as 11.0, 10.0, 1.0, and 1.0. No further parameters are taken into account.

In the following years a number of extensions and modifications to the Wu force field were published, taking into account an increased number of parameters. These extensions include improved force constants, torsions, mixed terms between second and third neighbors, mixed terms between distances and angles, and distance-to-angle variations. Except for the force field by Ceulemans et al., all above mentioned force fields have been exclusively designed for and applied to C$_{60}$. The program Fullerene$^{35}$ includes a 22-parameter force field with explicit dihedral angles. It takes into account three types of distances (bonds that are part of 0, 1, or 2 pentagons), two angles (part of a pentagon or a hexagon) and four types of improper dihedral angles, defined between a vertex and its three neighbors (where the starting vertex is part of 0, 1, 2, or 3 pentagons). It is applicable to all fullerenes and yields structures that are in very good agreement with DFT optimized structures. For example, for C$_{380}$-$^T$ $T$, the rms error between the B3LYP/SVP and the force field optimized structure is only 0.038 Å. For many purposes, the force-field optimized molecular geometry, which is computed in milliseconds, is good enough to be used instead of a more computer time intensive quantum chemical optimization (although semiempirical methods including density functional based tight-binding are also very computer time efficient and have yielded good results$^{122}$).

**Fullerene Symmetry**

There are two symmetry groups associated with a fullerene: the ideal or topological symmetry group of the fullerene graph, and the real or physical symmetry group of the molecule in 3D space. The physical symmetry group is a subgroup of the ideal one, due to (first or second order) Jahn-Teller distortions, variation of bond-lengths, or other effects from external perturbations. But because the shape is often only mildly altered from the ideal shape, the ideal symmetry group carries useful information about the physical system.

A surprising theorem by Mani$^{123}$ shows that any three-connected cubic planar graph can be embedded in space as a convex polyhedron, the point group of which realizes the full symmetry group of the graph. That is: every graph automorphism of a fullerene is also a rotation or reflection of its ideal polyhedral shape. In addition, it can be shown$^{124}$ that the point group for a fullerene must be one of the 28 point groups listed in Table 1. The largest group is the icosahedral group $I_h$, of order 120, and each of the 27 remaining attainable point groups are subgroups of $I_h$.

Can we automatically determine the symmetry group for the fullerene directly from the graph, without referring to spatial coordinates? (The latter requires first finding a maximally symmetric 3D embedding) For general graphs, this is a difficult problem. However, in the case of cubic polyhedral graphs, for which general face spirals determine isomorphism, it is both simple and efficient to do so.

Before moving on, we take note of an ambiguity in terminology. The terms symmetry group and automorphism group often refer simply to the group formed by operations that leave the system invariant. In the group theoretical sense, a group is fully defined by its multiplication table. Point groups, however, are not just groups, but are isometries of space and carry additional information about how they transform space. For example, the point groups $D_6$, $C_{6v}$, $D_{3h}$, and $D_{3d}$ are all the exact same group, but differ as point groups. It is unfortunate for this reason that ‘point groups’ are called ‘groups’. To make the distinction explicit, we will use the term abstract group for the symmetry group in the group theoretical sense, and either point group or symmetry group for the isometries.
Assume that we are given the fullerene graph dual \( G^* \), constructed from a generalized spiral \( S = (d_1, d_2, \ldots, d_F) \), and wish to compute the automorphism group \( G \) of \( G^* \). Since \( G^* \) is constructed from \( S \), the entries in \( S \) are the degrees of vertex number \( 1, 2, \ldots, F \) in \( G^* \). For every vertex \( v \) of degree \( d_1 \), we have \( 2d_1 \) different spiral starts: \( d_1 \) for clockwise and \( d_1 \) for counter-clockwise traversal. If a spiral start \((f_1, f_2, f_3)\) unwinds \( G^* \) to the input spiral \( S \), there is an automorphism of \( G^* \) that maps \((1,2,3)\)\( \mapsto (f_1, f_2, f_3) \). These are all the automorphisms, and the number of starts that unwind to \( S \) is the order \( |G| \) of the automorphism group. This construction lets us easily obtain the actual group: The spiral start determines the entire spiral, and by simply writing down the vertex names in \( G^* \) while unwinding the spiral, we obtain a permutation representation of the group element

\[
\pi_G(g) = \left( \begin{array}{cccc} 1 & 2 & 3 & \cdots & F \\ f_1 & f_2 & f_3 & \cdots & f_F \end{array} \right)
\]

(9)

This is a faithful (but not irreducible) representation of the group, and if we wish, we can easily build the multiplication table by composing all pairs of the permutations. Similarly, we can calculate characters, irreducible representations, and all other properties, and we can identify the group. There are a number of ways to make this efficient. In Wirz et al. (unpublished manuscript), we present a general, but efficient, \( O(N) \)-algorithm that works for all cubic polyhedral graphs, including all fullerenes and fulleroids.

The permutation representation \( \pi_G \) allows us to not only find the abstract symmetry group, but contains enough information to identify the point group, that is, the isometries of the three-dimensional fullerene polyhedron. Fowler and Manolopoulos published a program for identifying the point group of any spiralable fullerene in Ref 31, based on site symmetries and counting group orbits. The symmetry points of interest in a fullerene are the vertices, midpoints of edges, the barycenter of the polygons and the whole cage, the latter having the full symmetry of the point group. They have certain site symmetries according to the rotational axes or mirror planes going through these symmetry points,\(^{31,125,126}\) which are collected in Table 2. The full isometry group \( G \), which correspond to the rotations, reflections, roto-inversions, and inversions that leave the ideal polyhedron invariant, also act as permutations of the symmetry points,\(^{107,125,126}\) the action always being a subgroup of \( G \). The permutations of the symmetry points in fact completely determine the point group symmetry, and we can find the full point group of the graph \( G \) (or, equivalently, its dual \( G^* \)) as follows:

\[
\begin{align*}
\text{Vertices} & : \ C_{3v}(6), \ C_3(3), \ C_2(2), \ C_1(1) \\
\text{Edge centers} & : \ C_{5v}(10), \ C_5(5), \ C_2(2), \ C_1(1) \\
\text{Pentagon centers} & : \ C_{6v}(12), \ C_6(6), \ C_3(6), \ C_2(3), \ C_1(2), \ C_1(1) \\
\text{Hexagon centers} & : \ C_{6v}(24), \ C_6(12), \ C_3(12), \ C_2(6), \ C_1(6) \\
\text{Cage centers} & : \ \text{Full point group}
\end{align*}
\]

The order of the site-symmetry group is given in parentheses (the maximum value of any site symmetry group order is 12).

In step 1, one computes the face permutation representation \( \pi_F \) of the abstract group, as described above. From this one derives vertex and edge permutation representations \( \pi_V \) and \( \pi_E \) by acting with every group operation on the dual graph.

In step 2 one computes the vertex, edge, and face group orbits by acting with the permutation representations on every vertex, edge, and face.\(^{127}\) Each orbit belongs to a certain site-symmetry group as shown in Table 2. The site-symmetry groups are determined by the orbit sizes. For example, the site-symmetry group of the face \( f \) has order \( |G_f| = |G|/|G_f| \). By counting the number of sites belonging to each site-symmetry group, we obtain a signature that uniquely identifies the point group:

\[
\begin{align*}
\text{Vertices} & : \ m_V(k) = \left\{ v \in V \mid k = \frac{|G|}{|G_v|} \right\}, \\
\text{Edges} & : \ m_E(k) = \left\{ e \in E \mid k = \frac{|G|}{|G_e|} \right\}, \text{ and} \\
\text{Faces} & : \ m_F(k) = \left\{ f \in F \mid k = \frac{|G|}{|G_f|} \right\}
\end{align*}
\]

(10)

The information can be condensed to a site-symmetry count for each site-group order:

\[
m_s(k) = m_F(k) + m_E(k) + m_V(k)
\]

(11)

In the final step, the point group is determined by the site-symmetry counts by the decision tree structure shown in Figure 13. The method can be extended to any fullerene and fulleroid by using general spirals. However, for every possible point group, the site-symmetry signature must be worked out and added to the decision tree.

Once the point group has been determined, the number of infra-red and Raman active lines, as well as the \( ^{13} \text{C} \) NMR pattern can be derived.\(^{31,107}\) Moreover, point groups lacking an inversion center
are further divided into polar and chiral point groups. A chiral point group is one without any roto-inversion symmetry elements, and a polar point group allows for the fullerene to have a dipole moment. A point group with an inversion center or a mirror plane perpendicular to the axis of rotation cannot be polar. The nine chiral point groups for fullerenes are \( I, T, D_6, D_5, D_3, D_2, C_3, C_2, \) and \( C_1. \) The polar fullerenes belong to either of the point groups \( C_{3v}, C_3, C_{2v}, C_2, C_1 \) or \( C_1. \) For larger fullerenes, the fraction of low-symmetry to high-symmetry isomers grows rapidly, and the \( C_1 \) point group increasingly dominates. This can be intuitively understood combinatorially from distributing pentagons on a sphere. Already at \( C_{100}, \) more than 99% of the isomers are \( C_1. \)

The separation of spiral starts into equivalence classes by their spirals yields an interesting relation:

\[
|G| = \frac{6N}{N_S} \quad (12)
\]

where \( N_S \) is the number of symmetry distinct (general) spirals. For \( I_h \) symmetry we have \( |I_h| = 120, \) which gives \( N = 20N_S. \) Hence, \( I_h \)-fullerenes can only occur when \( N \) is a multiple of 20. In a similar way, for \( I \) symmetry we have \( N = 10N_S, \) and for \( T_d, T, D_6, \) or \( D_{6d} \) we obtain \( N = 4N_S. \) This explains why some of these high symmetry groups are not found in certain isomer lists.

In the case of icosahedral symmetry, we have a full characterization of when they occur: As described by Dutour and Deza, \( I_h, I, \) or \( I \) symmetry is a Goldberg-Coxeter transform of \( C_{20}. \) This means that they occur exactly when \( N = 20(k^2 + kl + l^2) \) for integers \( k \) and \( l. \) It is also possible to determine when these are of \( I_h \) and when they are of \( I \) symmetry. Halma and leapfrog transformations, which correspond to Goldberg-Coxeter transforms of \( l = 0 \) and \( k = l, \) respectively, both preserve symmetry. Hence, there is a fullerene with \( I_h \) symmetry at \( N = 20^2 \) and \( N = 20(3j^2) \) for every \( j \in \mathbb{N}, \) corresponding to a single Halma or leapfrog transformation on \( C_{20}. \) Consecutive application of the two shows that isomers with \( I_h \) symmetry are found for every \( N = 20(3j^2) \) \((j \in \mathbb{N} \text{ and } i = 0, 1). \) Moreover, general Goldberg-Coxeter \((k, l)\) transforms with \( k \neq l \) and \( l \neq 0 \) \((k > 0)\) break horizontal mirror plane symmetry.

**Shapes: Volume and Surface Area, Sphericity, and Convexity**

For ideal \( C_{60} - I_h, \) where all edges are of equal length, the geometric volume \( V, \) and surface area \( A \) can be obtained through simple algebraic and geometric considerations. The volume and surface are of a regular 20-sided polyhedron (icosahedron, see Figure 3) made out of equilateral triangles with edge length \( a \) is,

\[
V_{ico} = \frac{5}{12} (3 + \sqrt{5}) a^3 \quad \text{and} \quad A_{ico} = 5\sqrt{3}a^2 \quad (13)
\]

Cutting out 12 pentagonal pyramids at length \( a/3 \) gives the ideal truncated icosahedron \( C_{60} - I_h \) with

\[
V_{C_{60}} = \frac{1}{4} (53 + 43\sqrt{5}) r_e^3
\]

and \( A_{C_{60}} = 3r_e^2 \left( \sqrt{5} + 10\sqrt{5} + 10\sqrt{3} \right) \)

(14)
where \( r_e = a/3 \) is the length of the edge (C–C bond length). This gives the ratios \( V_{i60}/V_{i60} \cong 1.065 \) and \( A_{i60}/A_{i60} \cong 1.073 \) between the two volumes and surface areas. \( C_{60} \) has, however, two different bond lengths as the crystal structure shows,\(^1\) \( r_5 = 1.455 \) Å (edge at pentagon-hexagon fusion) and \( r_6 = 1.391 \) Å (edge at hexagon-hexagon fusion). A smaller bond distance in the hexagon implies that more of the original icosahedron is cut off,

\[
V_{i60} = \frac{5}{12} (3 + \sqrt{5}) (2r_5 + r_6)^3 - \frac{(5 + \sqrt{5})}{2} r_5^3 \quad \text{and} \quad A_{i60} = 3r_5^2 \sqrt{5^2 + 10\sqrt{5} + 5\sqrt{3}} (r_6^2 + r_5^2 + 4r_5r_6)
\]

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For \( r_5 = r_6 \) we obtain Eq (14). If \( r_6 = 0 \) \((r_5 = a/2)\) the hexagons become equalateral triangles and for \( r_5 = 0 \) we just get a hollow icosahedral \( C_{12} \) cluster. For \( C_{20} \) (equal edge lengths) the vertices lie in the center of each face of an icosahedron, and the volume and surface area are easily obtained,

\[
V_{i20} = \frac{15 + 7\sqrt{5}}{4} r_e^3 \quad \text{and} \quad A_{i20} = 3r_e^2 \sqrt{5^2 + 10\sqrt{5}}
\]

\(^1\) Mathematically, this translates into finding the solution to the problem\(^1\)\(^2\).\(^1\)\(^3\) Mathematically, this translates into finding the solution to the problem\(^1\)\(^2\).\(^1\)\(^3\) Mathematically, this translates into finding the solution to the problem\(^1\)\(^2\).\(^1\)\(^3\) Mathematically, this translates into finding the solution to the problem\(^1\)\(^2\).\(^1\)\(^3\)

For \( r_5 \neq r_6 \) and \( r_5 = 0 \) \((r_6 = a/2)\) the hexagons become equalateral triangles and for \( r_5 = 0 \) we just get a hollow icosahedral \( C_{12} \) cluster. For \( C_{20} \) (equal edge lengths) the vertices lie in the center of each face of an icosahedron, and the volume and surface area are easily obtained,

\[
V_{i20} = \frac{15 + 7\sqrt{5}}{4} r_e^3 \quad \text{and} \quad A_{i20} = 3r_e^2 \sqrt{5^2 + 10\sqrt{5}}
\]

\(^1\) Mathematically, this translates into finding the solution to the problem\(^1\)\(^2\).\(^1\)\(^3\) Mathematically, this translates into finding the solution to the problem\(^1\)\(^2\).\(^1\)\(^3\) Mathematically, this translates into finding the solution to the problem\(^1\)\(^2\).\(^1\)\(^3\)

How can we get the volume \( V \) and surface area \( A \) for any fullerene isomer? As fullerene cages are, however, a number of definitions according to which we can express both quantities. We could be used for calculating some important measures for fullerenes, such as the *sphericity* \( S \) (how spherical a fullerene is) and *convexity* \( C \) (how convex a fullerene is). The surface of a triangulated fullerene may not be convex. We can measure the non-convexity by comparing to the convex hull of the fullerene cage, which is the smallest convex polyhedron that contains all the points. The convex hull (CH) is uniquely defined, and there are several algorithms available, such as the incremental 3D convex hull algorithm.\(^1\)\(^2\) A measure for convexity \( C \) is obtained from the proportion of ‘empty space’ between the non-convex polyhedron and the convex hull by comparing the volumes or surface areas, that is, \( C_A = A/A_{CH} \) and \( C_V = V/V_{CH} \).

The simplest measure of sphericity is the *isoperimetric quotient* \( q_{IPQ} \),\(^1\)\(^3\) defined for a polyhedron as

\[
d_{IPQ} = \frac{36\pi}{A^2} \quad \text{with} \quad q_{IPQ} \in [0, 1]
\]

\(^1\)\(^3\) Mathematically, this translates into finding the solution to the problem\(^1\)\(^2\).\(^1\)\(^3\) Mathematically, this translates into finding the solution to the problem\(^1\)\(^2\).\(^1\)\(^3\) Mathematically, this translates into finding the solution to the problem\(^1\)\(^2\).\(^1\)\(^3\)

where \( q_{IPQ} = 1 \) for an ideal sphere and \( D_{IPQ} \) is a measure of the deviation from an ideal spherical shape. \( D_{IPQ} \) is shown for several fullerenes in Figure 14.

We usually regard \( C_{20} - I_h \) and \( C_{60} - I_h \) as almost spherical because all vertices lie on a (covering) sphere. We may therefore analyze whether or not the vertices of a given fullerene lie on a sphere. For this, we define the *minimum covering sphere* as a sphere of minimum radius that encloses all vertices in the polyhedral embedding (J. J. Sylvester, 1857).\(^1\)\(^2\)\(^3\) Mathematically, this translates into finding the solution to the problem\(^1\)\(^2\).\(^1\)\(^3\)

\[
R_{MCS} = \min_{\vec{c}} \max_{\vec{v}_i} ||\vec{v}_i - \vec{c}_{MCS}||
\]

\(^1\)\(^2\)\(^3\) Mathematically, this translates into finding the solution to the problem\(^1\)\(^2\).\(^1\)\(^3\) Mathematically, this translates into finding the solution to the problem\(^1\)\(^2\).\(^1\)\(^3\)

where \( || \cdot || \) denotes the Euclidian norm, \( \vec{v}_i \) is the location of the vertex \( v_i \) in 3D space, and \( \vec{c}_{MCS} \) is the center of the MCS with radius \( R \) and is usually close to the barycenter of the fullerene. The MCS is uniquely defined\(^1\)\(^2\).\(^1\)\(^3\) and can be obtained using an efficient algorithm, such as the one proposed by Yildirim.\(^1\)\(^3\) The convex hull of a 3D embedding of a graph \( G \) is contained entirely within the MCS. For fullerenes with an ellipsoidal shape like \( C_{72} - D_{6d} \), or with a cylindrical shape such as a nanotube\(^1\)\(^3\) (see Figure 2), it is more appropriate to use a minimum covering ellipsoid or cylinder instead of a sphere. The ellipsoidal problem has been addressed recently, and is known as the *minimum volume axis-aligned ellipsoid problem* (MVAE).\(^1\)\(^2\)\(^3\)

We can now define a number of useful measures for sphericity. The *MCS distortion* (normalized
Various deformation parameters $D$ (in percent) for a series of fullerenes selected according to stability. For larger fullerenes, Goldberg-Coxeter transformed structures of C$_{20}$ were chosen. Geometries were obtained from DFT (up to C$_{540}$) or force field optimizations. IPQ: isoperimetric quotient; MCS: Minimum covering sphere; MDS: Minimum distance sphere; DTP: Diaz-Tendero parameter; FAP: Fowler asymmetry parameter; TEP: $(1 - \rho)$ from the topological efficiency parameter $\rho$.

Figure 14 shows a comparison of sphericity parameters for a number of stable fullerenes. As C$_{20}$ is slightly deformed due to Jahn-Teller distortion, the vertices do not lie exactly on a sphere anymore, and $D_{\text{MDS}} \neq 0$. In contrast, the purely topological efficiency parameter $(1 - \rho)$ (see discussion in the topological indicator section) is exactly zero for C$_{20}$ (and for C$_{60}$, C$_{80}$, and C$_{240}$). For C$_{60}$, all vertices lie on a sphere and all deformation parameters are zero, except for the IPQ, which except for its simple definition is perhaps not the best measure for sphericity. All deformation parameters reveal the highly deformed fullerenes (spikes in Figure 14).

Toward the Solid State: Hilbert’s Problem and Space Fillings
Solid C$_{60}$ packs in a rotationally disordered face centered cubic (fcc) structure with a lattice constant of $a_{\text{fcc}} = 14.17 \pm 0.01$ Å, which according to Fischer et al. implies a close packing of spheres of a diameter of 10.02 Å. C$_{60}$ undergoes a phase transition below 255 K to a simple cubic (sc) structure, from which one can infer a lattice parameter of $a_{\text{sc}} = 14.04 \pm 0.01$ Å. The only other solid state fullerene obtained by experiment is that of C$_{70}$ - $D_{5h}$, which adopts a closed packed structure of either hcp or fcc symmetry with a fcc lattice constant of $a_{\text{fcc}} = 14.96$ Å. In the solid state, the interactions between the fullerenes are of Van der Waals type. Note that the polarizability and therefore the Van der Waals coefficients grow linearly with increasing number of carbon atoms in the fullerene cage, which should converge toward the graphene limit for the per-atom value. Experimentally, the solid state behavior of C$_{60}$...
has been studied in great detail. One interesting fact is that the bulk moduli of C₆₀ (B = 6.8 GPa for the simple cubic phase and 8.8 GPa for the fcc phase) and C₇₀ (11 GPa for the rhombohedral phase) are quite small as applied pressure will just squeeze out the compressible Van der Waals space. These bulk moduli are only twice as large compared to the rare gas solids, and much smaller than that of diamond (442 GPa). However, once the Van der Waals space is squeezed out and the fullerene cages touch, the bulk modulus increases substantially, as the fullerene cage is not easily compressible, similar to the high in-plane stiffness of graphite or graphene. We mention that it requires high pressures above 20 GPa to turn C₇₀ into graphite (using step-like shock-wave compression). For overviews on solid-state properties of fullerenes, see Zettl and Cumings or Fischer et al.

From the experimental data of C₆₀ - Ih, we can adjust the Van der Waals radius of the C₆₀ cage such that the experimental lattice constant is obtained, that is, we use a hard sphere model for periodic packing (fcc is the closest packing available for hard spheres as proven by computer by Hales in 2000). We can do this by using the MCS introduced in Eq (19), but we have to add the carbon Van der Waals radius (rᵥᵥ(C) = 1.415 Å) to the MCS radius,

\[ R_{ᵥᵥ}(C₇₀) = R_{MCS}(C₇₀) + r_{ᵥᵥ}(C) \]  

(25)

Figure 15 shows lattice constants for a number of fullerenes using this hard sphere model, which can be taken as an upper bound to experimental lattice constants. For the other experimentally known fullerene crystal structure, C₇₀, the fcc lattice constant afcc of 14.96 Å is in reasonable agreement with the hard-sphere value of 15.78 Å, considering the fact that the IPR C₇₀ isomer is slightly prolate and distorted. The dependence of the afcc lattice constant on the vertex number can be expressed as afcc = c√N + (4Rᵥᵥ(C))/√2, where the shift follows from the geometry of the fcc cell and the \( \sqrt{N} \)-term represents the dependence of the fullerene radius on N. In the hard sphere model we have afcc ∼ R, and the cage radius of the fullerene is R ∼ \( A^{1/2} \sim N^{1/2} \) (A being the surface area of the MCS).

For distorted fullerenes, the packing problem becomes far more complicated. For example, we could add the Van der Waals radius to every carbon atom and take the convex hull around this Van der Waals layer. We are then faced with close packing of complicated polyhedral structures, which is a difficult, unsolved problem. However, there exist algorithms for packing arbitrary polyhedra in finite spaces, and it is possible that one could approximate properties of the (periodic) close packing solution by making the finite cell large enough. For some fullerenes, ellipsoidal, or cylindrical covers are more appropriate as already mentioned. The close packing of ellipsoids was recently investigated by Donev et al., who showed that the maximum packing density can exceed that of a close sphere packing. Nevertheless, the question still remains how fullerenes of a certain topology pack in 3D space. This is related to one of Hilbert’s fundamental problems: How can one arrange most densely in space an infinite number of equal solids of given form, e.g., spheres, with given radii, that is, how can one so fit them together that the ratio of the filled to the unfilled space may be as great a possible? A more rigorous mathematical proof (beside Hales’ complete computer algorithm) for fcc packing of hard spheres as the most dense packing is still missing, and packing any other deformed spheres remains an open area of research. One possible approximation is to expand the shape of a fullerene in terms of multipole deformations and study the 3D packing of such smooth topologies.

**Topological and Chemical Indicators**

Chemists can deduce many useful properties of a chemical system just by looking at its structure. For fullerenes, the distribution of the 12 pentagons on a surface, for example, a sphere, can tell us qualitatively how stable the fullerene is or how it would pack in the solid state. Moreover, the symmetry of the underlying structure determines many useful spectroscopic properties. It is perhaps a realistic goal to connect the graph theoretical properties of a fullerene directly with its physical properties by mapping the fullerene graph G into a number describing that property. In more general terms, we define a topological indicator...
as a map $\tau_i$ from the graph $G$ into a finite series of numbers,

$$\tau_i : G \mapsto \{ x_1, \ldots, x_n \} \quad (26)$$

where the numbers $x_i$ are called topological indices and can be chosen to be integers, rationals, or reals. A topological index is called a chemical index if it is related to a chemical (or physical) property. Topological indicators may be placed in the same category as crude chemical bonding models, except that they can be strictly defined in graph theoretical terms and sometimes have interesting mathematical properties. There is no restriction that the mapping between fullerene graphs and any topological indicator be one-to-one, and indeed, most of the commonly used topological indicators are the same for many different isomers. Of course, any chemical or physical property is ruled by its underlying electronic structure governed by the Schrödinger equation. However, solving the electronic structure problem for any large fullerene is a daunting task and therefore, topological indices connected to the adjacency matrix are more often used.

The first topological index of chemical relevance was introduced by Wiener in 1947.\textsuperscript{164} He defined a so-called path number (Wiener index) as the sum of the entries in the topological distance matrix $d_{ij}$ consisting of the length of the shortest path between every pair of vertices in the chemical graph representing the non-hydrogen atoms in the molecule,

$$W(G) = \frac{1}{2} \sum_{i \in \mathbb{V}} d_{ij} \quad (27)$$

where $d_{ii} = 0$. The Wiener index provided a good measure of compactness for acyclic alkanes and gave a reasonable correlation to boiling points.\textsuperscript{164} The first application of the Wiener index to fullerenes came from Ori and co-workers, who obtained $W = 8340$ for $C_{60} - I_h$.\textsuperscript{163} For fullerenes, low Wiener indices provide a measure of high compactness of the cage. This can be seen from the scaling law, which has been found to behave like $\mathcal{O}(N^{5/2})$ for ‘spherical’ fullerenes\textsuperscript{166} and $\mathcal{O}(N^3)$ for fullerene nanotubes. For carbon nanotubes, analytical formulae in terms of polynomials in the vertex number are available, for example, for the smallest $D_{5b}/D_{5d}$ nanotubes we have $W(k) = 1/3(100k^3 + 1175k - 2100)$ with $N = 10k$ and $k \geq 5$,\textsuperscript{167} and for the smallest $D_{6h}/D_{6d}$ nanotubes we have $W(k) = 12(4k^3 + 69k - 136)$ with $N = 12k$ and $k \geq 7$.\textsuperscript{168} It is conjectured that the Wiener index grows like $\mathcal{O}(N^{2+1/d})$, where $d$ describes the dimensionality of the system (the Wiener dimensionality).\textsuperscript{166} The almost cubic growth with increasing number of vertices can make the Wiener index unwieldy for larger fullerenes. Therefore, Ori and coworkers defined a topological efficiency index $\rho$, derived from the Wiener index\textsuperscript{165,170}

$$\rho = 2 \frac{W(G)}{NW_{\text{min}}} \quad \text{with} \quad W_{\text{min}} = \min_{i \in \mathbb{V}} \left\{ \sum_{i \in \mathbb{V}} d(i, j) \right\} \quad (28)$$

which has the advantage that, unlike the Wiener index, it does not grow to large numbers. Small values of $\rho \geq 1$ indicate topologically efficient structures, for example, both $C_{20} - I_h$ and $C_{60} - I_h$ have $\rho = 1$.\textsuperscript{168} It can be seen as a measure of sphericity that does not involve 3D Cartesian coordinates, as shown nicely in Figure 14. Vukićević et al. showed that among the 4478 isomers of $C_{60}$, the joint information of the Wiener index and the topological efficiency index correctly identifies the $C_{60} - C_{20}$ (11) as the most stable molecule\textsuperscript{169} (as detected experimentally in endohedral $Sc_2@C_{60}$\textsuperscript{171}).

Many different topological indices have been introduced and studied since, mainly for the structural and statistical analysis of molecules, polyhedra, and graphs in general, often yielding interesting mathematical properties (for a list of topological indices see for example Ref 172). The distance matrix $d_{ij}$ gives rise to a number of very useful topological indices including the topological radius $R$ and diameter $D$,\textsuperscript{173} Hosoya polynomials,\textsuperscript{174} and related Wiener indices, the Szeged index $S$,\textsuperscript{175} and the Balaban index $J$,\textsuperscript{29,176} to name but a few (for fullerenes one has $W \sim S$, and $W \sim J$).\textsuperscript{29,177} There are also a number of topological indices connected to the adjacency matrix (or Hückel matrix), for example Hückel orbital energies and related properties (occupation numbers, band gap,\textsuperscript{31} spectral moments\textsuperscript{178} etc.),\textsuperscript{31} and the Estrada index.\textsuperscript{179,180} For a more detailed discussion of topological indices see Refs 29, 89, 173, and 181.

Some of these topological indices can directly be related to the stability of the fullerene cage. For example, the Fowler-Manolopoulos pentagon indices $\{p_n\mid n = 0, \ldots, 5\}$ define the number of pentagons attached to $n$ other pentagons ($n$ is called the neighbor index),\textsuperscript{31} that is, for IPR fullerenes we have $p_0 = 12$ and $p_n = 0$ for $n \geq 1$. For any fullerene, we of course must have $p_1 + \cdots + p_5 = 12$. Fowler-Manolopoulos
hexagon indices are similarly defined and useful for IPR fullerenes, that is, \( \{b_i| i = 0, ..., 6\} \), where \( b_n \) defines the number of hexagons with neighbor index \( n \). In an IPR fullerene every hexagon is adjacent to a minimum of three others and we can restrict the list of hexagon neighbor indices for example to \( (h_3, h_4, h_5, h_6) \).\(^{31}\)

We may contract the neighbor indices to one useful topological index describing the stability of fullerenes.\(^{31}\) The single pentagon signature \( P_1 \) is defined as

\[
P_1 = \frac{1}{2} \sum_{k=1}^{5} k p_k
\]

\( \text{max}(P_1) = 30 \) (for \( C_{20} \) only), and for IPR fullerenes such as \( C_{60} \) we have \( P_1 = 0 \). For IPR fullerenes a more useful single topological index is defined through the hexagon signatures. The standard deviation \( \sigma_h \) of the hexagon neighbor index distribution is defined as

\[
\sigma_h = \sqrt{\langle k^2 \rangle - \langle k \rangle^2}
\]

where

\[
\langle k^n \rangle = \frac{H_n}{H_0} \quad \text{with} \quad H_n = \sum_{k=0}^{6} k^n b_k \in \mathbb{N}
\]

and

\[
H_0 = \sum_{k=0}^{6} b_k = \frac{N}{2} - 10
\]

We call the topological index \( H_n \) the \( n \)-th moment hexagon signature. For general fullerene isomers, low \( P_1 \) values and high \( H_1 \) values correlate with high thermodynamic stability.\(^{31}\) It turns out, however, that \( H_1 \) stays constant for a given vertex count in IPR fullerenes, as this index is related to certain face patterns as we shall see. Fowler et al. advocated the use of the index \( \sigma_h \) instead, as low values of \( \sigma_h \) are better predictors for stability, and the strain is minimized when all hexagon neighbor indices are as similar as possible.\(^{31}\) For example, for \( C_{60} \) we have \( \{b_i\} = \{0, 0, 0, 20, 0, 0, 0\} \) giving \( H_1 = 60, H_2 = 180 \), and \( \sigma_h = 0 \). Fowler et al. devised the following scheme for estimating the internal order among fullerenes sorted by stability:\(^{173}\) First order the isomers according to lowest \( P_1 \)-value (each pentagon-pentagon fusion carries a penalty of about 20–25 kcal/mol). If multiple IPR fullerenes are present, that is, \( P_1 = 0 \), we order these first by largest \( H_1 \)-value, and finally by lowest Wiener index \( W \) (which indicates high compactness of the fullerene cage). The result is an efficient but effective screening method to find the most stable fullerene isomers.

Ju et al. presented a relationship between the Fowler-Manolopoulos hexagon indices and a particular hexagon structure count to provide a graphical interpretation.\(^{182}\) The complete list of different pentagon and hexagon combinations up to three connected faces are shown in Figure 16. Note that for connecting two or three faces we have

\[
n_{55} + n_{56} + n_{66} = E = \frac{3N}{2} \quad \text{and}
\]

\[
n_{555} + n_{556} + n_{566} + n_{666} = \frac{11}{2}N - 30
\]

where the notation is described in Figure 16. For example \( n_{556} \) is the sum of all occurring ring patterns containing the combination of two pentagons and one
hexagon. Ju et al. showed that for the second moment hexagon signature one gets\textsuperscript{182}

\[ H_{2}^{IPR} = 2n_{66} + 6n_{666} + 2\left(n_{666} + n_{6666}\right) = 2\left(n_{666} + n_{6666}\right) + 9N - 480 \]  

(33)

and

\[ H_{1}^{IPR} = 2n_{66} = 3\left(N - 40\right) \]  

(34)

Hence for IPR fullerenes, \( \sigma_{b} \) and \( H_{2} \) are related through the simple equation

\[ \sigma_{b}^{IPR} = \left(N - 20\right)^{-1} \left(2\left(N - 20\right)H_{2} - 36\left(N - 40\right)^{2}\right)^{1/2} \]  

(35)

and one requires only \( H_{2} \) to discuss the stability of IPR fullerenes. Stevanović extended these relations to general fullerenes (IPR or not),\textsuperscript{184}

\[ H_{1} = 3\left(N - 20\right) - n_{56} \]  

(36)

and

\[ H_{2} = 18\left(N - 20\right) - \left(6n_{56} + 2n_{666} + n_{6666} + n_{66666}\right) \]  

(37)

This not only illustrates nicely the current development in the area of topological indices, it also suggests that the stability of fullerenes can be approximated by counting different face patterns.\textsuperscript{183}

Indeed, Cioslowski et al.’s incremental scheme for the heat of formation \( \Delta H_{f}^{IPR} \) for IPR fullerenes\textsuperscript{33} uses 25 linearly independent face patterns (structural motifs) of up to 13 connected hexagons and pentagons to calculate the energy \( E_{FP} \), including an additional curvature term \( E_{C} \),

\[ \Delta H_{f} = E_{FP} + E_{C} = \sum_{i=1}^{25} m_{i} \varepsilon_{i} \]

\[ - 8050.751\left(N - 30.050\right)^{-1} \text{[kcal/mol]} \]  

(38)

where \( m_{i} \) stands for the total count of a specific face pattern (FP) that appears in the fullerene, \( \varepsilon_{i} \) is the FP energy contribution, and the second term in Eq (38) is the curvature term (\( C_{i} \)), which is repulsive for \( C_{20} \) and approaches zero for \( N \to \infty \). For example, for \( C_{60} - I_{b} \), we just have one pattern in Cioslowski et al.’s scheme consisting of a hexagon surrounded by 3 pentagons and 3 hexagons. There are 20 of these in \( C_{60} \), and with \( \varepsilon = 44.281 \text{ kcal/mol} \), we obtain \( \Delta H_{f} = 616.814 \text{ kcal/mol} \) for \( C_{60} \).\textsuperscript{33} This compares rather well with the estimated NIST listed heat of formation for \( C_{60} \) (612 ± 25 kcal/mol), although this value should probably be corrected downwards to a value of 602.7 kcal/mol as recently pointed out by Karton et al. using high-level theoretical procedures.\textsuperscript{185} The rms error for the standard enthalpies of formation for the 115 IPR isomers chosen was 4.0 kcal/mol compared to B3LYP/6-31G* calculations.

Alcami et al. devised a similar scheme for general fullerenes.\textsuperscript{32} By analyzing the most stable structures from B3LYP-DFT calculations of \( C_{20} \) to \( C_{72} \), and using only connections between four faces with varying number of pentagons with a total of 9 different motifs, an expression similar to Cioslowski et al. was used for \( E_{FP} \), but without the curvature term \( E_{C} \). This scheme gives \( \Delta H_{f} = 654.0 \text{ kcal/mol} \) for \( C_{60} - I_{b} \). For all fullerenes studied the rms error is 15.4 kcal/mol for the enthalpy of formation compared to DFT calculations, or 0.31 kcal/mol per carbon atom.

The question is: How well are topological indices suited to describe fullerene stability? Table 3 shows a comparison of stabilities for all \( C_{34} \) isomers obtained from DFT geometry optimizations (using the PBE functional\textsuperscript{186} and a def2-SVP basis set for carbon\textsuperscript{187}) in comparison with several topological indicators.\textsuperscript{35} According to the pentagon signature, isomer \( C_{34} - C_{3}(5) \) has the lowest \( P_{1} \) value, but with a very small HOMO-LUMO gap obtained from Hückel theory. This is indeed the case as the DFT results show. Alcami et al.’s stability indicator correctly predicts the most stable (\( P_{1} = 14 \)) and the least stable isomer (\( P_{1} = 17 \)). For the four isomers with pentagon signature \( P_{1} = 15 \), the two results (\( \Delta E_{DFT} \) and \( \Delta E_{A} \)) do not agree in their sequence (note that Alcami et al.’s scheme was not designed to distinguish between different isomers of a fixed vertex count). The different fullerene isomers are all very similar in surface area and volume. Concerning the HOMO-LUMO gap, the most stable isomer does not correspond to the one with the largest HOMO-LUMO gap. Moreover, these values are very sensitive to the method applied, and a better measure for the gap given by the singlet-triplet separation. The fullerenes with the smallest HOMO-LUMO gaps (isomers 2 and 6) prefer in fact the triplet ground state. The Hückel HOMO/LUMO gap on the other hand is only good for a qualitative discussion. We note that the two different fullerene isomers 3 and 5 have identical Wiener indices, and the isomer 6 with the smallest Wiener index is not the most stable isomer.

The first halma transform of \( C_{20} \) is \( C_{80} \) and therefore of \( I_{h} \) symmetry. There are seven IPR isomers out of a total of 31,924 isomers as possible candidates for the energetically most stable one. Table 4 shows...
the properties of all IPR isomers of C\textsubscript{80}. With larger fullerenes the band gap becomes very small (graphene is a semi-metal or zero-gap semiconductor\textsuperscript{189}), and one has to check for states of higher spin multiplicity. For C\textsubscript{80}, Huckel theory indicates an open-shell case that the triplet electronic state is taken instead of the singlet state. The results shown here clearly demonstrate that the topological indicators help enormously to sort out the most stable isomers. The situation is often quite complicated as seen from the DFT calculations here, or for example from the work of other authors who compared stabilities within a list of isomers.\textsuperscript{191−193} In the next section we introduce topological indicators that are connected to $\pi$-electron resonance structures (Kekulé structures) in fullerenes.

### TABLE 3 | Stability and Topological Indicators for the Six Isomers of C\textsubscript{34}

<table>
<thead>
<tr>
<th>$L_i$</th>
<th>PG</th>
<th>$P_1$</th>
<th>$\Delta E\text{DFT}$</th>
<th>$\Delta E^A$</th>
<th>$\Delta E^H_{HL}$</th>
<th>$\Delta E_{ST}$</th>
<th>$\Delta E\text{DFT}$</th>
<th>$\Delta E\text{DFT}$</th>
<th>$A$</th>
<th>$V$</th>
<th>$W$</th>
<th>$N_{HC}$</th>
<th>$N_{PM}$</th>
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<tr>
<td>1</td>
<td>C\textsubscript{2}</td>
<td>17</td>
<td>74.93</td>
<td>13.5</td>
<td>0.069</td>
<td>0.424</td>
<td>0.0267</td>
<td>81.41</td>
<td>60.30</td>
<td>1978</td>
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<td></td>
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<tr>
<td>2</td>
<td>C\textsubscript{3}</td>
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<td>28.10</td>
<td>4.2</td>
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<td>0.150</td>
<td>-0.153</td>
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<td>61.70</td>
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<td>58</td>
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<td></td>
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<td>3</td>
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<td>14</td>
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<td>-</td>
<td>-</td>
<td>-</td>
<td>81.32</td>
<td>61.69</td>
<td>1975</td>
<td>58</td>
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<td>62.80</td>
<td>1973</td>
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<td>0.2229</td>
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<td></td>
</tr>
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<td>0</td>
<td>0.225</td>
<td>0.0945</td>
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</tr>
</tbody>
</table>

$L_i$ is the numbering scheme according to the lexicographically ordered face spiral pentagon indices, PG the ideal point group, $P_1$ the pentagon signature, energy differences (in kcal/mol) $\Delta E\text{DFT}$ the DFT(PBE) energy difference to the most stable isomer (kcal/mol), $\Delta E^A$ the Alcami energy difference to the most stable isomer (kcal/mol), $\Delta E^H_{HL}$ the Alcami energy difference to the most stable isomer (kcal/mol), energy gaps (in eV) $\Delta \varepsilon_{HL}$ the Huckel HOMO-LUMO gap, $\Delta \varepsilon_{ST}$ the PBE HOMO-LUMO gap, $\Delta E\text{DFT}$ the singlet-triplet gap (in eV), $A$ the surface area ($\text{Å}^2$), $V$ the volume ($\text{Å}^3$), $W$ the Wiener index, $N_{HC}$ the number of Hamiltonian cycles, and $N_{PM}$ the number of perfect matchings. The symbol (T) indicates that the triplet electronic state is taken instead of the singlet state.

### TABLE 4 | Stability and Topological Indicators for the Seven IPR Isomers of C\textsubscript{80}

<table>
<thead>
<tr>
<th>$L_i$</th>
<th>PG</th>
<th>$H_2$</th>
<th>$\Delta E\text{DFT}$</th>
<th>$\Delta E^A$</th>
<th>$\Delta E^H_{HL}$</th>
<th>$\Delta E_{ST}$</th>
<th>$\Delta E\text{DFT}$</th>
<th>$\Delta E\text{DFT}$</th>
<th>$A$</th>
<th>$V$</th>
<th>$W$</th>
<th>$N_{HC}$</th>
<th>$N_{PM}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>D\textsubscript{5d}</td>
<td>500</td>
<td>0</td>
<td>4.34</td>
<td>2.12</td>
<td>0.220</td>
<td>0.355</td>
<td>0.168</td>
<td>203.70</td>
<td>255.16</td>
<td>17340</td>
<td>10450</td>
<td>270153</td>
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<tr>
<td>2</td>
<td>D\textsubscript{2}</td>
<td>496</td>
<td>2.63</td>
<td>3.0</td>
<td>0.75</td>
<td>0.528</td>
<td>0.460</td>
<td>0.293</td>
<td>203.85</td>
<td>257.72</td>
<td>17352</td>
<td>10642</td>
<td>237585</td>
</tr>
<tr>
<td>3</td>
<td>C\textsubscript{2v}</td>
<td>488</td>
<td>1.81</td>
<td>10.5</td>
<td>0.04</td>
<td>14.7</td>
<td>0.102</td>
<td>0.091</td>
<td>203.92</td>
<td>260.80</td>
<td>17412</td>
<td>9918</td>
<td>201623</td>
</tr>
<tr>
<td>4</td>
<td>D\textsubscript{3}</td>
<td>492</td>
<td>7.03</td>
<td>7.5</td>
<td>7.17</td>
<td>6.70</td>
<td>0.408</td>
<td>0.098</td>
<td>0.036</td>
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<tr>
<td>5</td>
<td>C\textsubscript{2v}</td>
<td>484</td>
<td>6.33</td>
<td>16.5</td>
<td>2.92</td>
<td>24.7</td>
<td>0.298</td>
<td>0.038</td>
<td>0.090</td>
<td>203.99</td>
<td>262.48</td>
<td>17454</td>
<td>9004</td>
</tr>
<tr>
<td>5(T)</td>
<td>C\textsubscript{2v}</td>
<td>484</td>
<td>4.25</td>
<td>16.5</td>
<td>2.92</td>
<td>24.7</td>
<td>0.298</td>
<td>0.038</td>
<td>0.090</td>
<td>203.99</td>
<td>262.48</td>
<td>17454</td>
<td>9004</td>
</tr>
<tr>
<td>6</td>
<td>D\textsubscript{5h}</td>
<td>480</td>
<td>6.81</td>
<td>22.5</td>
<td>0</td>
<td>37.7</td>
<td>0</td>
<td>0.081</td>
<td>-0.002</td>
<td>204.01</td>
<td>263.50</td>
<td>17500</td>
<td>10970</td>
</tr>
<tr>
<td>6(T)</td>
<td>D\textsubscript{5h}</td>
<td>480</td>
<td>6.38</td>
<td>22.5</td>
<td>0</td>
<td>37.7</td>
<td>0</td>
<td>0.081</td>
<td>-0.002</td>
<td>204.01</td>
<td>263.50</td>
<td>17500</td>
<td>10970</td>
</tr>
<tr>
<td>7</td>
<td>I\textsubscript{h}</td>
<td>480</td>
<td>17.49</td>
<td>30.0</td>
<td>14.2</td>
<td>63.4</td>
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<td>0.115</td>
<td>-0.003</td>
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<td>263.61</td>
<td>17600</td>
<td>10500</td>
</tr>
<tr>
<td>7(T)</td>
<td>I\textsubscript{h}</td>
<td>480</td>
<td>17.43</td>
<td>30.0</td>
<td>14.2</td>
<td>63.4</td>
<td>0</td>
<td>0.115</td>
<td>-0.003</td>
<td>204.02</td>
<td>263.61</td>
<td>17600</td>
<td>10500</td>
</tr>
</tbody>
</table>

$L_i$ is the numbering scheme according to the lexicographically ordered IPR face spiral pentagon indices, PG the ideal point group, $H_2$ the second moment hexagon signature, energy differences (in kcal/mol) $\Delta E\text{DFT}$ the DFT(PBE) energy difference to the most stable isomer (kcal/mol), $\Delta E^A$ the Alcami energy difference to the most stable isomer (kcal/mol), $\Delta E^H_{HL}$ the Alcami energy difference to the most stable isomer (kcal/mol), energy gaps (in eV) $\Delta \varepsilon_{HL}$ the Huckel HOMO-LUMO gap, $\Delta \varepsilon_{ST}$ the PBE HOMO-LUMO gap, $\Delta E\text{DFT}$ the singlet-triplet gap (in eV), $A$ the surface area ($\text{Å}^2$), $V$ the volume ($\text{Å}^3$), $W$ the Wiener index, $N_{HC}$ the number of Hamiltonian cycles, and $N_{PM}$ the number of perfect matchings. The symbol (T) indicates that the triplet electronic state is taken instead of the singlet state.
Kekulé Structures and Perfect Matchings, Clar and Fries Structures, and the Pauling Bond Order

Every chemist learns how to draw double bonds into an aromatic system called a Kekulé (or resonance) structure, sometimes also called a benzenoid graph. 194 For benzene there are only two possible Kekulé structures with three double bonds as we all know, but for C_{60} there are as many as 12,500 Kekulé structures with 30 double bonds (a fullerene has exactly \( n_{\text{DB}} = \frac{N}{2} \) double bonds), of which only 158 are symmetry distinct, that is, non-isomorphic. 195–197 A Kekulé structure is the same as a perfect matching in the chemical graph \( G = (\mathbb{V}, \mathbb{E}) \), where the edges of the matching correspond to double bonds. We write \( \mathcal{K}(G) \) for the set of all perfect matchings of the graph, and the number of different Kekulé structures \( K = |\mathcal{K}(G)| \) is called the Kekulé number. In graph theory, a perfect matching is a selection of edges such that every vertex of the graph \( G \) is part of exactly one edge in the matching. The edges of the matching correspond to the double bonds.

Chemists know from basic Hückel theory that for two different benzenoid graphs \( B_1 \) and \( B_2 \), \( K(B_1) > K(B_2) \) implies that \( B_1 \) is more stable than \( B_2 \) because of resonance stabilization. Schmalz et al. calculated the Kekulé number for several small fullerenes up to C_{84}. 50 Their work showed that for C_{60}, the least stable isomer C_{60}-D_{5h}(1) has \( K = 16,501 \), while for the most stable isomer C_{60}-I_6(1812) we have \( K = 12,500.50 \) In fact, Austin et al. showed that 20 isomers of C_{60} have a higher Kekulé numbers than C_{60}-I_6. 198 The Kekulé number is therefore not a good indicator for fullerene stability, but useful for rationalizing the different bond lengths in fullerenes as we shall see.

All cubic graphs have exponentially many perfect matchings and hence exponentially many Kekulé structures. 199 While the theoretical lower bounds proved in Refs 113, 199 have small exponents, the actual exponential behavior kicks in rapidly, as seen in Figure 17: The mean number of perfect matchings for a C_{N} isomer is approximately \( 20 \cdot 2^{(N-20)/4} \). In general, calculating the number of perfect matchings for graphs is intractable, but for planar graphs (such as fullerenes) the Kekulé number can be computed in \( \mathcal{O}(N^3) \) time using the Pfaffian matrix (the Fisher-Kasteleyn-Temperley algorithm). 200

Pauling, Brockway, and Beach introduced bond orders derived from Kekulé structures in order to rationalize the differences in the C–C bond lengths encountered for polycyclic aromatic compounds. 201 The Pauling bond order (PBO) \( P_{ij} \) of an edge \( v_i - v_j \) is defined as the sum over all appearances of double bonds in perfects matchings

\[
P_{ij} = \frac{1}{K} \sum_{M \in \mathcal{K}(G)} \beta_{ij}
\]  

(39)

where \( \beta_{ij} = 1 \) if the edge \( v_i - v_j \) has a double bond in the perfect matching \( M \), otherwise \( \beta_{ij} = 0 \). This gives \( 0 \leq P_{ij} \leq 1 \), with 0 being a pure C–C single bond and 1...
a pure C=C double bond. In C_{60}-I_h there are two different type of bonds and therefore two different PBOs, depending if the edge shares two hexagons ($P_{hh} = 11/25 = 0.44$) or a hexagon and a pentagon ($P_{hp} = 7/25 = 0.28$). Note that the Hückel bond orders derived from the eigenvectors of the adjacency matrix are generally larger with $P_{hh} = 0.6010$ and $P_{hp} = 0.4758$ for C_{60}-I_h.\(^{203}\) Narita et al. derived a linear relationship between the PBO and the bond distance in a fullerene obtained from X-ray diffraction,\(^{202}\)

\[ r_{ij} [\text{Å}] = 1.554 - 0.399P_{ij} \quad (40) \]

Figure 18 shows an AFM picture of one of the symmetry equivalent hexagons in C_{60} clearly showing the two different types of bonds. The measured bond lengths are $r_{hh} = 1.38(2)$ Å and $r_{hp} = 1.454(12)$ Å,\(^{204}\) in excellent agreement with the values obtained from Eq (40) ($r_{hh} = 1.378$ Å and $r_{hp} = 1.442$ Å).

Kekulé structures containing benzenoid moieties are of special interest. These are the Fries and Clar structures. Fries structures are Kekulé structures with maximum number of benzenoid hexagons. The Fries number $\text{Fries}(G)$ of a benzenoid graph is the maximum number of benzenoid hexagons over all of its Kekulé structures, and a Fries structure is a perfect matching that realizes this maximum (there could be many). Similarly, the Clar number $\text{Clar}(G)$ is the largest number of independent sets of benzenoid faces (separated from each other) over all Kekulé structures, and a Clar structure is a perfect matching that realizes this (there could be many). It is clear that a Clar structure localizes benzenoid structures in fullerenes. For C_{60}-I_h we have $\text{Fries}(G) = 20$ and $\text{Clar}(G) = 8$, and an example for a Fries and Clar structure of C_{60}-I_h is shown in Figure 19. One might naively assume that a set of Clar structures form a subset of Fries structures, but this is generally not the case for fullerenes.\(^{205}\)

We expect maximum stability for fullerenes with the highest Clar number. For example, C_{60}-I_h is unique among all other isomers in that it has a Fries structure where all hexagons contain three double bonds and all pentagons none. This is also seen as a reason for the unique stability of C_{60}. However, finding the Clar number is not a trivial problem as it is computationally NP-hard.\(^{206}\) Ye and Zhang recently showed that for C_{60} there are exactly 18 fullerenes with maximal Clar number of 8.\(^{207}\) Some of them contain as many as 6 fused pentagons. Hence the Clar number alone is not a good measure for stability. However, out of the 18 fullerenes with Clar number 8, C_{60} has the largest Kekulé count.\(^{208}\) Fowler showed that leapfrog transforms of fullerenes not only are closed shell, but also have the maximum proportion of benzenoid hexagons.\(^{209}\)

Figure 17(c) shows that the expected number of perfect matchings is no different for IPR fullerenes than it is for all fullerenes, except for small fullerenes for which it is larger than average. However, the maximum number is always significantly lower for IPR fullerenes. One could assume that the larger the Kekulé number is the more Clar sextets can be found and the more stable the fullerene is. While this has not been explored in detail, Table 4 shows that for C_{80} the most stable D_{5h} isomer has the lowest perfect matching count. There are, however, some good lower and upper bounds of these topological indices known. Došlić showed that the number of Kekulé structures grows exponentially with a lower bound of $2^{N/8}$.\(^{210}\)
This is not a very tight bound, however, as for C\textsubscript{60} it predicts at least 181 Kekulé structures, and there are as many as 12,500. The upper bound for Fries numbers is well known, \(Fries(G) \leq N/3\), which is seen to be the graphenic limit. Fries structures with a maximum \(N/3\) Fries number are called \textit{complete Fries structures}\textsuperscript{211} or \textit{perfect Clar structures}. This happens exactly for Clar structures that use every vertex. In that case, the Clar and the Fries structure is the same, and corresponds to a ‘soccer ball coloring’ of the faces. An interesting result comes from Pisanski who pointed out that the reduced \(p\)

An upper bound for Clar numbers has been obtained by Zhang and Ye, \(\text{Clar}(G) \leq (N/6 – 2)\).\textsuperscript{213} Both IPR fullerenes, \(C\textsubscript{60}\cdot I\textsubscript{h}\), and \(C\textsubscript{70}\cdot D\textsubscript{5h}\) with \(\text{Clar}(C\textsubscript{70}) = 9\) achieve this upper bound. Such fullerenes are called \textit{extremal fullerenes}.\textsuperscript{213}

Finally we mention that the Kekulé structures can be used to determine the resonance stability of fullerenes. To start with a very simple model, Randić et al. took the Kekulé structures of fullerenes ranging from \(C\textsubscript{20}\) to \(C\textsubscript{72}\), and calculated the \(\pi\)-\textit{content} \(P\pi\) of hexagons and pentagons, which is obtained by summing all the Pauling bond orders in a specific ring.\textsuperscript{197} For \(C\textsubscript{60}\cdot I\textsubscript{h}\) this gives \(P\pi = 3P\pi\textsubscript{hp} + 3P\pi\textsubscript{hp} = 54/25 = 2.16\) for a hexagon, and \(P\pi = 5P\pi\textsubscript{hp} = 35/25 = 1.40\). High hexagon \(\pi\)-count together with low pentagon \(\pi\)-count gives stability to a fullerene. High hexagon \(\pi\)-count increases the aromaticity in the system (for more details see Ref 197). Klein et al. applied Herndon’s resonance theory\textsuperscript{214,215} using counts of \(2m\)-resonance cycles to obtain the resonance energy.\textsuperscript{195} They correctly pointed out that the reduced \(p\pi\)-orbital overlap in strongly curved fullerenes will reduce the resonance energy and has to be taken into account.\textsuperscript{195} Here we should mention again that obtaining all Kekulé structures for a fullerene is an exponential problem, and it is more advantageous to apply methods that scale polynomially in time, such as Hückel or more sophisticated (semi-empirical) theories.

**Hamiltonian Cycles, IUPAC Rules, and Naming Fullerenes**

For naming an alkene, the IUPAC rules state that one has to find the longest continuing carbon chain containing as many double bonds as possible. For a cyclic system one chooses the longest cyclic chain, and if there are multiple longest cycles, one must choose the one that maximizes the number of double bonds along the chain. This is related to finding a \textit{Hamilton cycle}, which is a closed path in a graph that visits every vertex exactly once. If at least one such cycle exists, we say that the graph is \textit{Hamiltonian}. In this case, the Hamilton cycles are the longest carbon chains, and the optimal one must be chosen among these.

It is an open problem whether every fullerene has a Hamilton cycle.\textsuperscript{1} In practice, however, we find that fullerenes are not only Hamiltonian, but admit exponentially many Hamilton cycles. The mean number of Hamilton cycles taken over all \(C\textsubscript{N}\)-isomers grows approximately as \(8 \cdot 2^{(N/20)^{0.5}}\) for \(N\mod 4 = 0\) and as \(12 \cdot 2^{(N/20)^{0.5}}\) for \(N\mod 4 = 2\) (Schwerdtfeger et al., unpublished manuscript). Because of this, naming fullerenes according to the IUPAC alkene rules is a computationally heavy task: we must search through all the exponentially many Hamiltonian cycles (main rings) to find the one in which the secondary bridges are labeled in the lexicographically smallest way. This is computationally feasible only for small fullerenes such as the ones shown in Figure 20.

In addition to being infeasible to calculate, as \(N\) grows, the name resulting for a fullerene from the IUPAC alkene rules rapidly becomes long and unmanageable. Even for a relatively small fullerene such as \(C\textsubscript{60}\cdot I\textsubscript{h}\), the name\textsuperscript{8} is already unreadable, and the name length keeps growing with the fullerene size.\textsuperscript{216} The IUPAC alkene name for fullerenes is now seen mostly as a curiosity—and a good example of how a well meant rule has is limits.

IUPAC employs an alternative naming scheme for fullerenes: \(C\textsubscript{N}\cdot PG[5,6]\)fullerene, where PG stands for the (ideal) point group, and [5, 6] denotes a polyhedral cage with only pentagon and hexagon faces. For example, the IPR \(C\textsubscript{60}\) fullerene has the official IUPAC name \(C\textsubscript{60}\cdot I\textsubscript{h}[5,6]\)fullerene. While computing the name of a fullerene according to this scheme is easy and efficient, and contrary to the alkene naming scheme is short and easy to understand, we find that this scheme is not much better than the IUPAC alkene nomenclature. On the one hand it is unnecessarily long: ‘fullerene’ implies ‘[5, 6]’, and vice versa. On the other hand, it does not uniquely specify a fullerene, since many isomers share the same vertex count and symmetry. For example, while \(C\textsubscript{120}\cdot T\textsubscript{d}[5,6]\)fullerene happens to be unique, \(C\textsubscript{120}\cdot C\textsubscript{2}[5,6]\)fullerene refers to 10,787 different fullerene isomers, and \(C\textsubscript{120}\cdot C\textsubscript{1}[5,6]\)fullerene refers to no less than 1,660,007 different molecules. We therefore advocate the use of one of the following two names to uniquely specify fullerenes, based
on the (generalized) canonical face spiral pentagon indices (FSPI), see the chapter on face spirals above. Both schemes uniquely and compactly identify a specific fullerene molecule.

1. For moderate values of $N$, we can specify the fullerene by its canonical index, which is the lexicographic number of its FSPI among the isomers. For example, the IPR $C_{60}$ molecule can be written as $C_{60}^{-Ih}(1812)$ or $C_{60}^{-Ih}(IPR - 1)$, as it is the last fullerene of the 1812 $C_{60}$ isomers in the lexicographically ordered FSPI list.

2. For large $C_N$, for which it is not feasible to compile databases for all isomers, we annotate instead with the FSPI. For the vast majority of fullerenes, this is a list of 12 small integers. For the extremely rare cases where jumps are necessary (there are only two non-spiral fullerenes out of the $2.6 \times 10^{12}$ fullerenes up to $C_{400}$), two extra integers are required per jump. No currently known case requires more than one jump. With this scheme, $C_{60}^{-Ih}(IPR - 1)$ is written $C_{60}^{-Ih}(1, 7, 9, 11, 13, 15, 18, 20, 22, 24, 26, 32)$, and the unspirable $C_{380}$ is written $C_{380}^{-T}(110, 2; 45, 70, 71, 82, 83, 110, 119, 120, 144, 184, 185, 192)$. In the latter example, the two numbers before the semicolon denote a cyclic shift of length 2 before adding face number 110.

The canonical FSPI notation is especially advantageous for more reasons than being compact and complete without needing to refer to a precomputed database. The procedure to construct a fullerene graph from the FSPI (the ‘windup’ operation) is geometrically intuitive—visualized like peeling an orange—and the algorithm is so simple that it can be performed with pen and paper. Thus, one can even reconstruct fullerenes of moderate sizes from the FSPI by hand, without the help of a computer.

The algorithm for the windup operation is $\Theta(N)$, and the inverse operation, unwind, takes $\Theta(N)$ expected time to find a single generalized spiral, and $\Theta(N^2)$ to find the canonical one. The canonical FSPI constitutes a canonical labeling of fullerene graphs up to isomorphism. Hence one can check whether two fullerene graphs are isomorphic simply by testing whether they have the same canonical FSPI, and the canonical FSPI gives us a unique graph representative of each isomorphism class via the windup procedure.

As discussed later, the FSPI representation makes it easy to directly compute the ideal symmetry group of the fullerene. Because the generalized face spiral algorithm is complete for all connected planar cubic graphs, a compressed form of the face spiral similar to the FSPI can be used in general for fulleroids, to be introduced in the last chapter.

### Thermodynamic Stability and the Graphene Limit

The spherical shape of $C_{60}^{-Ih}$ with no adjacent pentagons is seen as the main reason for its unusual stability, which underlines the importance of Kroto’s isolated pentagon rule (IPR). For example, Nagase and co-workers investigated the stability of potential candidates for the lowest energy structure of $C_{74}$, with the sole IPR $D_{36}$ isomer being 16 kcal/mol lower in energy than the $C_2$ isomer containing two fused pentagons. It is therefore convenient to compare the stability of a fullerene to
C_{60}-I_h (relative fullerene stability, RFS) by using the isodesmic reaction C_{60}/60 \rightarrow C_N/N,

$$\Delta E^{RFS}(N) = \frac{1}{N}E[C_N] - \frac{1}{60}E[C_{60} - I_h]$$ (41)

where $E[C_N]$ is the total energy for a $C_N$ fullerene. For $N \rightarrow \infty$ we obtain the graphene limit, which can be estimated to be approximately $\Delta E^{RFS}(\infty) = -9$ kcal/mol.

As $E \sim N$, $E/N$ approaches a constant value for infinite systems, for example, the graphene limit if the pentagons are as separated as possible from each other, so that the polyhedral surface is mainly constituted of flat graphene sheets. Alternatively, for nanotube fullerenes with pentagon caps, the infinite particle limit is just the corresponding infinite nanotube-fullerene. For the graphene limit, the next dominant term in a $1/N$ expansion will be the surface tension or curvature term, which is $N^{-1}$ as for example used by Cioslowski et al.,$^{33}$ see Eq (38) (although a $N^{-1/2}$ law has been also suggested before$^{59}$). Indeed, as Figure 21 shows, the stability follows approximately a linear behavior in our $1/N$ fit (we set $\Delta E^{RFS} = 0$ for $N = 60$),

$$\Delta E^{RFS}(N) = 527.4N^{-1} - 8.79 \text{ (kcal/mol)}$$ (42)

that is fullerenes become more energetically stable with increasing vertex number $N$. It was already shown experimentally that $C_{70}$ is more electronically stable than $C_{60}$. Alcami et al.’s scheme predicts the correct graphene limit ($N \rightarrow \infty$), and Cioslowski’s scheme performs well for medium sized IPR fullerenes, but has an incorrect convergence behavior toward the graphene limit.$^{32,33}$ $C_{20}$ is the smallest member of the fullerene family, and as seen from Figure 21 also the least electronically stable one. It has been detected in 2000 by Prinzbach$^{16}$ (see also Ref 220), and a current review on the state of affair concerning $C_{20}$ is given by Fei et al.$^{221}$

It becomes more and more difficult to extract the most stable isomers from the huge isomer space as the size of the fullerene increases. Besides using topological indicators, Hückel theory can be used to determine the resonance stability of a fullerene. It requires diagonalization of the adjacency matrix $A_{ij}$ (the Hückel matrix is defined as $H_{ij} = a\delta_{ij} + \beta A_{ij}$, where $a$ is the Coulomb and $\beta$ the resonance integral, and $\delta_{ij}$ is the unit matrix), and the occupied Hückel orbitals determine the Hückel resonance stability (HRS) of a fullerene (in units of $\beta$),

$$\Delta E^{HRS}(N) = E_\pi - N$$

with $E_\pi = \sum_{i=1}^{occ} n_i\varepsilon_i \quad (\varepsilon_i \in [-3,+3])$ (43)

where $n_i = 0,1,2$ are the occupation numbers, $\varepsilon_i$ the orbital energy for Hückel orbital $i$, and the last term comes from subtracting the non-resonant ethene value (single double bond). This gives $E_{HRS} = 33.161\beta$ for $C_{60}$. Per $\pi$-electron this gives a resonance energy of $0.5527\beta$ much higher compared to benzene with exactly $\beta/3$. This should indicate that $C_{60}$ is more aromatic than benzene, which has been a matter of some debate in the past (see the review by Bühl and Hirsch$^{223}$). The problem is that the $\pi$-overlap in non-planar systems is not ideal and there is substantial mixing (hybridization) with the $p_e$ orbitals of carbon, thus questioning the $\sigma$-$\pi$-separability in Hückel theory. Hence there should be a correction accounting for the curvature at each carbon atom. Such curvature corrections, for example through using the $\pi$-orbital axis vector method (POAV), have been considered before.

**FIGURE 21** The stability of fullerenes $C_N$ in comparison to $C_{60}$ obtained from density functional calculations up to the graphene limit ($N \rightarrow \infty$). Topological stability indices from resonance energies$^{222}$ (TRE using $\beta = -216 \text{ kcal/mol}$) or ring patterns by Alcami$^{32}$ and Cioslowski$^{33}$ are also shown. The graphene limit is estimated from the heat of formation of $C_{60}$.$^{185}$
with some success, and the different approaches are discussed in detail by Bakowies and Thiel. For example, Haddon et al. showed that C_{240} is significantly more stable than C_{60}, which is indeed the case as Figure 21 shows. Clearly, as the fullerene system becomes larger, the curvature term becomes smaller. Moreover, for similar shaped fullerenes the Hückel resonance energy might still be a good approximation. Indeed, as Table 4 for C_{60} shows, the first two isomers are the most stable ones predicted by the simple Hückel resonance energy term in agreement with DFT calculations, and isomer 7 is the least stable one. The Hückel method only requires a diagonalization of the adjacency matrix, and for obtaining bond orders and resonance energies it is computationally more efficient than sorting out Kekulé structures. We mention that the Hückel spectrum \( \{ \varepsilon_i \} \) for fullerenes shows some interesting mathematical properties. The largest eigenvalue (which translates into the lowest occupied level as the resonance integral \( \beta \) is negative) in a fullerene is always \( \varepsilon_{\text{max}} = +3 \). The smallest eigenvalue (highest unoccupied level) is largest for C_{20} with \( \varepsilon_{\text{min}} = -\sqrt{5} \), and for the IPR isomers C_{60} we have \( \varepsilon_{\text{min}} = -\left( 1 + \sqrt{5} \right)^2 /4 \). As a geometric consequence, the famous golden ratio \( (1 + \sqrt{5}) /2 \) appears here as it does in the volume and surface area calculations discussed above. We also mention that in general, different graphs can have the same Hückel spectrum, and this is the case also for fullerenes: the eigenvalues do not uniquely determine the fullerene graph.

It is well known that the Hückel scheme for canonical resonance structures is of rather limited use, and Aihara and Gutman et al. introduced a different reference system by defining the topological resonance energy (TRE) for a graph \( G \) as

\[
\Delta E^{\text{TRE}} (G) = E^\pi (G) - E^{\text{MP}} (G)
\]

where the reference energy \( E^{\text{MP}} (G) \) is the sum of all roots given by the matching polynomial, which for a fullerene is

\[
P^{\text{MP}} (G) = \sum_{k=0}^{N/2} (-1)^k c (G, k) x^{N-2k} \quad (c^{\text{MP}} (G, k) \in \mathbb{Z})
\]

Here the polynomial coefficients \( c(G, k) \) define the number of ways of choosing \( k \) non-adjacent edges from the graph \( G \). Balasubramanian gave polynomial coefficients \( c(G, k) \) for a number of fullerenes up to C_{50}, Babič determined the coefficients for C_{70} and Salvador for C_{60} and C_{70} to C_{100}. The first few polynomial coefficients are known as they are independent of the isomers for a specific vertex count, that is, \( c(G, 0) = 1 \), \( c(G, 1) = -3N/2 \), \( c(G, 2) = 3N(3N - 10)/8 \), \( c(G, 3) = -(9N^3 - 90N^2 + 232N)/16 \). The last coefficient \( c(G, N/2) \) is just the number of perfect matchings in the graph. The computation of these coefficients soon becomes computationally intractable. Babič et al. found, however, a good correlation between \( E^{\text{MP}} \) and \( E^\pi \) values for fullerenes, and the \( E^\pi \) value suffices to calculate the topological resonance energy approximately as (in units of \( \beta \)),

\[
\Delta E^{\text{TRE}} = 1.024296 E^\pi - 1.512148N
\]

In order to give reasonable results for the isodesmic reaction energy (Eq 41), a rather low value of \( \beta = -216 \) kcal/mol has to be applied. This scaled TRE plot is shown in Figure 21, which is in reasonable agreement with the results obtained from DFT calculations.

The five or six Pauling bond orders in a pentagon or hexagon respectively can be added to give the Pauling ring bond order. For C_{60}, this gives 1.40 (5 \times 11/25) for a pentagon and 2.16 (3 \times 11/25 + 3 \times 7/25) for a hexagon compared to 3.00 for benzene. According to Randić this suggests only moderate aromaticity for C_{60}. We are not going into the controversy of aromaticity of fullerenes, but refer to a recent paper by Schleyer and co-workers, who find that both C_{60} and C_{60} are not spherically \( \pi \) aromatic but spherically \( \pi \) anti-aromatic, which for C_{60} explains the large heat of formation. In this respect it is interesting that Schleyer and co-workers conclude that fullerenes are not highly stable molecules, which perhaps is in line with the data shown in Figure 21 showing no ‘magic’ stability for C_{60} compared to the other fullerenes. We finally mention that fullerene cage abundance is not only guided by thermodynamics, but mostly by kinetic stability. For example, the relative isomer abundance of fullerenes and carbon nanotubes correlates well with kinetic stability.

**Electronic Aspects to Structure and Stability**

So far we have discussed topological aspects to determine the structure and stability of fullerenes and basic Hückel theory gives us a first insight into the electronic structure. For example, Fowler and co-workers have shown that leapfrog fullerenes adopt a closed-shell structure, and have equal numbers of positive and negative eigenvalues.
For a more detailed discussion on this subject see Ref. 243. However, as we have already seen, more detailed quantum chemical calculations are required to describe the bonding in fullerenes accurately (see also the work by Thiel and co-workers244–246). For example, the IPR isomer C_{80}-C_{2v}(5) shown in Table 4 prefers a triplet ground state (at the DFT-PBE level of theory) contrary to Hückel theory, which predicts a singlet state. This is perhaps not surprising as already Hückel theory predicts a small HOMO/LUMO gap. As the band gap closes further with an increasing number of vertices (see for example the recent paper by Noël et al.247), we cannot expect anymore to predict the correct spin ground state from simple topological arguments. Moreover, the system becomes more multi-reference in nature, making even a quantum theoretical treatment difficult. As an example, C_{50}-D_{5h}(271) was investigated by Lu et al. who showed that two singlet states with different HOMO-LUMO symmetries are quasi-degenerate.248

In addition, some fullerenes may undergo first- or second-order Jahn-Teller (JT) distortions (which could, however, be very small and almost undetectable for larger fullerenes), that is, they can distort to subgroups of the ideal point group symmetry given by the fullerene topology.249,250 For example, C_{20} is well known to distort away from the ideal I_{h} symmetry, which has led to much discussion and debate over the correct electronic ground state.251 The JT distortion in C_{20} is a consequence of two electrons being distributed over four energetically degenerate g_{u}-orbitals, resulting in a rather complicated topology of the JT energy hypersurface.248,250,252–254 JT distortions for highly symmetric open- or closed-shell fullerenes may require a multi-reference treatment. It is therefore often difficult to predict the correct electronic ground state and corresponding physical point group symmetry of a fullerene.

The Gas Phase Formation of Fullerenes

Since the discovery of fullerenes almost 30 years ago, there has been considerable activity, both from the experimental and theoretical side, to gain a detailed understanding of the fullerene formation in the gas phase. However, the formation mechanism and especially the high yield of C_{60}-I_{h} and C_{70}-D_{5h} remains elusive and somewhat controversial.

Fullerenes can be produced by (a) evaporating a carbon target (graphite, amorphous carbon, fullerenes), optionally with addition of metal oxides with a laser,5 (b) an electric arc between carbon electrodes,9 or (c) by partial combustion of carbon rich organic compounds.255 Each of these methods can be adjusted by several experimental parameters (gas pressure, carrier gas, rod feeding rate, amount of oxygen, etc., and therefore indirectly the carbon vapor concentration, expansion rate, annealing time, etc.). Furthermore, fullerenes are found in space,14 at meteor impact sites, after lightnings and bush-fires, and soot from household candles.99 These conditions are similar in that carbon vapor is formed at very high temperatures with a deficit of possible reactants like oxygen or hydrogen. The distribution of yielded fullerene cage sizes depends on the production method and the above-mentioned experimental parameters; however, C_{60}-I_{h} and C_{70}-D_{5h} are always among the most abundant species.

A large number of formation mechanisms have been proposed.99 First, inspired by their resemblance to graphene sheets or nanotubes, it was suggested that graphene curls or nanotubes break apart to form fullerene cages, followed by various bottom-up strategies that suggest the successive addition of faces (party line, pentagon road, or polyyne rings (ring stacking) before the cage is closed, the folding of chains to form a cage (ring fusion zipper), or the growth of already existing cages (fullerene road, closed network growth), and many others. The currently most widely accepted mechanism comes from Irle, Zheng, Wang, and Morokuma, and is called the ‘shrinking hot giant road’.99 It is based on the concept of self-organizing structures under non-equilibrium conditions, backed by extensive QM/MD simulations.

This mechanism can be divided into five phases: First, linear polyyne chains and cycles form. In the second stage (nucleation), entangled carbon chains rehybridize and form faces. It is to be noted, that pentagons and hexagons are close in energy at the given temperatures,99 however, smaller and larger faces are formed as well. Third, more carbon dimers attach to the side chains of an existing nucleus (growth), allowing for the formation of additional faces. Fourth, after the formation of sufficiently many faces of sizes <6, the cage may close spontaneously (cage closure), leaving a carbon cage with face sizes not restricted to 5 and 6 and with polyyne side chains attached to it. The final step is the ejection of carbon dimers off side chains and the cage in combination with rapid isomerization of the cage structure resulting in a fullerene without side chains and faces of sizes 5 and 6 only. Fullerenes can not only shrink but also grow in steps of C_{2},5,98,262 and the existence of C_{2} and C_{3} fragments is backed by spectroscopy.263 The addition and ejection of small carbon fragments (especially C_{2}) to and from forming fullerene cages is a fast equilibrium. While the non-cage carbon concentration is high addition prevails; as the carbon vapor
expands ejection begins to dominate. The first four steps are exothermic, while the last step is endothermic but increases the overall entropy.

As shown in Figure 21, large fullerenes are more electronically stable than small ones, with graphene having a lower energy than any fullerene. In an equilibrium one would therefore anticipate the formation of graphene—the formation of strained cages and especially the high yield of C_{60}^{-1} I_b and C_{70}^{-1} D_{5h} require further explanation. The experimental conditions of the cooling and expanding carbon vapor are, however, far from an equilibrium, and the whole formation must be understood as a process of self-organization that is governed by kinetics more so than thermodynamics. Once fullerenes have formed they are subject to restricted equilibration only. Curl et al. have shown numerically that exchange of carbon dimers between existing cages is sufficient to explain the high yield of C_{60}^{-1} I_b and C_{70}^{-1} D_{5h} as well as a broad distribution of larger cages. They show, that the driving forces for the C_2 exchange are the energy difference between C_N and C_{N+2} and the high stability of C_{60}^{-1} I_b and C_{70}^{-1} D_{5h} relative to their isomers rather than entropy; even at 4000 K entropic effects are not strong enough to explain the existence of small cages.

The experimental observation of endohedral metallofullerenes supports a top-down formation mechanism in two ways: As any enclosed fragments need to enter the cage before it is closed, fullerenes containing fragments that use up most of the space available in the carbon cage must either be formed top down or their existence implies the breaking and reforming of carbon bonds to open and close the cage after it was formed. Secondly, enclosed metal fragments may stabilize otherwise unstable fullerenes: Zhang et al. report the finding of non-IPR M$_1$C$_2$@C$_{84}$(51383), which they interpret to be a missing link in the top-down road, and which would have ejected further C$_2$ fragments without the stabilizing metal carbide.

### How Much Space Is Inside a Fullerene?

**Endohedral Fullerenes and Buckyonions**

We consider only topological aspects here as there are several reviews on the chemistry of endohedral fullerenes available. We only note that endohedral enclosure can change the stability between different isomers. For example, Tb$_3$N@C$_{84}$ has been isolated and shows one fused pentagon pair. Rodriguez-Fortea et al. argued that if the enclosed metal atom or cluster donates electrons to the fullerene cage, the negative charge resides mostly at the pentagons. As this creates a Coulomb repulsion between the pentagons, one can introduce the inverse pentagon separation index (IPS),

$$\text{IPSI} = \sum_{i=1}^{12} \sum_{j>i}^{12} R_{ij}^{-1}$$

with $R_{ij}$ being the Euclidian distance between the pentagons (although the topological distance might be used instead). Small IPSI-values are preferred as these correspond to a small Coulomb repulsion.

It is clear that the cavity in the fullerene cage should be large enough to encapsulate atoms, molecules, or even smaller fullerenes. The size of the cavity can be estimated from fitting shapes such as spheres or ellipsoids inside a fullerene cage as discussed above.

Conversely to the minimum covering sphere problem (MCS), we wish to find the largest sphere that is fully contained within the polyhedron. This is called the maximal inscribed sphere (MIS), or simply inscribed sphere, and its radius for a given polyhedron $P$ is

$$R_{\text{MIS}} = \max_{\mathbf{x} \in P} \min_{\mathbf{c} \in \text{MIS}} ||\mathbf{x} - \mathbf{c}||$$

where $\mathbf{c} \in \text{MIS}$ is constrained to the points interior to $P$. In the case of convex polyhedra, the MIS is unique, and can be computed directly by finding the Chebyshev centre of the polyhedron. The inscribed radius is then determined by a linear programming problem. However, in the non-convex case, the MIS is no longer unique: Consider two overlapping circles in 2D space with the points lying on these circles. We now have two equivalent possibilities for placing our inner circle. The same argument holds for peanut-shaped fulleroids, which are introduced in the last chapter. Hence, in the non-convex case, we only search for one of the possibly many largest spheres contained entirely inside the polyhedral, giving us a numerical optimization problem.

Once the MIS has been obtained, we can estimate whether an atom or molecule fits inside the fullerene without coming close to the repulsive wall of the fullerene cage. For example, the space available inside C_{60}^{-1} I_b is roughly $R_{\text{MIS}}(C_{60}) - R_{\text{VdW}}(C) \approx 3.550 - 1.415 \approx 2.135 \AA$. If we compare this to the Van der Waals radii for the rare gas atoms, which are 1.40 Å for He, 1.54 Å for Ne, 1.88 Å for Ar, 2.02 Å for Kr, and 2.16 Å for Xe, we see that they all fit into C_{60}^{-1} I_b as experimentally already verified (even for the borderline case of Xe as theoretical investigations show). Even molecules like water fit into C_{60}.

If we take C$_{20}$ as a endohedral host molecule...
we arrive at a different picture as $R_{\text{MIS}}(C_{20}) - R_{\text{VdW}}(C) = 2.040 - 1.415\,\text{Å} = 0.635\,\text{Å}$, and even He does not fit anymore into the $C_{20}$ cage. We note the important role of dispersion interactions in endohedral fullerenes. Further, $C_{60}$ is a very stiff molecule, and the cage cannot so easily be expanded by endohedral atoms or molecules. In other words, the endohedral system can become repulsive very soon if the size of the endohedral atom or molecule becomes too large.

As a further example we consider hyperfullerenes (also called buckyonions), that is, fullerenes which contain smaller fullerenes inside their cage. They have already been detected and studied by theoretical methods. In a similar way to the rare gas estimates, we can derive approximately the following (conservative) condition for a non-repulsive enclosing hyperfullerene,

$$\Delta R_{N,M} = R_{\text{MIS}}(C_{N>M}) - R_{\text{MCS}}(C_M) - 2R_{\text{VdW}}(C) > 0$$ (49)

This only holds for ideal spherical systems, but can be seen as a lower limit for the guest $C_M$ fullerene. For $C_{20}@C_{60}$ we get $\Delta R_{60,20} = (3.553 - 2.084 - 2 \times 1.415)\,\text{Å} = -1.361\,\text{Å}$ and it is therefore predicted to be thermodynamically very unstable. Indeed, DFT(B3LYP) calculations predict $C_{20}@C_{60} \rightarrow C_{20} + C_{60} - 123.0\,\text{kcal/mol} (-15\,\text{kcal/mol per carbon atom})$, the high energy required to expand the $C_{60}$ cage. In fact, analyzing $C_{20}@C_{60}$ we find that the carbon atoms of $C_{20}$ close bonds with the carbon of the $C_{60}$ cage, that is, the structure should be considered as a carbon cluster rather than a hyperfullerene. Moreover, the volume of $C_{20}@C_{60}$ increases by 22% compared to $C_{60}$. In contrast, $C_{60}$ fits nicely into $C_{240}$, and $C_{240}$ just into $C_{540}$ in agreement with the analysis of Bates and Scuseria (see Figure 22). Here we get $\Delta R_{240,60} = 0.583\,\text{Å}$ and $\Delta R_{540,240} = 0.066\,\text{Å}$.

The buckyonions, which have been considered in the past, are all Goldberg-Coxeter transforms of $C_{20}$ and have the most spherical appearance as they are of icosahedral symmetry. It has been suggested that the rather high sphericity observed for buckyonions in experiments in contrast to the faceted polyhedral structure predicted (see Figure 22) is due to $C_2$ removal in a [c5666] fragment consisting of 1 pentagon and 3 hexagons (notation of Figure 16 is used here) introducing one heptagon and an additional pentagon, that is, [l575]. As heptagons introduce local negative Gaussian curvature (see discussion below), the fragment flattens out and the buckyonion becomes more spherical in shape.

For the buckyonions one can establish a simple relationship between the number of vertices $N > M$ of the buckyonions $C_{M}@C_N$. As $R \sim \sqrt{N}$ ($R$ being the radius of the cage), an analysis of $G_{C_{k,l}}$ transforms of fullerenes up to $k = 6$ shows that $R_{\text{MIS}} \approx 0.424\sqrt{M}\,\text{Å}$. This gives a very simple estimate from Eq (49) for such fullerenes,

$$\sqrt{N} \gtrsim 6.68 + \sqrt{M} \quad (N > M)$$ (50)

and we see that this is fulfilled for the cases already discussed, and for the largest buckyonion $C_{60}@C_{240}@C_{540}@C_{960}@C_{1500}$ considered by Bates and Scuseria. In fact, Casella et al. recently showed by using dispersion corrected DFT that the formation of $C_{60}@C_{180}$ from the two fullerenes is endothermic while $C_{60}@C_{240}$ is exothermic, and in the latter case only because dispersion interactions have been accounted for. We finally note that (possibly damaged) buckyonions are the main product in most gas phase fullerene generation procedures.

**Weird Fulleroidal Shapes: Generalizing Fullerene Structures**

Finally, what happens if we relax the rules a little bit, and allow for other types of three-valent ($sp^2$) carbon frameworks? There are many generalizations that lead to structures of beautiful shapes that have both elegant mathematical theory and physical realizations: allowing for polygons with faces different from pentagons and hexagons; for surfaces other than a sphere (genus 0), such as a torus (genus 1), a Klein bottle or a double
focus article

We will discuss mainly two types of carbon frameworks: fulleroids, which are fullerene-like structures, and Schwarzites, which are periodic negative-curvature surfaces. Different authors employ a wide range of working definitions for fulleroids. We advocate the following definition, generalizing fullerences: A fulleroid graph is a three-connected trivalent polyhedral graph. The definition is identical to that of fullerences, except that there is no restriction on face sizes. Hence, a fullerene is a fulleroid with only pentagons and heptagon faces. We further generalize to genus-\(n\) fulleroids, abbreviated \(G_{n}\)-fulleroids, defined as the three-connected trivalent genus-\(n\) graphs.

What kind of fulleroids are allowed? Can we tile a sphere or a torus with heptagons only (the answer is \(\text{no}\)), or with only pentagons and heptagons? (the answer is \(\text{yes}\), with the right number.) And how can we construct such fulleroids?\(^{310}\) In order to answer these questions, we need to generalize Euler’s equation to the Euler-Poincaré polyhedral formula for orientable surfaces,

\[
N - E + F = 2 (1 - g) = \chi \tag{51}
\]

where \(g\) is the genus of the graph,\(^m\) and the corresponding value \(\chi\) is called the Euler characteristic.\(^{39}\) The Klein bottle is a non-orientable surface (like the Möbius strip), and the corresponding (non-orientable) genus is \(g' = 2\) with an Euler characteristic of \(\chi = 0\) (here we take \(\chi = 2 - g'\) instead).

What happens when we allow vertex degrees \(r > 3\)? Consider a possibly irregular genus-\(n\) graph with \(N\) vertices. Let \(N_r\) be the number of vertices with degree \(r\), and \(F_n\) be the number of \(n\)-gonal faces. Then we trivially have

\[
F = \sum_n F_n, \quad N = \sum_r N_r, \quad \text{and} \quad 2E = \sum_n nF_n = \sum_r rN_r \tag{52}
\]

Combining this with the Euler-Poincaré polyhedral formula, we obtain

\[
2 \sum_{r \geq 3} (3 - r) N_r + \sum_{n \geq 5} (6 - n) F_n = 12 (1 - g) = 6 \chi
\]

\[
(53)
\]

If we only allow for three-valent graphs (fulleroids), the first term on the left hand side of Eq (53) conveniently vanishes (hence their special place in graph theory). This gives,\(^{311}\)

\[
3F_3 + 2F_4 + F_5 + \sum_{n \geq 7} (6 - n) F_n = 12 (1 - g) = 6 \chi
\]

\[
(54)
\]

From this equation we make a number of observations. a) We can play with as many hexagons as we want to build fullerenes or fulleroids, but the number of \(n\)-gons with \(n \neq 6\) is limited by Eq (54) or (53). b) While both equations allow for a large variety of face combinations, certain combinations are not allowed. The hand-shaking lemma (last formula in Eq (52)) allows only for an even number of vertices. We already know that there exist no polyhedral structures of the type \(C_{22}[5,6]\). To give another example, for \(C_{N}[5,7]\) polyhedra with pentagons and heptagons only (no hexagons) one requires at least two heptagons since, analogous to \(C_{22}\), the combination of 13 pentagons and one heptagon is not valid despite being allowed by Euler’s formula. Because of this, the smallest [5,7]-fulleroid is \(C_{28}[5,6,7]d\) with 14 pentagons and 2 heptagons, as shown in Figure 23.\(^n\) Structures containing only one heptagon are not forbidden, however, but requires introducing hexagons to form \(C_{N}[5,6,7]\)-structures. These can be obtained by replacing a \([c5666]\) fragment by \([l575]\),\(^{303}\) and can even be energetically favorable. For example, Fowler and co-workers showed that \(C_{62}\) consisting of one heptagon, 13 pentagon, and 19 hexagon faces is of lower energy than all the 2385 regular fullerene

**FIGURE 23** | \(C_{28}[5,7]-D_{7d}\), the smallest fulleroid with heptagon extension, compared to the most stable (by \(\Delta E = -102\) kcal/mol) \(C_{28}-T_d[5,6]\) fullerene isomer.
isomers of C\textsubscript{62}.\textsuperscript{312} c) For cubic planar graphs with only one type of face we get \( F_3 = 4 \) (tetrahedron), \( F_4 = 6 \) (cube), and \( F_5 = 12 \) (dodecahedron), which are the three three-valent polyhedra of the five Platonic Solids, the other two, the octahedron and icosahedron, require \( r = 4 \) and \( r = 5 \), respectively, in Eq (53); d) If we allow for two different \( n \)-gons, we arrive at the 13 Archimedean solids including our famous buckyball C\textsubscript{60}\textsuperscript{Ih}. e) Of course, we can tile a surface with triangles and the duals of fulleroids are prime examples, but in order to do this beside the tetrahedron, we must allow for different valencies of the vertices. f) We can tile a toroid \((g = 1)\) with hexagons only, consistent with Eq (54), see Figure 24. The torus is the only closed, orientable surface that can be tiled exclusively with hexagons. However, the plane, which is not closed, can be tiled with hexagons as well (graphene), as can the Klein-bottle (which is not orientable). Introducing pentagons into a toroid requires higher \( n \)-gons such as heptagons, since its total Gaussian curvature must be \( 2\pi(2 - 2 \cdot 1) = 0 \). The Euler-Poincaré’s formula allows to introduce pentagons into toroids if we introduce an equal number of heptagons, for example, we can tile a torus exclusively with pentagons and heptagons. Such ‘defects’ are well known in carbon nanotubes,\textsuperscript{313} which can be realized as infinite toroids with \( g = 1 \) and \( \chi = 0 \). Tiling in a pattern with alternating pentagons and heptagons yields a toroidal arrangement of azulenes, called azuloids.\textsuperscript{314}

A few examples should be mentioned here. The smaller fullerenes \((N < 60)\) are not very stable compared to C\textsubscript{60} because of the energy penalty one has to pay for fusing pentagons. As a result, structures deviating from the classical fullerenes can be more stable energetically or lie close by. For example, for C\textsubscript{26}, An et al. found six low lying isomers with the lowest structure being the fullerene C\textsubscript{26}\textsuperscript{D\textsubscript{3h}}\[5,6\], but only 3 kcal/mol lower in energy than the C\textsubscript{26}\textsuperscript{C\textsubscript{2}}\[4,5,6\], a fullroid that contains one square.\textsuperscript{315} Some combinations with heptagons are, however, disfavored energetically. For example, a Stone-Wales transformation\textsuperscript{9} of four connected hexagons to two pentagons and two heptagons in C\textsubscript{540}\textsuperscript{D\textsubscript{2h}} costs 260 kcal/mol.\textsuperscript{317}

Turning to toroidal fullerenoids (also called toroids) with \( g = 1 \) and \( \chi = 0 \), there are different techniques to construct such structures using either combinatorial or geometric approaches.\textsuperscript{318–320} Like in the nano-tubes, the hexagons can be oriented in different ways around the torus ring. For example, Figure 24(a) shows hexagons with 1/3 of the edges aligned tangential to the toroidal direction. We could also chose the hexagons to be aligned with 1/3 of the edges perpendicular to the toroidal direction (Figure 24(b)), and similar to nanotubes there are also chiral alignments depending on the chirality vector. Kirby and Pisanski showed how 2D graph drawings of toroids can be obtained.\textsuperscript{314} Borštnik and Lukman\textsuperscript{321} as well as Diudea and Kirby\textsuperscript{322} considered the structure and stability of such toroids using simple molecular mechanics. Faghani analyzed the symmetry of toroidal fullerenoids.\textsuperscript{126} Kang analyzed the band gap in such toroids by graph theoretical means.\textsuperscript{323} Deza et al. pointed out that leapfrog toroids and Klein-bottles have equal numbers of positive and negative eigenvalues, but with 4 and 2 eigenvalues being zero respectively.\textsuperscript{242} Interestingly, the leapfrog transformation performed on toroids (such as the one shown in Figure 24) results in an open shell toroid.\textsuperscript{324}

As discussed earlier, the positive curvature in fullerenoids originate from the pentagons, because sheets of hexagons like to be planar. Introducing heptagons or even octagons into a fullerene requires additional faces of size less than 6 to outweigh them, and results in a highly non-convex structure with often strong negative local Gaussian curvature,\textsuperscript{325} introducing a saddle-type topology to the structure. Two examples of such fullerenoids are shown in Figure 25. In this figure, every patch consisting of a pentagon surrounded by hexagons is replaced by six pentagons surrounded by

\[ \text{FIGURE 24} \] Torus consisting of hexagons only with 1/3 of the edges tangential (a) and perpendicular (b) to the toroidal direction.
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(a) (b) (c)

FIGURE 25 | Spiky fulleroids with negative curvature containing heptagons derived from fullerenes through patch replacement. (a) $C_{260}^{-}\{5, 7\}$ fulleroid derived from $C_{140}$. (b) $C_{300}^{-}\{6, 7\}$ derived from $C_{180}^{-}\{6\}$. (c) $C_{310}^{-}\{5, 6, 7\}$ derived from $C_{190}^{-}\{6\}$.

(a) (b)

FIGURE 26 | Peanut shaped fulleroids. (a) $C_{120}^{-}\{5, 6, 7\}$ with 10 heptagons and 22 pentagons. (b) $C_{168}^{-}\{5, 6, 7\}$ with 18 heptagons and 30 pentagons.

Five heptagons, for three different fullerenes, yielding characteristic ‘spiky’ fulleroids. In Figure 26, heptagons are introduced in a fashion that leads to peanut shaped fulleroids, and in fact, many types of interesting shapes can be constructed.

More interesting fulleroid shapes can be found in a recent paper by Diudea et al. Negatively curved fulleroids of higher genus (so-called perforated fullerenes) have been explored by Terrones and Terrones, who studied their electronic structures. Under specific conditions fullerenes can be combined by coalescence, forming larger carbon structures such as deformed nano-tubes or nano-peapods. For an overview of such carbon nano-peapods see Ref 331.

Structures of negatively curved graphitic carbon, which can be periodically extended to a lattice, have been proposed in 1991 by Mckay and Terrones and subsequently explored by experiment. These so-called ‘spongy’ carbon frameworks or Schwarzites were named in honor of the mathematician Hermann Schwarz (1843–1921), who investigated minimal surfaces with zero mean Gaussian curvature.
Schwarzites are structures containing (beside hexagons) heptagons and/or octagons and are of genus \( g \geq 3 \). These Schwarzites can be periodically continued through so-called junctions as shown in Figure 27 for the case \( g = 3 \) (Figure 27(a)). The two structures in Figure 27 are termed P-type and D-type, and can be approximately represented by the simple formulae derived from the Weierstrass-Enneper parameterization for minimal surfaces,

\[
\cos(x) + \cos(y) + \cos(z) = 0 \quad \text{(P-type)} \tag{55}
\]

\[
\sin(x)\sin(y)\sin(z) + \sin(x)\cos(y)\cos(z) + \cos(x)\sin(y)\cos(z) + \cos(x)\cos(y)\sin(z) = 0 \quad \text{(D-type)} \tag{56}
\]

Relevant reviews on such minimal surface carbon networks have been given by Terrones and Mackay\(^{311}\) and by Terrones and Terrones.\(^{338}\)

Lenosky et al. have performed local density functional calculations for two different \( g = 3 \) periodic Schwarzite structures with 216 atoms and a tessellation of hexagons and 24 heptagons within the unit cell.\(^{339}\) The vertices on such surfaces can be obtained through the Weierstrass representation.\(^{340,341}\) These calculations reveal a higher thermodynamic stability compared to \( C_{60} \), but not surprisingly lower compared to the graphene sheet (or graphite). D-type Schwarzites have the structure of a ‘diamond shaped’ lattice so that the unit cell can be split into two identical elements having 12 heptagons each, whereas P-type Schwarzites have the structure of a simple cubic lattice (see Figure 27). In principle, Schwarzites can be associated to any kind of lattice, either periodic or amorphous, the latter are realized in so-called random Schwarzites observed experimentally.\(^{332-335}\) Such random Schwarzites show rather nice transmission electron microscope (TEM) images (Figure 28) rather different to periodic Schwarzites. A recent comprehensive review over experimental and theoretical studies on Schwarzites has been given by Benedek et al.\(^{335}\)

Finally, we consider non-cubic (\( n \)-valent) polyhedral structures. In chemistry this is realized by using elements from the periodic table which can share more (or less) than three bonds with its neighbors. Prime examples are all-boron ‘fullerenes’,\(^{342}\) gaudiene \( C_{72} \cdot O_6 \) (with two- and three-valent carbon as analogs...
FIGURE 29 | All boron fullerene B_{40} (adapted from Ref 350). (a) 3D structure; (b) 2D graph.

CONCLUSIONS

To review the many developments in the topology and graph theory of fullerenes or fulleroids would be a monumental task. We have only outlined a few important concepts in order to give the reader a good introduction into this exciting field. There is certainly the need of a more comprehensive review or book. Nevertheless, we hope to have shown that the interplay between mathematics and structural chemistry is both interesting, rich and well alive. As more sophisticated method in the synthesis of carbon and other materials become available, we hope that some of the exotic, but very beautiful, structures become accessible with many useful applications in chemistry and materials science.

NOTES

a If we grow Goldberg-Coxeter transforms of C_{20} to infinity we obtain spherical graphene sheets connected by 12 pentagons. For definitions see below.
b An interesting side aspect is that fullerene-like shapes are realized also in viral shapes.\cite{20}
c Most of the useful graph theoretical and topological aspects discussed here are implemented into a Fortran/ C++ program called Fullerene, an open-source code freely available at the Massey University website.\cite{35}
d The embedding is unique in the following sense: All the planar embeddings of a three-connected graph are topologically equivalent, which means that they can be continuously deformed into each other without crossing any edges. Consequently, every planar embedding of such a graph defines the same set of faces. The converse it true as well: only three-connected graphs have this property.
Fulleroids are cubic graph fullerene-like structures where we allow also for other polygons than pentagons or hexagons.

In the mathematical sense that they are dense in this set, i.e. any smooth non-negative curvature genus 0 surface is the limit of a sequence of genus 0 triangulations with vertex degree six and less.

The interested reader is directed to the Gauss-Bonnet theorem, which generalizes Euler’s theorem to arbitrary Riemann surfaces. In the discrete case they are the same, except that Gauss-Bonnet is stated in terms of the Gaussian curvature. From this identification, we can see that every $n$-gon, or vertex of degree $n$ in the dual, contributes the angle $2\pi(6-n)/6$ to the Gaussian curvature, which must sum to $2\pi(2-2g)$, where $g$ is the surface’s genus.

When there are multiple exponentially crowded centers, the Tutte-embedding is still guaranteed planar, but the mapping to the sphere can have crossings. In practice, however, it turns out to not matter.

Counting perfect matchings for general graphs is \#P-complete. #P is the complexity class of counting solutions for decision problems in NP. While only exponential algorithms exist both for NP-complete and #P-complete problems, in practice #P-complete problems are dramatically harder than NP-complete.

It is known that not all connected cubic planar graphs are Hamiltonian, and that it is an NP-complete problem to determine whether any one cubic planar graph admits a Hamilton cycle. However, it is a long standing but yet unproven conjecture that fullerene graphs, a small subset of the cubic planar graphs, are all Hamiltonian. It has been verified by Brinkmann, Goedgebeur, and McKay for all $2.4 \times 10^{11}$ fullerene isomers up to $C_{316}$.

In Schwerdtfeger et al. (unpublished manuscript), we investigate in detail the Hamiltonian cycle counts admitted by fullerene graphs.

Fulleroids may in general have point group symmetries not admissible to the regular fullerenes. There are many other operations which can be performed on polyhedra resulting in transformed fullerenes or fulleroids, see the review by Vizitiu and Diudea.

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