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Calculating the number of Hamilton cycles in layered polyhedral graphs

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Abstract
We describe a method for computing the number of Hamilton cycles in cubic polyhedral graphs. The Hamilton cycle counts are expressed in terms of a finite-state machine, and can be written as a matrix expression. In the special case of polyhedral graphs with repeating layers, the state machines become cyclic, greatly simplifying the expression for the exact Hamilton cycle counts, and let us calculate the exact Hamilton cycle counts for infinite series of graphs that are generated by repeating the layers. For some series, these reduce to closed form expressions, valid for the entire infinite series. When this is not possible, evaluating the number of Hamiltonian cycles admitted by the series’ \( k \)-layer member is found by computing a \((k-1)\)\textsuperscript{th} matrix power, requiring \(\log_2(k)\) matrix-matrix multiplications. We demonstrate our technique for the two infinite series of fullerene nanotubes with the smallest caps. In addition to exact closed form and matrix expressions, we provide approximate exponential formulas for the number of Hamilton cycles.

KEYWORDS
cubic graph, finite-state machines, fullerene graph, Hamilton cycle, layered graph

1 INTRODUCTION

A Hamilton cycle is a spanning cycle in a graph \( G = (V, E) \), that is, a closed path that visits every vertex exactly once. In general, determining whether a given graph has at least one Hamilton cycle is \(NP\)-complete, including for cubic polyhedral graphs.\textsuperscript{1-3} Thus, counting all Hamilton cycles is \(#P\)-complete, the complexity class for counting solutions to \(NP\)-complete problems: a computationally much harder problem.\textsuperscript{4,5} Figure 1 shows the most famous example of a cubic polyhedral graph that does not admit a Hamilton cycle. For a history of Hamiltonian graphs, see Wilson.\textsuperscript{6}

Within the class of cubic polyhedral graphs, there is at least one subclass of \textit{layered cubic planar graphs}, defined by Franzblau, which are always Hamiltonian.\textsuperscript{9} Those are graphs that can be completely decomposed into—not necessarily identical—concentric cycles, where neighboring cycles are connected by two or more edges and where the first and last cycles are identical to a face.

In the special case of layered graphs that are the underlying graphs of closed nanotubes (a subclass of fullerenes), we can decompose the graph into layers, such that a sequence of layers is identical and consists only of hexagons. These identical layers are characterized by the chiral vectors \( C_h = (n, m) \).

Since layered cubic planar graphs are always Hamiltonian, counting their Hamiltonian cycles is no longer \#P-complete, opening for the possibility of discovering more efficient algorithms.
FIGURE 1  The Tutte graph ((A) a planar drawing of the graph, and (B) a 3D embedding produced using the software in References 7 and 8) is an example of a cubic polyhedral graph that is not Hamiltonian. In general, determining whether a given polyhedral graph admits a Hamilton cycle is an $NP$-complete problem, whereby counting the number of Hamilton cycles is a $\#P$-complete problem.

FIGURE 2  Illustration of the correspondence between Hamilton cycles and the bi-coloring defined in Lemma 1 for a Hamilton cycle for the dodecahedron, the $C_{20}$ fullerene.

In the following (Section 2), we first describe a method for counting Hamilton cycles in any cubic polyhedral graph. This method is applied to the special case of layered polyhedral graphs, for which it efficiently computes the number of Hamilton cycles. The method works not just for single graphs, but applies to infinite inductively defined series, yielding exact closed form expressions in simple cases, and expressions in terms of matrix powers for more difficult cases. In Section 3, we demonstrate it on the two infinite series of layered fullerene graphs with the smallest caps. A proof for the main theorem, showing that the method indeed produces exact Hamilton cycle counts, is given in Section 4.

2  COUNTING HAMILTON CYCLES IN A CUBIC POLYHEDRAL GRAPH

We introduce here a general technique for counting the Hamilton cycles in cubic polyhedral graphs. As the general problem is $\#P$ complete, it will in the fully general case require exponential time in the graph size, as it must. However, for layered graphs of bounded layer size, it reduces down to polynomial time. While we begin by describing how the method works for a single polyhedral graph, the method's strength—and the motivation for us to introduce it—is for finding exact expressions for the Hamiltonian cycle count for entire infinite series of layered polyhedral graphs.

We begin by identifying Hamilton cycles with certain bi-colorings. Figure 2 illustrates this correspondence. While this identification has been used before for layered graphs, we make it precise and prove its correctness for all cubic polyhedral graphs.

Lemma 1 (Hamiltonian cycles as bi-colorings). Given a cubic polyhedral graph $G$, and a choice of one of its faces $F_0$, there is a bijection between the set of Hamilton cycles and the set of face bi-colorings on $G$ that have the following three properties:

(a) $F_0$ is unshaded, (b) Every vertex is adjacent to at least one shaded and one unshaded face, and (c) all shaded and all unshaded faces are connected.

Proof. Let there be given a cubic polyhedral graph $G$ and a choice for $F_0$ (which will be the “outer face” in the planar drawings used to illustrate the method), and consider a Hamiltonian cycle $\gamma$ on $G$. By Jordan's curve theorem, $\gamma$ separates the surface uniquely into two connected regions, since it traces a simple closed curve on $G$'s surface. Let the entire region containing $F_0$ be white (without loss of generality), and the other region shaded. Hamiltonicity implies that every vertex of $G$ must lie on the boundary between the shaded and unshaded region. Hence, this bi-coloring fulfills properties (a), (b), and (c). It is unique since the color of $F_0$ is fixed, whereby it yields a well-defined mapping $f$ from the Hamilton cycles to the set of colorings that fulfill (a), (b), and (c). Finally, the edges comprising the Hamilton cycle are the boundary between the two regions, any two distinct cycles yield distinct colorings, whereby $f$ is injective.
Conversely, let there be given a bi-coloring of $G$'s faces fulfilling properties (a), (b), and (c). By property (c), the edge set comprising the boundary forms a simple cycle $\gamma$, which by property (b) visits every vertex in $G$, that is, it is a Hamilton cycle. Hence, this bi-coloring is the value $f(\gamma)$, whereby $f$ is also surjective.

In all, $f$ is bijective, so we can identify the Hamiltonian cycles through $f$ with the bi-colorings that fix the color of $F_0$ and fulfill (b) and (c).

**Definition 1** (Cap). A cap is a subgraph that has exactly one boundary. It must have at least one face and no leafs.

**Definition 2** (Layer). A layer is a subgraph that has exactly two boundaries which do not have any vertices in common. It may not have any leaves. That is, it has exactly two holes with respect to the parent graph.

For a polyhedral graph, exactly two caps can be defined, corresponding to two opposite ends of the polyhedron. The caps are not uniquely defined, however, but can be chosen in many ways, as can the subdivision of the remaining graph into layers between the two caps. In the following, we consider a polyhedral graph $G$ and assume that this choice has been made, such that $G$ is composed of a start-cap $A$, an end-cap $Z$, and in between them: $k$ layers. Each layer has an inner and an outer boundary cycle. If $k = 1$, the outer boundary is the boundary of $A$, and the inner cycle is the boundary of $Z$. If $k \geq 2$, the two terminal layers share a boundary with $A$ and $Z$, respectively, and for $1 \leq i < k$, the inner boundary of layer $i$ is the outer boundary of layer $i + 1$.

We write $S \oplus T$ for the combined graph of subgraphs $S$ and $T$, merged by identifying the inner boundary of $S$ with the outer boundary of $T$. $A$ and $Z$ only have one boundary each, respectively considered “inner” and “outer.”

Consider now a polyhedral graph that is partitioned into two caps and zero or more layers. We do not address here the question of how to optimally decompose a graph into caps and layers.

The second important step needed to construct the Hamilton cycle counting method is the following locality condition, which ensures that we can count incrementally by way of a finite state-machine. We start from one cap and end in the other:

**Lemma 2** (Locality). When shading in a graph which is decomposed into layers according to the rules defined in Lemma 1, the legal colorings of each layer depend only on the previous layer and which of the shaded and unshaded faces in the previous layer are already connected, that is, which shaded faces in the previous layer are connected by shaded faces in the subgraph that consists of all previous layers and the starting cap (and respectively for unshaded faces).

**Proof.** By Lemma 1(c), for a shading of faces to correspond to a Hamilton cycle, it is required that all shaded and all unshaded faces, respectively, are connected after all faces are colored in.

During the process of shading in the graph layer-wise, the set of shaded faces may temporarily be disconnected. However, in order to connect them through the following layers, each shaded face must through other shaded faces be connected to the boundary where new layers are added. This analogously applies to unshaded faces.

In order to fulfill this requirement, we must include information on the connectedness of shaded faces into a local layer description.

Lemma 1 requires that each vertex is adjacent to a shaded and an unshaded face. As all vertices in a graph that is decomposed into layers are either interior to the layer or lie on the boundary between one layer and the next, the face adjacency of each vertex is fully determined by the two layers and their orientation and does not depend on any other layers.

Because of the locality condition of Lemma 2, we can model the counting as a finite state machine, in which the finitely many states are the legal shadings of single layers together with the information about which shaded faces are connected. We represent the state machine as a directed multigraph $M$, in which the edges represent transitions to states that are accessible from the corresponding state. The directed subgraph that describes the transition from one layer to the following one is denoted as $M^{(p)}$. If one state can be followed by another state in more than one way (due to symmetry), they are connected by multiple directed edges. We call the vertices that correspond to starting points initial states (typically every valid shading of the first cap), and those that can be final layers in a Hamilton cycle accepting states (typically every valid shading of the last cap).

The number of Hamilton cycles admitted by a graph that consists of two caps and $k$ layers is then the number of paths of length $k + 1$ in $M$ that originate in an initial state (start-cap bi-coloring) and end in an accepting state (end-cap bi-coloring). See Figure 3(B).

The state-machine can equivalently be represented by a very sparse square matrix $M$, with $M_{st} = n$ if $s \rightarrow^nt$ is an arc of $M$, and $M_{st} = 0$ otherwise. In the $k$th power of $M$, the entry $M_{st}^k$ is the number of length-$k$ paths from state $s$
FIGURE 3 (A) The polyhedral graph is subdivided into two caps A and Z, separated by k layers \( L_1, \ldots, L_k \). We identify A with \( L_0 \) and Z with \( L_{k+1} \). (B) For each layer \( L_p \), the symmetry-distinct shadings that do not violate Lemma 1 are considered “valid” and are added as a state in the DFA. Lemma 1 then determines the valid combinations of shadings from \( L_p \) to shadings in \( L_{p+1} \), defining the arcs in the DFA. (C) The DFA can be described by a large matrix \( M \), which is zero everywhere except in the blocks \( M^{(p)} \) just above the diagonal, which count the valid combinations of \( L_p \)-shadings to \( L_{p+1} \)-shadings; here shown for \( k = 3 \).

to state \( t \). Hence, the number of Hamilton cycles can be counted by raising \( M \) to the \( k^{th} \) power and summing the elements of the AZ-subblock, consisting of the initial-state rows and final-state columns. This sum is the number of Hamilton cycles.

\( M \) is zero everywhere except in the blocks \( M^{(p)} \) directly above the diagonal, representing the directed subgraphs \( M^{(p)} : L_p \rightarrow L_{p+1} \) that link layer \( p \) to layer \( p + 1 \), as shown in Figure 3(C) (where we identify \( A = L_0 \) and \( Z = L_{k+1} \)). The small blocks \( M^{(p)} \) are still sparse, but rectangular with dimensions \( N_p \times N_{p+1} \), where \( N_p \) is the number of valid shadings for layer \( p \) (with \( A \) and \( Z \) as \( p = 0 \) and \( p = k + 1 \)).

Because of this, we can calculate the number of Hamilton cycles as a single length-(\( k + 1 \)) product of the much smaller blocks \( M^{(p)} \). The entry \( M^{(p)} \) in the adjacency matrix is the number of single-step transitions from state \( s \) in layer \( p \) to state \( t \) in layer \( p + 1 \). In the product of two consecutive matrices \( M^{(p)} \) and \( M^{(p+1)} \), the entry \((s, t)\) is the number of length-2 paths from state \( s \) in layer \( p \) to state \( t \) in layer \( (p + 2) \), and so forth. The product of the \( k + 1 \) matrices \( M^{(p)} \) directly yields the AZ-subblock of \( M^k \), and thus sums to the number of Hamilton cycles. This representation makes calculations convenient, shown in Equations (1) and (3) below.

2.1 An acyclic finite state-machine that counts Hamilton cycles in polyhedral graphs

We are now ready to construct a finite state machine that counts the Hamiltonian cycles in infinite series of cubic layered-graphs.

The following procedure constructs a deterministic finite automaton (DFA), such that paths of length \( k + 1 \) starting in an initial state and ending in a final state count the number of Hamilton cycles in \( G_k \):

**Definition 3** (Hamilton cycle-counting DFA). Let \( A \) and \( Z \) be two caps, and let \( G_k \) be a graph that is decomposed into \( A \), followed by \( k \) layers, and end-capped with \( Z \). Construct a finite state machine as follows:

**Initialization:**
- I.1 Define the initial states as the symmetry-distinct shadings of \( A \) that do not violate Lemma 1.
- I.2 Initialize the set \( S \) of states to be equal to the initial states.

**Recursion:**
- R for each of the \( k \) layers and \( Z \):
- (a) for each state \( s \) in \( S \):
  - i. Write the associated shading as \( S(s) \).
For each connectedness-labeled shading \( t \) of the current layer for which \( S(s) \oplus S(t) \) fulfills L1(a)–(b) (and if we have reached the cap \( Z \), also fulfills (c)):

A. If \( t \) is not already an existing state in \( \mathcal{T} \) (up to symmetry), create it and add it to \( \mathcal{T} \).
B. Draw an arc \( s \to t \).
C. Label the arc with the number of legal ways to identify the inner boundary of \( s \) with the outer boundary of \( t \).

(b) move \( \mathcal{T} \) to \( S \)

**Theorem 1** (The Hamilton cycle-counting DFA counts Hamilton cycles). Let \( \{ G_k \}_{k \in \mathbb{N}_0} \) be a graph which is decomposed into \( k \) layers and the two caps A and Z, and let \( M \) be the state machine of Definition 3. Then (i) each Hamilton cycle of \( G_k \) corresponds to a length-(\( k + 1 \)) path from an initial state in \( M \) to a final state in \( M \), and (ii) the product of the labels along the path counts how many Hamilton cycles correspond to this path. (iii) Equivalently, let \( M \) be the matrix such that \( M_{st} = n \) if \( s \to^n t \) is an arc of \( M \), and \( M_{st} = 0 \) otherwise. Writing \( P = M^{k+1} \) for the \( (k + 1) \) matrix power of \( M \), the total number of Hamilton cycles in \( G_k \) is:

\[
N_{HC}(G_k) = \sum_{s \in \text{Initial}} \sum_{t \in \text{Final}} P_{st}. \quad (1)
\]

Writing \( M^{(0)}, M^{(1)}, \ldots, M^{(k)} \) for the \( k + 1 \) blocks of \( M \) that link consecutive layers (see Figure 3(C)), we can write \( P_{st} \) as a product of these smaller rectangular matrices:

\[
P_{st} = (M^{(1)} \times \cdots \times M^{(k+1)})_{st}. \quad (2)
\]

The proof is given in Section 4 below.

### 2.2 A cyclic finite state-machine that counts Hamilton cycles in infinite series of polyhedral graphs

The real strength of the method becomes apparent when we consider an infinite series of layered polyhedral graphs, in which identical layers are repeated arbitrarily many times. That is, instead of considering a single polyhedral graph, we consider an entire series. This can be modeled simply by adding cycles to the state machine. In the general case, this yields a matrix product that can be evaluated to yield the exact Hamilton cycle count for any member of the series, and in neater cases such as shown in Section 3, we can obtain closed form expressions that apply to the entire series.

To simplify the following definition, we may fuse any number of non-repeating layers with A or Z, such that the obtained state machine contains only one type of layer of which there are zero or more instances. One can shade these layers separately and produce the product matrices \( M_{AL} \) and \( M_{LZ} \) layer-wise for efficiency.

The two caps A and Z together with a single layer \( L \) define an infinite series of polyhedral cubic graphs in the following way:

**Definition 4** (Layered polyhedral graph series). The layered polyhedral graph series for two caps A and Z and a single layer \( L \) is an infinite series of polyhedral graphs \( \{ G_k \}_{k \in \mathbb{N}_0} \) such that \( G_k \) consists of A and Z connected by \( k \geq 0 \) instances of the layer \( L \). The repeated layer must have a total Gaussian curvature of zero, such that a variable number of layers is permissible without violating the discrete Gauss-Bonnet theorem.

The total number of Hamilton cycles in \( G_k \) can now be simplified from Equation (1), where we needed \( k + 1 \) matrix blocks, to just:

\[
N_{HC}(G_k) = \sum_{s \in \text{Initial}} \sum_{t \in \text{Final}} (M_{AL} \times M_{LL}^{k-1} \times M_{LZ})_{st}, \quad (3)
\]

where \( M_{AL} \) describes the transition from A to \( L \), \( M_{LL} \) is the square matrix describing the transition between two instances of \( L \), and \( M_{LZ} \) describes the transition from \( L \) to Z, see Figure 4(C).
FIGURE 4 (A) When polyhedral graphs can be decomposed into two caps A and Z, separated by k repetitions of the identical layers L, the situation simplifies greatly. (B) The repeated layer is modeled with cycles in the DFA. Again, Lemma 1 determines the legal shadings, as well as the arcs that connect layer shadings that can combine to eventually form Hamilton cycles. (C) The matrix form of the cyclic DFA has three (sparse) nonzero blocks: the two super-diagonal blocks representing combining cap A with the first layer and the last layer with cap Z, and in addition an $L \rightarrow L$ block on the diagonal, which captures how the repeated layer $L$ combines with itself.

\[
\begin{array}{c|cc}
A & 0 & M_{AL} & 0 \\
L & 0 & M_{LL} & M_{LZ} \\
Z & 0 & 0 & 0 \\
\end{array}
\]

FIGURE 5 Two representative polyhedra for (5,0) fullerene nanotubes, (A) $C_{70}$-$D_{5h}$ and (B) $C_{60}$-$D_{5d}$. Pentagon caps are distinguished from the hexagons by shading.

We can thus either compute $N_{HC}(G_k)$ directly for a particular $k$ with $O(\log_2(k))$ matrix-matrix multiplications by the standard recursive algorithm for computing powers, or for all of $k = 1, \ldots, K$ with $K + 1$ matrix-matrix multiplications in total. We will see that we can sometimes obtain either exact or approximate closed form expressions, when the infinite layered graph series under study is sufficiently well-behaved.

The correctness of this cyclic DFA follows by construction from the fact that it is a special case of the state machine defined in Definition 3 which counts the number of Hamilton cycles according to Theorem 1.

3 | EXAMPLES

As examples we examine two series of graphs: those of the nanotubes with the smallest diameters. In both cases, the graphs fall into the class of layered graphs as defined by Franzblau. In the first case, the chiral vector is $C_h = (5,0)$, implying caps that consists of six fused pentagons and no hexagon, i.e., they have 10 vertices each. Each layer consists of 5 hexagons and adds 10 vertices, yielding a total of $N = 20 + 10k$ vertices with $k = 0, 1, 2, \ldots$ for the k-layer graph. Each layer induces a twist of $\pi/5$ of the end-pentagons with respect to each other. Since the main axis of symmetry runs through the ends, the result is that the symmetry alternates between $D_{5d}$ for even $k$, and $D_{5h}$ for odd $k$. We denote this series as $D_{5+}$. Figure 5 shows two representative examples of this series, $C_{70}$-$D_{5h}$ and $C_{60}$-$D_{5d}$.

The smallest $D_{5+}$ nanotube is $C_{20}$, which is also the smallest fullerene. It has exactly 30 Hamilton cycles, which are all equivalent under $I_h$-symmetry, but up to the symmetry of the $D_{5+}$-nanotube series (5-fold rotation and reflection), the three cycles shown in Figure 6 are distinct.
The three $D_3$-symmetry distinct Hamilton cycles out of in total 30 for $C_{20}$-$I_h$. The Hamilton cycle traces the boundary of the green shaded area. Note that, in the full $I_h$-symmetry, these are symmetry-equivalent as well.

The second series has a chiral vector of $(6,0)$ and caps consisting of one hexagon which is surrounded by six pentagons. Here the caps consist of 12 vertices each and each layer contributes 12 vertices, yielding a total of $N = 24 + 12k$ vertices per graph. Members of this series alternatingly have $D_{6d}$ (for even $k$) and $D_{6h}$ (for odd $k$) point group symmetry. We denote this series as $D_{6+}$. Figure 7 shows two representative examples of this series, $C_{60}$-$D_{6h}$ and $C_{72}$-$D_{6d}$.

In the remainder of this section, we derive expressions for the number of Hamilton cycles in these two series of layered graphs. Following the same approach, expressions for any other series of layered graphs can be constructed.

### 3.1 Syntax

In order to keep track of the information needed in Lemma 2, we introduce the following notation for a partial coloring of a layer in a layered graph: (a) All faces in the layer are numbered counter clockwise, starting at 1. (b) Each set of shaded faces that is connected, either because they are neighbors, or through shaded faces in previous layers, is written as a comma separated list enclosed in parenthesis. The elements in the lists are ordered ascendingly. (c) The connected groups are ordered ascendingly by their respective first elements. (d) The starting point of the counter clockwise numbering is arbitrary, but needs to be chosen consistently for all layers in the graph (up to rotation and reflection with respect to the connections between consecutive layers). If the selected first face is not unique due to symmetry, it is chosen such that we start counting in the largest connected group, and each following face is given the smallest possible number, while larger connected groups have preference to get smaller indices and hence be listed first. If the layer has reflection symmetry, both clockwise and counter clockwise numbering has to be considered, that is, mirror images are given the same label.

It is not necessary to include the total number of faces in a layer, because only one size occurs in a chosen layered graph. Furthermore, it is not necessary to specify the connectedness of the unshaded faces, because it follows implicitly from the shaded faces and the fact that we are dealing with planar graphs.

For example, “(1, 2) (4)” denotes a layer with two neighboring shaded faces that are connected to each other and a third face at position “4” which is not connected to them through previous layers. “(1, 2, 4)” denotes a layer that is locally identical, but where all three faces are connected through previous layers. In the examples in Section 3, the layers have 5-fold and 6-fold rotational symmetry, respectively, as well as reflection symmetry. Therefore, all faces are equivalent with respect to the connections to the previous and the following layer, and we chose the starting point and orientation of the numbering such that indices are minimized as explained above.

As is conventional, the initial states are indicated by an arrow, and the final states are drawn with double circles.
3.2 The number of Hamilton cycles in $C_{20+10k}^{-}D_{5+}(5,0)$ nanotubes

We now turn to the series of $C_{20+10k}^{-}D_{5+}(5,0)$ fullerene nanotubes. This is a $k$-layered polyhedral graph for every $k = 0, 1, \ldots$ (where $k$ counts the inner layers). The smallest specimen is $C_{20}$, with two layers and $k = 0$.

**Theorem 2.** The $D_{5+}$ fullerene nanotube $C_{20+10k}$ ($k \in \mathbb{N}_0$) admits exactly

$$N_{HC}(k) = 5 \left[ 2^{k+1} + 4 \cdot 12^k \chi(k) \right]$$

Hamilton cycles, where $\chi(k) = 1$ for even $k$, and $\chi(k) = 0$ for odd $k$.

**Proof.** Consider a $C_{20+10k}^{-}D_{5+}$ fullerene nanotube. In the planar drawing below, let $F_0$ be the outer face, unshaded. Up to the $D_5$-symmetry common for the nanotube-series, there are two inequivalent ways to shade the faces in the first cap of the graph that do not violate Lemma 2, each of multiplicity 5.

All other colorings of the cap yield either vertices that are not adjacent to a shaded face, or the unshaded faces are separated into two disconnected areas (by shading in all five faces that are adjacent to the boundary). It will be seen below that Hamilton cycles starting in cap-type $A$ give rise to the term $5 \times 2^{k+1}$, and type $B$ to the term $5 \times 4 \times 12^k \chi(k)$.

Assume that the cap is shaded as $A$. One can easily show using Lemma 2 that only two layer types need to be considered for shading of all remaining layers (up to reflection). Using the notation introduced above, these are $(1)$ and $(1, 2, 3, 4)$:

A $(1, 2, 3, 4)$-layer must be followed by a $(1)$-layer in one of two ways, and a $(1)$-layer must be followed by a $(1, 2, 3, 4)$-layer in one of two ways:

for which the adjacency matrix is:

$$A = \begin{pmatrix} 0 & 2 \\ 2 & 0 \end{pmatrix}$$

The possible end-cap shadings that can follow states $(1)$ and $(1, 2, 3, 4)$ are $W$ and $X$, respectively. $W$ can follow state $(1)$ in two ways and $X$ can follow state $(1, 2, 3, 4)$ in two ways.
As state (1) can follow on cap A in one of two ways, and for each of (1) and (1, 2, 3, 4), there is a cap which may follow in one of two ways, we obtain the two terminal adjacency matrices \((2 0)\) and \((0 2 0)\). The number of Hamilton cycles of \(C_{20+10k}\) that originate from case A is:

\[
N_{\text{HC}}^{(A)}(k) = 5 \times \sum \left( 2 0 \right)^{k-1} \times \left( 0 2 \right) = 5 \times 2^{k+1}. \tag{10}
\]

When case B is chosen for the cap, there are four layer types to be considered for shading in the remainder of the graph: (1, 3), (1, 3), (1, 2, 4), and (1, 2, 4).

This is described by the following DFA, which with the order ((1, 2, 4), (1, 2, 4), (1, 3), (1, 3)) has the adjacency matrix B:

\[
\begin{align*}
(1,2)(4) & \quad \rightarrow \quad (1,2,4) \\
2 & \quad \rightarrow \quad 3 \\
2 & \quad \rightarrow \quad 3 \\
3 & \quad \rightarrow \quad 2 \\
(1,3) & \quad \rightarrow \quad (1,3) \\
\end{align*}
\]

\[
B = \begin{pmatrix} 0 & 2 & 2 & 0 \\ 0 & 0 & 0 & 3 \\ 3 & 0 & 0 & 0 \\ 0 & 2 & 2 & 0 \end{pmatrix}
\tag{13}
\]

with accepting states corresponding to columns 1 and 4.

Cap B may be followed by (1, 2, 4) and (1, 3) in one of two ways each.

States (1, 2, 4) and (1, 3) may be followed by either of caps Y and Z in one of two ways each. No other caps are possible because shading the central last face differently would either yield multiple cycles by disconnecting the shaded or unshaded area, or exclude one vertex from the cycle, both violating Lemma 1.

Thus, the number of Hamilton cycles in \(C_{20+10k}\) that start from case B is

\[
N_{\text{HC}}^{(B)}(k) = 5 \times \sum \left( 0 2 2 0 \right)^{k-1} \times B^{k-1} \times \left( 2 2 \right) = 5 \times (B_{1,2}^{k+1} + B_{1,3}^{k+1}). \tag{15}
\]
To understand the alternating term in Equation (4), we compute the first few powers of $B$:

$$B^2 = \begin{pmatrix}
6 & 0 & 0 & 6 \\
0 & 6 & 6 & 0 \\
0 & 6 & 6 & 0 \\
6 & 0 & 0 & 6
\end{pmatrix}, \quad B^3 = \begin{pmatrix}
0 & 24 & 24 & 0 \\
18 & 0 & 0 & 18 \\
18 & 0 & 0 & 18 \\
0 & 24 & 24 & 0
\end{pmatrix}, \quad B^4 = \begin{pmatrix}
72 & 0 & 0 & 72 \\
0 & 72 & 72 & 0 \\
0 & 72 & 72 & 0 \\
72 & 0 & 0 & 72
\end{pmatrix}. \quad (17)$$

We now prove the even and odd cases separately by induction. Assume for some odd $k \geq 1$ that $B^{k+1}$ is of the form

$$\begin{pmatrix}
x & 0 & 0 & x \\
0 & x & x & 0 \\
0 & x & x & 0 \\
x & 0 & 0 & x
\end{pmatrix}.$$  

Then the next odd $k$ yields:

$$B^3B^{k+1} = \begin{pmatrix}
6 & 0 & 0 & 6 \\
0 & 6 & 6 & 0 \\
0 & 6 & 6 & 0 \\
6 & 0 & 0 & 6
\end{pmatrix} \begin{pmatrix}
x & 0 & 0 & x \\
0 & x & x & 0 \\
0 & x & x & 0 \\
x & 0 & 0 & x
\end{pmatrix} = \begin{pmatrix}
12x & 0 & 0 & 12x \\
0 & 12x & 12x & 0 \\
0 & 12x & 12x & 0 \\
12x & 0 & 0 & 12x
\end{pmatrix} = 12B^{k+1}. \quad (18)$$

Since $B^2$ (the case $k = 1$) is of this form, we obtain by induction that $B^{k+1}$ is of this form for every odd $k \geq 1$, by which we obtain $B^{k+1} + B^{k+1} = 0$ for every odd $k \geq 1$. That is, the number of Hamiltonian cycles starting in configuration $B$ is 0 whenever the number of layers $k$ is odd.

For the even case, assume for some even $k \geq 0$ that $B^{k+1}$ is of the form

$$\begin{pmatrix}
0 & x & x & 0 \\
y & 0 & 0 & y \\
y & 0 & 0 & y \\
0 & x & x & 0
\end{pmatrix}.$$  

Then the next even $k$ gives:

$$B^2B^{k+1} = \begin{pmatrix}
6 & 0 & 0 & 6 \\
0 & 6 & 6 & 0 \\
0 & 6 & 6 & 0 \\
6 & 0 & 0 & 6
\end{pmatrix} \begin{pmatrix}
0 & x & x & 0 \\
y & 0 & 0 & y \\
y & 0 & 0 & y \\
0 & x & x & 0
\end{pmatrix} = \begin{pmatrix}
0 & 12x & 12x & 0 \\
12y & 0 & 0 & 12y \\
12y & 0 & 0 & 12y \\
0 & 12x & 12x & 0
\end{pmatrix} = 12B^{k+1}. \quad (19)$$

Since $B (B^{k+1} \text{ for } k = 0)$ is of this form, we obtain by induction that this holds for every even $k \geq 0$. How many Hamilton cycles start in configuration $B$ for $k$ inner layers when $k$ is even? For $k = 0$, we read off the number to be $5 \times (2 + 2) = 5 \times 4 \times 12^{\frac{k-1}{2}}$. Finally, assuming the count is $5 \times 4 \times 12^{\frac{k-2}{2}}$ for some even $k - 2 \geq 0$, the count for $k$ (by Equation 19) is

$$N_{HC}^{(B)}(k) = 5 \times 4 \times 12^{\frac{k-2}{2}} \times 12 = 5 \times 4 \times 12^{\frac{k}{2}} \text{ (k even)}, \quad (20)$$

whereby, by induction, the formula holds for all even values of $k \geq 0$. Together, Equations (10) and (20) complete the proof.

3.3 The number of Hamilton cycles in $C_{24+12k-D_{6+}(6,0)}$ nanotubes

We next turn to the number of Hamilton cycles in the $C_{24+12k-D_{6+}(6,0)}$ fullerene nanotube series, the next-thinnest class of layered nanotubes.

**Theorem 3.** The $C_{24+12k-D_{6+}(6,0)}$ fullerene nanotube ($k \in \mathbb{N}_0$) admits

$$N_{HC}(k) \approx 16 \times 2^k + 55.67911535839 \times 2^{2.1679870k} \quad (21)$$

Hamilton cycles. The exact expression for the Hamilton cycle count is given in Equation (38).
Proof. There are four inequivalent options for shading in the first cap of the graph up to $D_6$-symmetry, with multiplicities 6, 6, 3, and 2, respectively:

\begin{align*}
\begin{array}{cccc}
A & B & C & D \\
\times 6 & \times 6 & \times 3 & \times 2
\end{array}
\end{align*}

Starting from the first case (A) we need to consider only two different layer shadings (in analogy to case A of the previous proof), namely, $(1, 2, 3, 4, 5)$ and $(1)$:

\begin{align*}
\begin{array}{c}
(1, 2, 3, 4, 5) \\
(1)
\end{array}
\end{align*}

Both layers may follow each other according to the following graph:

\begin{align*}
\begin{array}{c}
(1) \\
\overset{2}{\rightarrow}
\end{array}
\begin{array}{c}
(1, 2, 3, 4, 5)
\end{array}
\end{align*}

which can be written as the adjacency matrix $U = \begin{pmatrix} 0 & 2 \\ 2 & 0 \end{pmatrix}$.

State $(1, 2, 3, 4, 5)$ may follow on $A$ in two ways, and states $(1, 2, 3, 4, 5)$ and $(1)$ may be followed by caps $V$ and $W$ respectively in two ways each:

\begin{align*}
\begin{array}{c}
V \\
\end{array}
\begin{array}{c}
W
\end{array}
\end{align*}

The number of Hamilton cycles starting from cap $A$ is therefore

\begin{align*}
N_{HC}^{(A)}(k) &= 6 \times \sum \left( \begin{pmatrix} 2 & 0 \\ 2 & 0 \end{pmatrix} \times \begin{pmatrix} 0 & 2 \\ 2 & 0 \end{pmatrix}^{k-1} \times \begin{pmatrix} 0 & 2 \\ 2 & 0 \end{pmatrix} \right) \\
&= 6 \times (U_{1,1}^{k+1} + U_{1,2}^{k+1}) = 6 \times 2^{k+1}.
\end{align*}

Caps $B$ and $C$ can be treated together because starting from either leads to the same options for shading in the remainder of the graph. We need to consider $(1, 2, 3, 5)$, $(1), (1, 2, 4), (1, 4)$, and $(1, 2, 4, 5)$.

\begin{align*}
\begin{array}{cccc}
(1, 2, 3, 5) & (1) & (1, 2, 4) & (1), (1, 2, 4, 5)
\end{array}
\end{align*}
We obtain the following graph $V$ and adjacency matrix $V$:

![Graph Image]

$$V = \begin{pmatrix} 0 & 2 & 2 & 0 & 0 \\ 2 & 0 & 0 & 0 & 1 \\ 0 & 1 & 3 & 1 & 0 \\ 0 & 2 & 0 & 0 & 2 \\ 0 & 0 & 4 & 2 & 0 \end{pmatrix} \quad (29)$$

Cap $B$ may be followed by $(1) (3)$ and $(1, 2, 4)$ in two ways each, while cap $C$ may be followed by states $(1, 2, 4)$ and $(1) (4)$ in four and two ways, respectively.

There are two end caps. $X$ may follow on states $(1, 2, 3) (5)$ and $(1, 2, 4)$ in two and one way, respectively, while $Y$ may follow $(1, 2) (4, 5)$ and $(1, 2, 4)$ in two and one way, respectively.

![Diagram Image]

Keeping in mind the multiplicities of the first layers, we get

$$N_{HC}^{(B)}(k) + N_{HC}^{(C)}(k) = \sum \left( \begin{pmatrix} 6 & 3 \\ 0 & 0 \\ 0 & 4 \\ 2 & 0 \end{pmatrix} \times \begin{pmatrix} 0 & 2 & 2 & 0 & 0 \\ 0 & 0 & 4 & 2 & 0 \end{pmatrix} \times V^{k-1} \times \begin{pmatrix} 2 & 0 \\ 0 & 0 \\ 1 & 1 \\ 0 & 0 \\ 0 & 2 \end{pmatrix} \right) \quad \text{with} \quad (31)$$

$$N_{HC}^{(B)}(k) = 6 \times (V_{1,2}^{k+1} + V_{1,4}^{k+1}) \quad \text{and} \quad (32)$$

$$N_{HC}^{(C)}(k) = 3 \times (V_{5,2}^{k+1} + V_{5,4}^{k+1}). \quad (33)$$

Cap $D$ only allows for choosing state $(1) (3) (5)$ for all following layers.

![Diagram Image]

However, there are two possible orientations for each layer.

![Diagram Image]
There is one possible cap $Z$ that may follow (1) (3) (5) in either of two ways.

The adjacency matrix is (2), yielding

$$N_{HC}(D_d)(k) = 2 \times (2)^{k-1} \times (2) = 2 \times 2^{k+1}$$

Hamilton cycles starting from cap $D$. In total the number $N_{HC}$ of Hamilton cycles in $C_{24+12k}(6,0)$ nanotubes is

$$N_{HC}(k) = N_{HC}^{(A)}(k) + N_{HC}^{(B)}(k) + N_{HC}^{(C)}(k) + N_{HC}^{(D)}(k)$$

$$= 6 \times 2^{k+1} + 6 \times (V_{1,2}^k + V_{1,4}^k) + 3 \times (V_{5,2}^k + V_{5,4}^k) + 2 \times 2^{k+1}.$$  \hspace{1cm} (38)

Due to the complicated behavior of the matrix powers $V^k$, it is not possible to derive a simple closed form expression. However, while the exact counts have many complicated terms, they do asymptotically approach simple exponentials: $(V_{1,2}^k + V_{1,4}^k) \to 1.0862679 \times 2.1679870^k$, and $(V_{5,2}^k + V_{5,4}^k) \to 1.9573872 \times 2.1679870^k$, yielding 16-digit accuracy already at $k = 35$, improving as $k$ grows (tested up to $k = 3000$). Hence, a simple approximate expression can be found:

$$N_{HC}(k) \approx 16 \times 2^k + 55.6791153839 \times 2.1679870^k.$$  \hspace{1cm} (39)

### 3.4 Long-range exponential behavior of $C_{20+10k}$-$D_5^+$ nanotube series versus $C_{24+12j}$-$D_6^+$ series

Our original interest in finding expressions for the Hamiltonian cycle counts of the infinite series was due to our work in Reference 11, which originally suggested that the $C_{20+10k}$-$D_5^+$ series defined both strict upper and lower bounds to the number of Hamiltonian cycles admitted by any fullerene graphs. This was verified for the tens of millions of graphs up to $C_{120}$. However, while the conjecture that this $D_{5d}$ nanotube series maximizes Hamiltonian cycle count looked extremely plausible, the exact expressions allowed us to produce a counter-example—in a region that could never have been checked exhaustively by existing methods.

The crossover-point is $C_{1020} = C_{24+83 \times 12} = C_{20+100 \times 10}$ (i.e., $k = 100$ and $j = 83$). From $C_{1020}$ on, every $C_{24+12j}$-$D_6^+$ break the conjectured upper bound due to the larger exponent taking over. $C_{1020}$-$D_{5d}$ has

$$N_{HC}(C_{1020}-D_{5d}) = 18,200,087,300,004,299,546,655,183,227,191,134,932,148,320,173,545,226,240$$  \hspace{1cm} (40)

![FIGURE 8](image-url) Hamilton cycle counts for the $D_{5d}$ (solid) and the $D_6$-series (dashed). Near to the crossing, the two exponentials are very similar (left), but as $N$ grows, so does the gap between them (right).
Hamiltonian cycles (i.e., $1.820 \times 10^{55}$) but $C_{1080}D_{6d}$ has

$$N_{HC}(C_{1020}D_{6d})=18,250,747,738,371,979,597,388,745,176,681,058,911,549,835,175,634,326,272$$

(41) (i.e., $1.825 \times 10^{55}$). After that, the $D_{6+}$ series takes over forever (but may be overtaken by other series), as seen in Figure 8.

4 | PROOF OF THEOREM 1

Proof. (i) Consider a Hamilton cycle admitted by $G_k = A \oplus L_1 \oplus \cdots \oplus L_k \oplus Z$ represented through Lemma 1 as a face shading $S$. We can decompose $S$ into shadings $S_1, \cdots, S_k$ represented in the subgraphs components, $S = S_1 \oplus S_2 \oplus \cdots \oplus S_k \oplus Z$.

Each shaded subgraph corresponds to a unique node in the DFA connected through a path from start to end. We prove the property by induction. For notational convenience, write $S_0 = S_A, S_{k+1} = S_Z$, and $S_i = S_{L_i}$ for $i = 1, \cdots, k$.

Since by Lemma 1, $S$ fulfills property (a)–(c), so must also $S_A, S_Z$, and $S_1, \cdots, S_k$ (where, for property (c), we consider two faces connected in a sub-component if they are either connected within that component, or determined to be so from the connectedness-label). In particular, $S_A = S_0$ and $S_Z = S_{k+1}$ are nodes in the DFA by Step I.1.

We show that the appropriately connectedness-labeled shadings $S_1, \cdots, S_k$ are nodes in the automaton, and $S_0 \to S_1 \to \cdots \to S_{k+1}$ is a path over it. This is done by induction over the length $i$ of the sub-path $S_0 \to S_i$.

$i = 0$: The node for $S_i$ is created in Step I.1 if $k = 0$, or R.1 in the first iteration if $S_i$ is an internal layer. In both cases, the arc $S_0 \to S_1$ is created in the first iteration of Step R.1.

$i \Rightarrow i+1$: Assume now for $1 \leq i \leq k$ that the path $S_0 \to S_1 \to \cdots \to S_i$ exists. Then, at some step $j \leq i$, the connectedness-labeled shading $S_j$ is extracted in Step R.1, and since $S_i \oplus S_{i+1}$ must fulfill L1 (a)–(c) (or $S$ would not be a Hamilton cycle), the appropriately connectedness-labeled $S_{i+1}$ is added together with the arc $S_i \to S_{i+1}$ at latest in Step R.3 of iteration $j \leq i$.

The caps are not connectedness-labeled, but are not needed to be, as this information is only needed to maintain Lemma 1(c) through Lemma 2 in the induction step.

(ii) We show that every path beginning in an initial state and ends in a final state corresponds to a Hamilton cycle, and the product of the arc labels along the path is the number of Hamilton cycles that correspond to this path through (i).

Consider such a path $S_0 \to \cdots \to S_{k+1}$. By construction, a node for the connectedness-labeled shading $S_i$ is created if either $S_i$ is a cap-shading, or if $S_{i-1} \oplus S_i$ fulfills property (a) and (b), and does prevent (c) in the future. By Lemma 2, this then extends to all of $S_0 \oplus S_1 \oplus \cdots \oplus S_i$. The final arc to the end-cap $S_k \to S_{k+1}$ is placed if and only if it closes correctly, so that $S = S_0 \oplus S_1 \oplus \cdots \oplus S_k \oplus S_{k+1}$ fulfills all of (a), (b), and (c). Thus, $S$ represents a Hamilton cycle by Lemma 2.

Since at each step, the arc is labeled with the number of equivalent ways to orient the shading $S_{i+1}$, the product yields the total number of Hamilton cycles that are represented by this path.

(iii) By the above, every Hamilton cycle maps uniquely to a path $S_0 \to S_1 \to \cdots \to S_{k+1}$ starting in an initial state $S_0$ and ending in a final state $S_{k+1}$ such that the product $m_{01}m_{12} \cdots m_{k,k+1}$ is the number of Hamilton cycles that map to this same path. The total number of Hamilton cycles admitted by $G_k$ is then the sum of these products over all such paths of length $k$. Letting $M$ be the adjacency matrix of the state machine $M$ with $m_{st}$ nonzero whenever $s \to^e t$ is an arc in $M$, this is exactly calculated by

$$N_{HC}(G_k) = \sum_{s \in \text{Initial}} \sum_{t \in \text{Final}} N_{HC}(G_k)$$

(42)

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CONFLICT OF INTEREST

This work does not have any conflicts of interest.

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