

Geometric and Topological Approaches to Crystallography

—Connections between Symmetry, Lattices, Crystalline Structures

Ellie Richwine, Lucian M. Ionescu

Department of Mathematics, Illinois State University, Normal, IL, USA

Email: errichw@ilstu.edu, lmiones@ilstu.edu

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Abstract

This article explores the mathematical structures underpinning crystalline materials, bridging the gap between pure mathematics and materials science. To clarify the scope of this work, the paper is divided into a tutorial exposition and an original theoretical contribution. The tutorial components provide a rigorous yet accessible introduction to the geometric and topological modeling of crystals, building upon Toshikazu Sunada's breakthrough framework of topological crystallography and subsequent formalizations by John C. Baez. We examine polyhedral geometry, duality, and lattice arrangements such as the Eisenstein and triangular lattices, framing them within the context of covering maps and Abel-Jacobi maps. Our original mathematical contribution is presented in Section 11, where we advance this foundation by introducing a simplified formulation of Graph Cohomology. The precise novelty of this section lies in its use of short exact sequences of graphs to categorify the connectivity of crystalline networks. By applying integer cohomology to quotient maps, this approach provides a unifying architectural template capable of rigorously tracking lattice defects as non-trivial cohomology classes. We conclude by discussing the broader heuristic applications of these tools in molecular biology, theoretical physics, and fault-tolerant quantum engineering.

Keywords

Topological Crystallography, Graph Cohomology, Bravais Lattices, Abel-Jacobi Maps, Materials Science, Crystal Networks

1. Introduction

This report examines mathematical structures that underpin crystalline materials with a focus on polyhedral geometry, lattice arrangements, and symmetry. The study

is based primarily on guidance from the second author and insights from a key advancement of the topic by Toshikazu Sunada on crystallography [1] [2]. The article responds at the same time to the “plea for more awareness of the work by mathematicians and materials scientists” [3], whose authors further “beg mathematicians to make their work more accessible to physical scientists (e.g. by including pictures such as the beautiful one presented by Sunada)” (loc. cit., p. 8000), being well aware of the importance of the breakthrough work of Sunada, bridging crystallography and Material Sciences, and the modern mathematical framework offered by Homological Algebra and Algebraic-Geometry.

Regarding this later goal of our article, the informal and intuitive statements are aimed towards a larger audience consisting of graduate students and Post-Doc students, in case their advisors would choose this exciting topic as a subject to be developed by their students. This matches the tandem structure of the authorship of the article. The well-seasoned researcher, most likely, will skip them for the more technical part of our tutorial synthesis of Toshikazu Sunada’s work, further detailed by John Baez’s article cited in the bibliographic section, together with our own novel mathematical framework through the addition of Graph Cohomology theory (a secondary goal of our article). The latter is an extension of Sunada’s Homological Algebra and AG-approach to the intrinsic level of graphs, before placements in Euclidean spaces.

In addition to the contributions to the formalism and rigor from [4], we will point towards the developments by the second author of Graph Cohomology [5] [6], which already led to advancements in Quantum Field Theory, in connection with the understanding of renormalization, pioneered by A. Connes and D. Kreimer [7] and notable results like the proof of the Formality Conjecture by M. Kontsevich [8] [9].

A full development of the work in [1], with contributions from [4], in the light of Graph Cohomology, is beyond the scope of this article, and will only be highlighted for other mathematicians to be aware of, and hopefully further developed as an application to the theory of building blocks of periodic crystals, by Sunada.

By connecting theoretical concepts to concrete crystalline structures, the project aims to develop a deeper understanding of how geometry and topology describe the organization of matter. As a follow-up project, application of Graph Cohomology [5] is expected to extend the results already obtained by Sunada, and provide a unifying framework for further applications (see 14).

This report investigates Platonic and Archimedean solids, dual graphs of space-filling polyhedra, and lattice arrangements such as the Eisenstein and triangular lattices. These structures illustrate how symmetry and space-filling constraints give rise to highly ordered arrangements in crystals. Additionally, homological concepts are discussed to provide an algebraic perspective on periodic structures and global connectivity. One of the goals is to emphasize the interplay between mathematical theory and crystallographic application, demonstrating how abstract structures can inform our understanding of real-world materials. Supple-

mentary material is included at the end of the article to maintain clarity and focus in the main discussion.

To directly address the scope and intent of this manuscript: Sections 2 through 10 serve as a tutorial exposition. They function to review, synthesize, and visually illustrate established mathematical frameworks—ranging from dual graphs to Abel-Jacobi maps—making them accessible to a broader scientific audience. Section 11 constitutes our original mathematical contribution. Here, we introduce a novel, simplified formulation of Graph Cohomology tailored for crystallography, utilizing short exact sequences of graphs to formalize the internal topology of fundamental building blocks. Finally, Sections 12 and 13 provide a forward-looking, heuristic discussion on how these homological tools might be applied to advanced topics like time crystals and quantum engineering. To cater to our diverse readership, we have structured the exposition to balance heuristic physical interpretations with formal mathematical rigor. For the physical scientist, the visual and conceptual descriptions provide a bridge to topological methods. For the mathematically inclined reader, we have included formal definitions and theorems summarizing the foundational results of Sunada [1] [2] and Baez [4] where appropriate. The full technical development of our novel Graph Cohomology framework, complete with specific functional setups and boundary operators, is explicitly developed in Section 11.

We proceed in this first part of the article with a leisurely presentation of the work of Sunada, extracting some key aspects from *Lectures on Topological Crystallography* [1].

Before considering periodic graphs, a preliminary recall of the history of geometric patterns relevant to crystal structures includes, of course, Platonic and Archimedean finite geometries in the sense of Felix Klein: “An abstract geometry is a group acting on a space”.

2. Polyhedra

A significant portion of crystallographic structure is described through the language of polyhedra and three-dimensional solids. A natural starting point is the set of Platonic solids, **Figure 1**, which are perfectly regular polyhedra composed of exactly one type of congruent polygonal face, with identical face arrangements at every vertex. There exist exactly five such solids, distinguished by their exceptional symmetry. An important feature of Platonic solids is their duality, where vertices and faces may be interchanged to produce another Platonic solid. While this concept will be developed more in later sections, it is worth noting here that duality emerges at the most fundamental level of polyhedral geometry and plays a recurring role in crystallographic descriptions.

Extending beyond this idealized class leads to the *Archimedean polyhedra* (**Figure 2**), a collection of thirteen convex solids constructed from regular polygonal faces but allowing more than one polygon type within a single polyhedron. Unlike Platonic solids, Archimedean polyhedra are not face-transitive, but they retain

vertex transitivity and exhibit high degrees of symmetry. This distinction makes them particularly relevant to the modeling of naturally occurring crystal structures, where perfect regularity is often relaxed in favor of geometries that better accommodate spatial constraints.

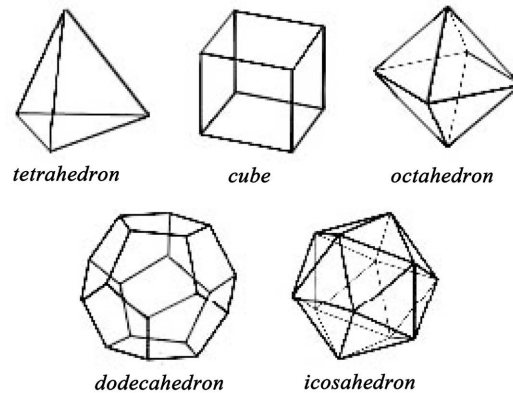


Figure 1. The Platonic solids.

Archimedean solids

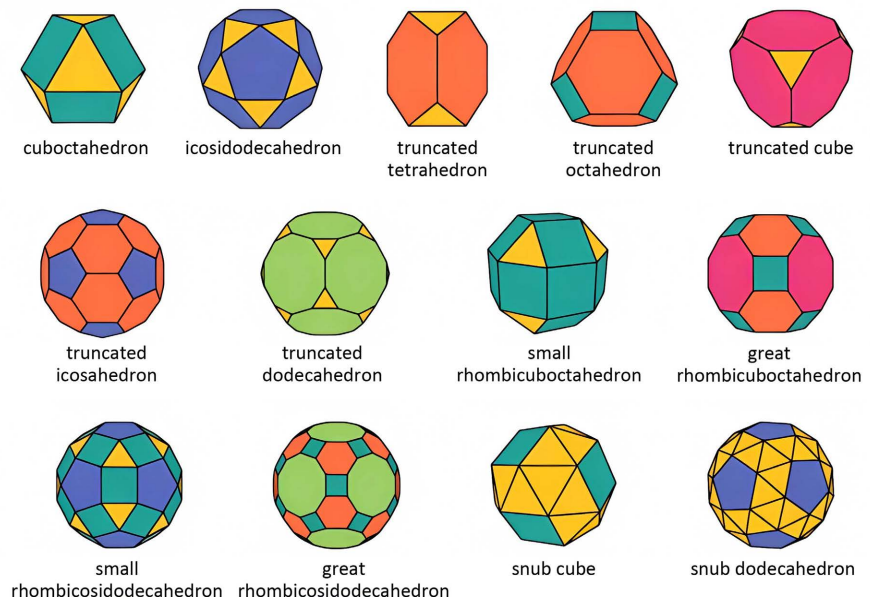


Figure 2. The Archimedean solids.

Among these solids, the truncated octahedron is of special importance, as it tiles three-dimensional Euclidean space without gaps or overlaps. This property aligns directly with the defining characteristics of crystalline materials as periodic, space-filling lattices.

The significance of Platonic and Archimedean solids, therefore, extends beyond the classification of geometric forms. Together, they contribute to a conceptual framework that links abstract geometry with physical and natural organization. Through these polyhedra, principles such as symmetry, packing efficiency, and

energetic optimization can be expressed in precise geometric terms, providing a bridge between mathematical idealization and the structure regularities observed in crystalline matter.

3. Duality and Dual Graphs

Duality in polyhedral geometry provides a natural extension of the symmetry principles introduced by Platonic and Archimedean solids. Given a convex polyhedron, its dual is constructed by interchanging vertices and faces while preserving edge connectivity. This operation produces a new polyhedron whose combinatorial structure reflects that of the original yet emphasizes complementary geometric features. In the case of Platonic solids, duality maps each solid to another Platonic solid, reinforcing the fundamental role of symmetry at this level of idealized geometry.

Beyond its geometric definition, duality offers a powerful interpretive lens for crystal structures. Polyhedra that emphasize vertex coordination may be transformed, through duality, into solids that instead describe spatial partitioning. This shift perspective is particularly relevant in crystallography, where both atomic coordination environments and the division of space into repeating regions are essential for understanding structure. As later sections will demonstrate, this dual relationship closely parallels constructions such as Voronoi tessellations and Wigner-Seitz cells, which arise naturally in periodic lattices.

Duality functions in this way not merely as a formal operation but as a conceptual bridge between local and global descriptions of order. By allowing structural information to be viewed from complementary geometric perspectives, it prepares the groundwork for interpreting how ideal polyhedral symmetries are adapted within real space-filling systems.

When this notion of duality is extended from isolated polyhedra to space-filling systems, it provides a particularly transparent way to analyze honeycomb structures. A honeycomb (Figure 3) may be viewed as a tiling of space by congruent polyhedral cells, and its dual graph is obtained by placing a vertex at the center of each cell and connecting vertices whose cells share a common face (Figure 4). In this construction, we see that it forms a partition of a network that encodes its connectivity.

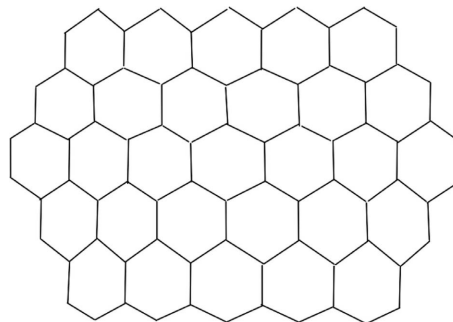


Figure 3. Original honeycomb tree graph.

The resulting dual graph captures how space is organized at a global level while remaining directly linked to the local geometry of each cell.

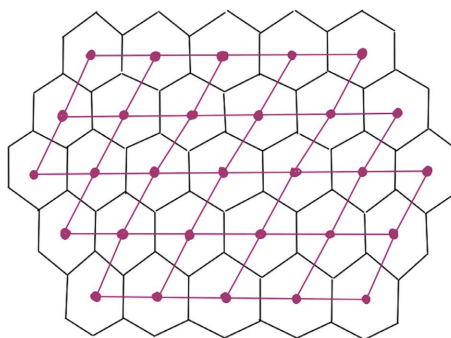


Figure 4. Honeycomb with dual graph overlay.

Framed this way, the dual of a honeycomb (compare **Figure 4** and **Figure 5**) serves as an intermediate representation between full geometric realization and abstract lattice models, making it especially useful for interpreting coordination, transport pathways, and symmetry in periodic structures. Upon closer examination of the dual graph associated with the honeycomb, a familiar structure emerges.

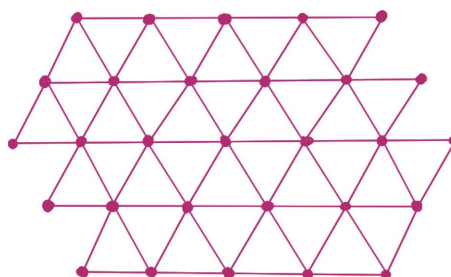


Figure 5. Eisenstein lattice.

The dual of this honeycomb realizes an Eisenstein lattice, a two-dimensional triangular lattice embedded in three-dimensional space, characterized by its angular relationships and uniform coordination. This identification reveals that the connectivity of the honeycomb encodes a highly symmetric lattice arrangement, linking local face-sharing relations to a well-known crystalline geometry. The implications of this correspondence for crystalline structures and lattice optimality will be discussed in greater detail in a later section.

4. Deck Transformations and Fundamental Groups

Deck transformations provide an essential algebraic viewpoint on the symmetries of covering spaces, extending the duality principles discussed above into the realm of topology. Given a covering map $p: \tilde{X} \rightarrow X$, a deck transformation is a homeomorphism of the covering space that preserves fibers, meaning $p \circ g = p$ for

every such transformation g . The collection of all deck transformations forms a group under composition, and this group encodes how local geometric information lifts to global periodic structure.

When the covering space is universal, the deck transformation group is naturally isomorphic to the fundamental group $\pi_1(X)$. This gives a geometric interpretation of loops in the base space, traversing a non-trivial loop lifts to a path in the covering space whose endpoint differs from the starting point by a deck transformation. So, homotopy classes of loops become concrete symmetries acting on the lifted space [7].

In the context of crystalline materials, the covering space often corresponds to an idealized infinite lattice, while the base space represents a fundamental domain or quotient by translational symmetry. The fundamental group $\pi_1(X)$ acts naturally on the covering space, and in many cases, the deck transformation group is isomorphic to a quotient of $\pi_1(X)$. This correspondence provides a rigorous way to interpret lattice periodicity as arising from the algebraic structure of loops in the quotient space.

For example, a two-dimensional Bravais lattice may be modeled as $\mathbb{R}^2/\Lambda$, where Λ is a lattice of translations. The quotient is topologically a torus, whose fundamental group is $\mathbb{Z}^2$. Each generator corresponds to translation along one independent lattice direction. In three dimensions, the analogous quotient yields a 3-torus with fundamental group $\mathbb{Z}^3$, matching the three translational periods of a crystal lattice.

For honeycomb structures and their dual lattices, deck transformations describe how repeating polyhedral cells tile in space, while the fundamental group captures the global constraints imposed by periodicity. This interplay between algebraic and geometric viewpoints prepares the groundwork for later discussions of Abel-Jacobi maps and homological invariants in periodic systems.

5. The Eisenstein Lattice and Its Role in Crystalline Structure

The Eisenstein lattice occupies a central position in both mathematics and the theory of crystalline order, representing one of the most symmetric two-dimensional lattice arrangements. Geometrically, it is realized (as aforementioned) as a triangular lattice in which each lattice point has six nearest neighbors arranged at 60° intervals. This configuration yields the highest possible rotational symmetry and packing efficiency among planar Bravais lattices, properties that make it a natural organizing principle for ordered systems (Figure 6).

Before embedding them geometrically in Euclidean space, these lattices exist intrinsically as algebraic structures pertaining to Number Field extensions.

From an algebraic perspective, the Eisenstein lattice arises from the ring of Eisenstein integers $\mathbb{Z}[\omega]$, where $\omega = e^{\frac{2\pi i}{3}}$. In this formulation, lattice points correspond to integer linear combinations of two basis vectors separated by 60° ,

linking geometric symmetry with complex multiplication and number-theoretic structure.

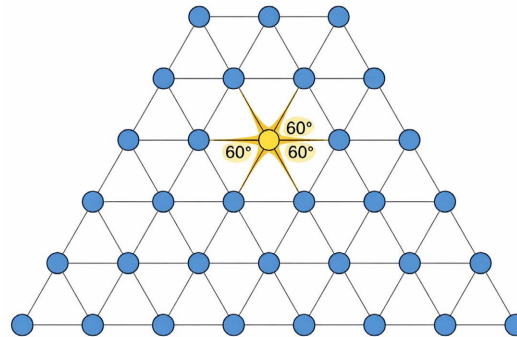


Figure 6. Rotational symmetry in an Eisenstein lattice.

Similarly, the square lattice arises from the Gaussian integers $\mathbb{Z}[i]$. The compositum of these fields yields a Galois extension containing both $\mathbb{Z}[i]$ and $\mathbb{Z}[\omega]$, revealing how pure Algebraic Number Theory dictates permissible symmetries. At this stage, Minkowski's Lattice Point Theorem from the Geometry of Numbers becomes fundamentally important. It provides the rigorous bridge between the algebraic volume of these integer rings and the geometric existence of lattice points within convex bodies, ultimately governing how these algebraic structures can be physically packed or embedded in $\mathbb{R}^n$.

This dual geometric-algebraic characterization underlies the lattice's prominence in areas ranging from complex analysis and modular forms to sphere packing and coding theory.

In crystallography, the Eisenstein lattice is significant because it provides an optimal framework for uniform coordination and efficient space utilization. When a honeycomb structure admits a dual graph that realizes an Eisenstein lattice (See **Figure 4**), the local face-sharing relationships of the cells give rise to a globally ordered triangular network.

This correspondence clarifies how highly symmetric lattices can emerge naturally from space-filling polyhedral arrangements, rather than being imposed as abstract models.

Interpreted in this way, the Eisenstein lattice provides a unifying geometric framework that links space-filling polyhedral arrangements with the algebraic description of crystalline order. Its emergence as the dual graph of a honeycomb structure illustrates how local geometric constraints can propagate into globally optimal and highly symmetric lattice configurations. This perspective underscores the role of duality in revealing how ideal mathematical lattices are naturally realized within physical crystal structures.

6. Topological and Geometric Properties of the Triangular Lattice

To prevent the conflation of related but distinct objects, we must explicitly distin-

guish the structures discussed thus far, specifying their dimensionality. The hexagonal honeycomb is a purely planar, two-dimensional graph; however, it is not a true Bravais lattice because the local environment of its nodes alternates. Its dual, the triangular lattice, is a genuine two-dimensional planar Bravais lattice. The Eisenstein lattice, as discussed above, is the specific, optimal algebraic realization of this triangular lattice embedded in the complex plane. When materials like graphite are modeled, these purely planar 2D sheets are then embedded and stacked within three-dimensional (3D) Euclidean space.

The triangular lattice occupies a distinguished position among two-dimensional periodic networks due to its simultaneous optimality in topology, packing efficiency, and connectivity. These properties jointly explain its frequent emergence as the dual structure of hexagonal honeycombs and related cellular systems.

From a graph-theoretic perspective, the triangular lattice is maximally planar, meaning it saturates the upper bound on the number of edges permissible for a planar graph at a fixed node number. Every face is also a three-cycle, the shortest possible closed loop in a planar embedding, which enforces strong local constraints and eliminates internal degrees of freedom. As a result, the lattice is globally rigid under generic conditions with no floppy modes arising from topology alone. Short cycle length also minimizes ambiguity in force balance and transport pathways, yielding unique and stable solutions to equilibrium and flow problems.

Geometrically, the triangular lattice (**Figure 7**) realizes the densest possible packing of congruent disks into two dimensions. This property implies optimal use of space and near-isotropic local environments, as each node is surrounded symmetrically by its neighbors. In physical systems governed by short-range interactions or exclusion restraints, such isotropy minimizes directional bias and leads to uniform stress, transport, or interaction fields. Consequently, the triangular lattice often corresponds to a minimum-energy configuration under fixed density or spacing constraints.

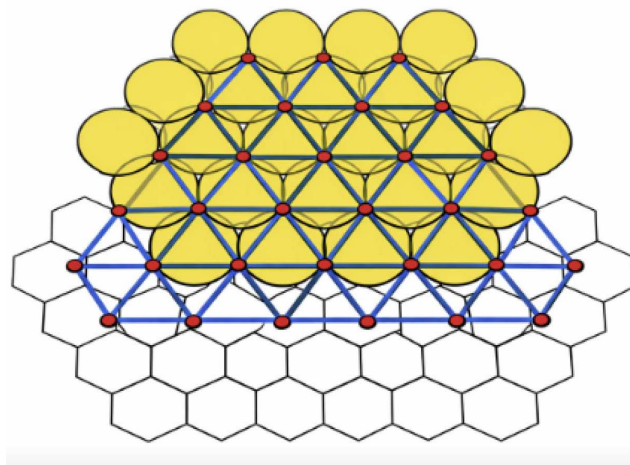


Figure 7. Combined illustration of a hexagonal honeycomb lattice, its dual triangular lattice, and an optimal circle packing.

For a planar lattice with uniform edge length, the triangular lattice achieves the highest possible coordination number, with each node connected to six nearest neighbors. This maximal connectivity enhances redundancy and robustness, distributing loads, flows, or signals over multiple independent pathways. High coordination also reduces characteristic path lengths and suppresses localization phenomena, improving mechanical stiffness and transport efficiency relative to square or hexagonal lattices.

Taken together, these properties render the triangular lattice an optimal carrier of constraints in two dimensions. When a system separates spatial efficiency from mechanical or topological constraints, such as in hexagonal honeycombs, the dual triangular lattice naturally emerges as the structure that maximizes rigidity, isotropy, and robustness while remaining planar.

7. Bravais Lattices

In the language of mathematics and solid-state theory, a Bravais lattice is defined as an infinite, periodic arrangement of discrete points in Euclidean space such that each point possesses an identical local environment (**Figure 8**). Formally, a Bravais lattice consists of all integer linear combinations of a set of linearly independent basis vectors.

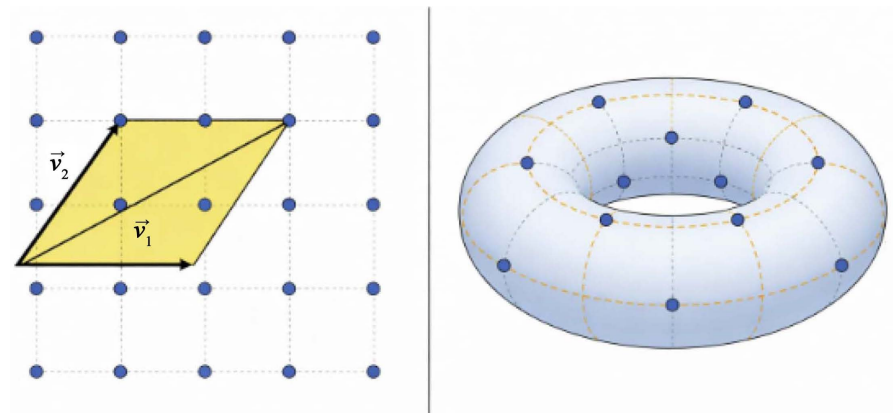


Figure 8. Two views of a 2d Bravais lattice. Left: representation as a periodic lattice. Right: the same lattice as a torus by identifying opposite edges of the fundamental domain.

This structure ensures that the lattice is invariant under discrete translations, meaning that if the entire configuration is shifted by a lattice vector, it coincides perfectly with itself. In other words, the symmetry of the lattice is purely translational, and every lattice point is indistinguishable from any other when viewed within the global structure.

This concept plays a foundational role in crystallography and materials science. In three-dimensional space, Bravais demonstrated that there are exactly 14 distinct lattice types that classify all possible periodic crystal structures. These lattices provide the framework on which crystals are built, allowing one to describe atomic arrangements in terms of repeating unit cells. By reducing a complex solid to a

repeating geometric pattern, the Bravais lattice offers a precise mathematical model for understanding symmetry, periodicity, and spatial organization in crystalline materials.

A particularly important example relevant to this discussion is the Eisenstein lattice, as previously mentioned. It arises from the ring of the complex numbers of the form $a + b\omega$, where a and b are integers and ω is a primitive cube root of unity. It is important to this structure because of its high degree of symmetry and optimal packing properties.

Understanding Bravais lattices is essential for interpreting crystal maps, as these maps encode the translational symmetries that define crystalline order. Moreover, the lattice structure provides the algebraic and geometric foundation necessary for extending the discussion into homological concepts. When studying homology in a lattice context, one examines how cycles, boundaries, and higher-dimensional analogues behave within periodic spaces. The regularity and translational invariance of Bravais lattices make them particularly well-suited for such topological analysis, serving as a bridge between geometric symmetry and algebraic topology.

8. Homology and Periodic Structure

Homology provides the algebraic infrastructure needed to pass from geometric intuition to a rigorous description of structure within periodic systems. While the preceding sections have emphasized symmetry, duality, and lattice optimality, homology works to address a complementary question: how do cycles, connectivity, and global constraints behave within repeating spaces? In the context of crystallography and lattice models, this perspective is essential for understanding how local configurations propagate through an infinite periodic structure.

At its core, homology studies the relationship between cycles and boundaries in space. Informally, a cycle is a closed configuration, such as a loop in a graph or a closed surface in three dimensions, while a boundary is a cycle that arises as the edge of a higher-dimensional object. Homology distinguishes between cycles that are merely boundaries and those that represent genuine topological features of the space. The resulting homology groups measure the presence of “holes” in various dimensions and remain invariant under continuous deformations.

When applied to lattice graphs, this theory becomes especially transparent. A periodic lattice may be viewed as an infinite graph equipped with translational symmetry. The translational subgroup acts on the graph, producing a natural quotient. The infinite periodic structure can be reduced to a finite fundamental domain together with algebraic data describing how edges wrap around the domain. In this setting, the infinite graph serves as a covering space, while the finite quotient graph encodes its essential combinatorial structure.

To formalize this mathematically, we rely on the precise definitions established by Sunada [1] and expanded by Baez [4]:

Definition 1 (Topological Crystal [1]). *A topological crystal is defined as a pair (X, Λ) , where X is an infinite, connected topological graph, and Λ is a free*

abelian group of finite rank acting freely on X by graph automorphisms, such that the quotient graph X/Λ is a finite graph.

We invite the reader to browse Section 7 from loc.cit., which contains the fundamental concepts regarding the graph theory and algebraic topology aspects of Sunada's approach to crystallography (e.g., covering spaces).

Proposition 1 (Periodic Realization [4]). *If (X, Λ) is a topological crystal, the finite quotient graph X/Λ serves as the fundamental building block. The infinite crystal X can be uniquely reconstructed as a regular covering space of X/Λ with deck transformation group Λ .*

A classical illustration arises from considering the infinite cyclic group $\mathbb{Z}$ acting by translation. If one restricts attention to translations by multiple of 6, the subgroup $6\mathbb{Z}$ defines a periodic identification. The quotient group $\mathbb{Z}/6\mathbb{Z}$ then captures the structure of a sixfold cyclic system. Graphically, an infinite linear lattice can be “folded” into a finite cycle of length six. You can envision this as the image below displays, a rope that is coiling over itself in loops of length 6 (**Figure 9**). The homological structure detects the persistence of the fundamental loop in this quotient, although locally the graph appears linear, globally it contains a non-trivial one-dimensional cycle.

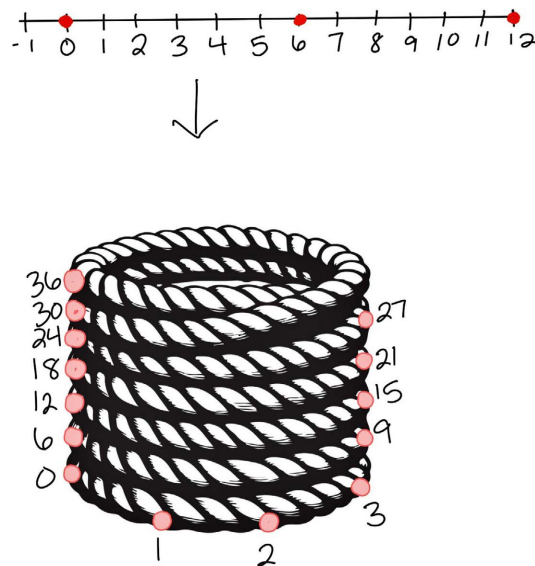


Figure 9. Illustration of the wrapping of $6\mathbb{Z}$.

This subobject-extension-quotient viewpoint generalizes naturally to high-dimensional crystal lattices. A Bravais lattice in three-dimensional space may be interpreted as a free abelian group of rank three acting on Euclidean space by translations. Passing to a quotient by a full-rank sublattice yields a compact torus-like structure, whose first homology group will reflect the independent translational cycles of the lattice. In this way, homology provides an algebraic record of periodicity itself.

The importance of this perspective for crystallography lies in modeling. Crystal

maps encode local adjacency and coordination data, but when embedded into three-dimensional space, they inherit global topological constraints. Homology formalizes how these local connections assemble into extended networks and identifies which cycles represent genuine structural features rather than artifacts of a particular embedding.

From a broader viewpoint, homology serves as the bridge between geometric symmetry and algebraic topology. While group theory classifies the allowable symmetries of a lattice and duality clarifies complementary geometric descriptions, homology captures the invariant connectivity of the resulting periodic network. Together, these tools help to provide the conceptual building blocks in a tangible way for modeling crystalline materials in a three-dimensional space, ensuring both local coordination and global structure are treated within a unified mathematical architecture.

9. Abel-Jacobi Maps and Crystalline Periodicity

The Abel-Jacobi map provides a bridge between homology and geometric realization by embedding cycles of a graph or lattice into a torus constructed from its first homology group. For a periodic lattice graph, the first homology group captures independent cycles that persist under translational symmetry. The Abel-Jacobi map assigns to each vertex a point in the torus, effectively encoding the cumulative displacement along fundamental cycles.

In classical algebraic geometry, the Abel-Jacobi map sends divisors of degree zero on an algebraic curve into its Jacobian variety, where path integrals of holomorphic differentials record their accumulated displacement. In discrete graph theory, analogous constructions exist for metric graphs and periodic networks, where combinatorial cycles replace continuous curves [10].

To define this rigorously for discrete networks, we utilize the standard homological setup for graphs:

Definition 2 (Discrete Abel-Jacobi Map [2]). *Let $Y = X/\Lambda$ be the finite quotient graph of a topological crystal. The discrete Abel-Jacobi map $\Phi: Y \rightarrow J(Y)$ embeds the graph into its Jacobian torus, defined as $J(Y) = H_1(Y, \mathbb{R})/H_1(Y, \mathbb{Z})$. The map assigns to each vertex a position in the torus by integrating a canonical basis of harmonic 1-forms along paths in the graph.*

As mentioned at the end of the Section 1 Introduction, the Algebraic-Geometric framework for studying the properties of such mathematical models for crystals require more advanced concepts of Algebraic-Geometry, yet provide a direct path to understand their quantum properties, as relating to a *finite Feynman Path Integrals* formalism, providing quantum amplitudes in a gauge theory interpretation on top of the mathematical model, periods related to their material science properties¹.

Theorem 1 (Standard Realization [1]). *The Abel-Jacobi map lifts to a Λ -equivariant map $\tilde{\Phi}: X \rightarrow H_1(Y, \mathbb{R})$, providing a canonical geometric realization of the*

¹We foresee applications in several modern areas of research, e.g. Spintronics, Magnetronics etc. relevant to new QC hardware solutions.

abstract topological crystal X into a Euclidean space. This realization uniquely minimizes the energy of the periodic graph.

In crystallographic settings, this construction yields a canonical way to represent the “phase” of a point within the periodic structure. The resulting torus parametrizes the global degrees of freedom of the lattice, while the embedding reflects how local connectivity propagates through the entire crystal.

This framework is especially useful in the presence of defects or quasi-periodicity (see **Figure 10**). Dislocations and vacancies may appear locally minor while producing non-trivial winding behavior globally. The Abel-Jacobi image records these cumulative offsets by measuring how paths fail to close trivially in homology. Consequently, it provides a rigorous tool for studying transport channels, phase shifts, and topological obstructions in periodic media.

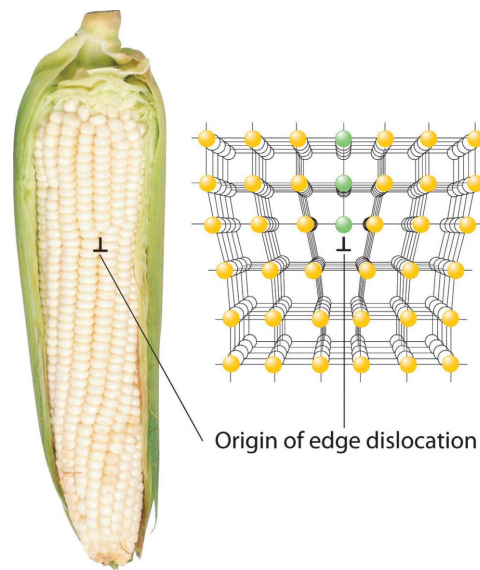


Figure 10. Crystal deformation example using the Abel-Jacobi map.

This perspective aligns naturally with the interpretation of Bravais lattices as quotients of Euclidean space by a discrete translation group, and it provides a powerful tool for analyzing defects, quasi-periodicity, and transport phenomena.

10. Covering Maps and Crystalline Symmetry

Covering maps arise naturally in the study of periodic structures because they formalize the relationship between an infinite lattice and its finite fundamental domain. A covering map allows one to treat the infinite periodic structure as a geometric unfolding of a compact quotient. In crystallography, this quotient often corresponds to a unit cell or a toroidal representation of the lattice.

More precisely, a covering map is locally a *homeomorphism*, meaning every small neighborhood in the quotient lifts to disjoint copies in the covering space (**Figure 11**). This models the physical principle that each unit cell of an ideal crystal has the same local environment as every other unit cell.

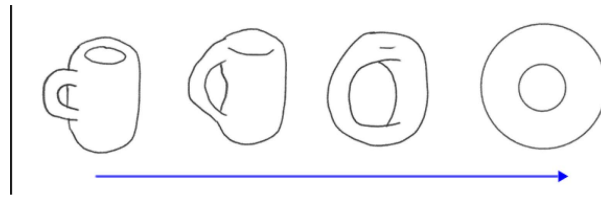


Figure 11. Classic homeomorphism example.

This viewpoint clarifies how local geometric constraints, such as coordination number or face-sharing relations, extend globally through repeated application of deck transformations.

For translationally periodic materials, Euclidean space serves as the universal cover, while the quotient by the translation lattice yields a torus. In this way, standard crystallographic concepts such as primitive cells, lattice vectors, and repeated coordination environments admit a natural topological interpretation [11] [12].

Moreover, covering maps provide a natural setting for analyzing defects: a dislocation, for example, corresponds to a failure of the covering structure to remain globally consistent, reflected algebraically in a non-trivial element of the fundamental group.

Physically, a circuit around a dislocation core may fail to return to the original lattice position, producing a Burgers vector. Topologically, this corresponds to non-trivial monodromy in the covering structure, linking classical defect theory with algebraic topology [12].

This topological framework requires a careful distinction of dimensionality. The infinite periodic network, acting as the universal cover, is analyzed as a purely planar (2D) or spatial (3D) structure. Conversely, the fundamental building block—the quotient—“lives” on a compact Riemann Surface (such as a 2-torus for a 2D planar lattice). The finite quotient graph embedded on this surface acts as a topological skeleton. This deep connection between graphs embedded on Riemann Surfaces and algebraic curves directly parallels Grothendieck’s theory of *Dessins d’Enfants*, hinting at profound, yet underexplored, applications of Algebraic Geometry in classifying the fundamental building blocks of matter [13].

The use of homological algebra techniques in Topological Crystallography invites an investigation into its relation with *Graph Cohomology*. Originally formulated as an intrinsic algebraic approach to Quantum Field Theory—notably for understanding renormalization, as pioneered by A. Connes and D. Kreimer [7]—where *Graph Cohomology* abstracts the combinatorial properties of networks.

Rather than relying heavily on the advanced machinery of Differential Graded Lie Algebras (DGLAs) used in the proof of the Formality Conjecture [9], we can distill the essence of this theory using fundamental concepts from Category Theory. By defining “extensions” of graphs, the second author [5] provided a framework where graphs themselves can be treated as algebraic objects subject to the rules of homological algebra.

11. Highlights of Graph Cohomology

In mathematics, complex objects are frequently understood by breaking them down into simpler components via the concept of an *extension*, much like the theory of group extensions. This foundational idea can be implemented in the category of graphs and graph morphisms by adapting the concept of a *short exact sequence*.

11.1. Category of Graphs and Exact Sequences

To elevate the heuristic description to a rigorous homological framework, we must first specify the category of graphs we are working within, as well as the precise nature of its morphisms [5].

Definition 3 (Category of Graphs for Cohomology). *Let **Graph** denote the category whose objects are finite, directed graphs (with multiple edges and loops permitted). A morphism $f : G \rightarrow H$ in **Graph** consists of a pair of maps $f_V : V(G) \rightarrow V(H)$ and $f_E : E(G) \rightarrow E(H) \cup \{*\}$, where V and E denote the vertex and edge sets respectively, and $*$ denotes a contracted state. The maps must preserve incidence relations: if an edge e connects v_1 to v_2 in G , then $f_E(e)$ either connects $f_V(v_1)$ to $f_V(v_2)$ in H , or $f_E(e) = *$ implying that $f_V(v_1) = f_V(v_2)$.*

This allowance for edge contraction is the critical feature that permits quotient operations within the category.

Definition 4 (Monomorphisms and Epimorphisms). *Within this category of graphs:*

- A **monomorphism** $i : G_1 \rightarrow G_2$ is an injective graph homomorphism corresponding to a strict subgraph inclusion. No edges or vertices are contracted ($i_E(e) \neq *$ for all $e \in E(G_1)$).
- An **epimorphism** $q : G_2 \rightarrow G_3$ is a surjective graph homomorphism. In our context, this corresponds specifically to a graph contraction map, where a well-defined subgraph $G_1 \subset G_2$ is collapsed into a single vertex in G_3 , and all internal edges of G_1 are mapped to the contracted state $*$.

With these morphisms established, we can define the quotient graph, denoted G_2/G_1 , as the graph obtained by collapsing the entire subgraph G_1 into a single node. The edges in G_2 that previously connected a vertex outside G_1 to a vertex inside G_1 now connect directly to this new, collapsed node.

Definition 5 (Short Exact Sequence of Graphs [5]). *A sequence of graphs and morphisms*

$$0 \rightarrow G_1 \xrightarrow{i} G_2 \xrightarrow{q} G_2/G_1 \rightarrow 0 \quad (1)$$

is defined as a **short exact sequence** if i is a monomorphism (inclusion of the subgraph G_1), q is an epimorphism (contraction of G_1 to a single macroscopic node), and the image of i is exactly the pre-image of the contracted node under q . Here, 0 represents the trivial graph consisting of a single vertex and no edges.

From a physical or crystallographic perspective, this exact sequence mathemat-

ically models coarse-graining or block-spin transformations. The subgraph G_1 might represent a complex fundamental building block (such as a unit cell or a local geometric defect). The quotient G_2/G_1 represents the macroscopic lattice where the internal details of G_1 are ignored, and G_1 is treated simply as a point-like node in the broader network.

Conversely, the sequence can be read as a substitution process: substituting a single vertex in the macroscopic graph G_2/G_1 with a more complex internal structure G_1 yields the refined graph G_2 .

11.2. From Exact Sequences to Cohomology

Once graphs are organized into these short exact sequences, the standard machinery of *Homological Algebra* can be applied. We can define chain complexes where a boundary operator, typically denoted ∂ , acts on a graph by systematically collapsing its subgraphs.

If we denote C_n as the space spanned by graphs with a specific complexity rating (e.g., the number of independent cycles or edges), the boundary map $\partial: C_n \rightarrow C_{n-1}$ reduces complexity by quotienting out simple subgraphs. The fundamental property of any boundary operator, $\partial^2 = 0$, allows us to define *Graph Homology* as the kernel of ∂ modulo the image of ∂ . Passing to the dual vector spaces naturally yields *Graph Cohomology*.

In this simplified formulation, graph cohomology measures the topological invariants of how graphs can be nested and glued together. It categorizes the connectivity of the network, detecting structural “holes” not in a physical spatial embedding, but in the combinatorial space of graph substitutions.

11.3. Quotient Maps, Building Blocks, and Cohomology with Integer Coefficients

To fully appreciate the utility of graph cohomology in this setting, it is highly instructive to compare the exact sequences introduced above with the quotient maps utilized in Topological Crystallography.

In Sunada’s framework [1], a crystal is modeled as a periodic realization of an infinite graph X . The translational symmetries of the crystal are captured by a free abelian group Λ (typically isomorphic to $\mathbb{Z}^3$) that acts on X . The quotient space X/Λ yields a finite graph, which Sunada identifies as the “building block” or the topological unit cell of the crystal.

In real-world physical systems, the vertices of this building block are not identical; they represent distinct atomic species, and the edges represent specific chemical bonds. This chemical identity is mathematically formalized as a “coloring map” or “atomic labeling” on the quotient graph.

As elaborated in the work of Baez [4], who rigorously advanced Sunada’s formalism, this labeling process is intimately connected to the cohomology of graphs. Rather than treating atomic identities merely as qualitative names or colors, we can assign them integer values. By doing so, the coloring map on the vertices be-

comes a zero-cochain, an element of $C^0(X/\Lambda; \mathbb{Z})$, utilizing integer coefficients. Consequently, bond properties, directional shifts, and the “wrapping” of the lattice map naturally to the first cohomology group $H^1(X/\Lambda; \mathbb{Z})$.

Comparing this with the short exact sequences of graphs, we observe a powerful unifying theme. The exact sequence $0 \rightarrow G_1 \rightarrow G_2 \rightarrow G_2/G_1 \rightarrow 0$ models coarse-graining by collapsing a complex localized subgraph into a single node. Conversely, the Sunada-Baez quotient map “folds” an infinite periodic structure onto a finite topological summary.

In both approaches, passing to a quotient simplifies the raw geometry. Applying cohomology with integer coefficients allows us to systematically recover and track the rich, “colorful” internal structure—whether that structure represents the hidden internal topology of the collapsed subgraph G_1 , or the distinct atomic identities and bonding rules within the crystalline lattice.

12. Applications in Topological Crystallography and broader Fields

The cohomological framework introduced above extends far beyond the classification of ideal crystals. By treating lattice structures as topological complexes, we open up new pathways for both theoretical exploration and practical application, transforming abstract mathematical invariants into predictive physical tools.

12.1. Theoretical Advancements: A Unifying Framework

At the theoretical level, graph cohomology offers a rigorous language for bridging discrete microscopic data with macroscopic continuous phenomena. The exact sequences and quotient maps described previously do not just model geometric crystals; they provide a structural template that can be applied across scientific disciplines.

This mathematical architecture highlights profound unifying trends across physics, chemistry, and biology. In molecular biology, for example, complex macromolecules and aperiodic biological structures can be analyzed using the same homological principles that govern periodic crystals, a rapidly growing area of study often framed within persistent homology [14]. Furthermore, in theoretical physics, interpreting discrete networks through a cohomological lens aligns with modern paradigms where macroscopic space-time itself can be viewed as an emergent property of underlying discrete, quantum-computational networks [15].

By analyzing the homological properties of these lattices, one can also extract crucial spectral data. The spectral zeta functions associated with specific lattice operators are deeply tied to the underlying topology of the graph, dictating the physical resonances, thermodynamic stability, and phase transitions within the material [16].

12.2. Applied Advancements: Defect Analysis and Quantum Engineering

On the applied side, graph cohomology—particularly when utilizing integer coef-

ficients—provides an exceptionally precise tool for tracking lattice defects. Dislocations, impurities, or vacancies disrupt the perfect periodicity of a crystal. Rather than viewing these simply as geometric flaws, they can be rigorously quantified as non-trivial cohomology classes. This allows material scientists to mathematically calculate how localized defects will propagate and ultimately alter the global mechanical, thermal, or electronic pathways of the material.

Additionally, this topological framework is highly relevant to the rapidly advancing field of quantum engineering. As researchers design synthetic quantum circuits and robust architectures, the underlying hardware frequently relies on highly symmetric lattice arrangements. Applying these topological and cohomological tools allows for the identification of topologically protected states—quantum configurations that remain inherently stable against local perturbations due to their global network properties [17]. This intersection of abstract topology and structural crystallography is poised to be instrumental in the development of fault-tolerant quantum systems and advanced materials.

In this part, we point to other applications of Topological Crystallography, from geometry to dynamics, notably the concept of a time crystal, and use its discrete approach to suggest its uses in Quantum Computing, hardware, and software.

13. Time Crystals, Vanishing Homology, and Quantum Applications

Time crystals represent a remarkable extension of crystallographic ideas into the temporal domain. Unlike conventional crystals, which exhibit periodicity in space, time crystals display periodicity in time, breaking continuous time-translational symmetry into a discrete one.

Originally proposed by Frank Wilczek in 2012 [18], time crystals were later realized in driven quantum systems such as trapped ions and spin chains [19]. Their defining feature is persistent subharmonic oscillation: the system responds with a period larger than that of the external drive.

Building on the rigorous spatial formalisms established by Sunada and Baez [1] [4], we propose an educated intuition for extending these structures into the temporal domain. In this framework, oscillations and vibrations can be modeled as universal covers where time serves as a “labeling dimension,” entirely analogous to periodicity in physical space.

Mathematically, the stability of this temporal periodicity can be interpreted heuristically through the lens of vanishing homology groups associated with the system’s configuration space. When certain homology groups vanish, the system lacks continuous deformations that would otherwise eliminate temporal periodicity, thereby stabilizing discrete time evolution.

While vanishing homology is not yet standard terminology in condensed matter physics, it can be interpreted as the absence of deformation modes that would continuously unwind the periodic orbit.

In lattice models, time-crystalline behavior often emerges from Floquet systems,

where periodic driving induces an effective covering structure in time analogous to spatial coverings in crystalline materials. The “fundamental group in time” can be viewed as generated by one period of evolution, and the stability of the time crystal corresponds to non-trivial deck transformations acting on the space of states.

If the system evolves with period nT rather than the driving period T , the motion defines a non-trivial orbit under repeated temporal translation.

This perspective naturally aligns with emerging theoretical frameworks where relativistic time and space-time itself are not fundamental background arenas, but rather emergent properties arising from quantum phase, the algebraic-geometric Theory of Periods, and quantum computing formalisms [20] [21].

These ideas have direct implications for quantum computing. Time crystals offer potential pathways for constructing robust qubits whose states are protected by temporal symmetry rather than solely spatial or energetic barriers. The periodic structure in time can suppress decoherence by constraining the evolution of quantum states to a discrete set of symmetry-protected configurations².

On the hardware side, engineered quantum systems, such as trapped ions, superconducting qubits, and Rydberg arrays, provide platforms where time-crystalline phases have already been experimentally realized. On the software side, algorithms that exploit periodicity, such as those based on Floquet engineering or topological quantum error correction, may benefit from the inherent stability of time-crystalline phases [22] (Figure 12).

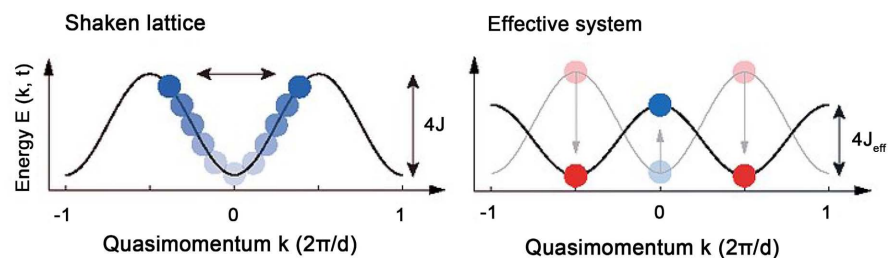


Figure 12. Example of Effectiveness using the Floquet System [22].

Ultimately, time crystals extend the geometric and topological themes of this report into the quantum domain, illustrating how symmetry, periodicity, and homological structure can stimulate educated intuitions that shape the frontier of quantum technologies.

14. Conclusions and Further Developments

The foundational work of Sunada brings forward modern mathematical tools for modeling crystals and similar structures, using Algebraic Topology (Homology and Cohomology) and Algebraic Geometry.

On the “intrinsic” side, the theory of periodic nets and building blocks formu-

²Specifically, the recent error correcting technique referred to as *Quantum Echo* [23] supports considerations of “retro-causality” which are indicative of the universal cover role of Lab time-parameter in otherwise periodic phenomena, modulo I/O-interactions.

lated by Sunada can be developed using *Graph Cohomology*, a general framework developed in [5]. This is a project that will be considered elsewhere. Further applications may benefit from *Topological Crystallography*, including perhaps Biology, with structures which are not necessarily periodic, but rather correspond to the building blocks themselves: RNA, DNA, proteins, etc. In other words, the general framework of Topological Crystallography viewed in the larger context of Graph Cohomology and Algebraic Geometry, may be the right “tool” for Mathematical Modeling in Chemistry, Biology, as well as in Elementary Particle Physics [10].

Author Contributions

The present article resulted from an independent study of the first author, MAT 400 Applied Group Theory at Illinois State University, equivalent to a full course on the presented topics, conducted by the second author (Lecture and Recitation 1:1). It is based on the notes and additional research done by the first author, with essential contributions of the second author to Sections 11 and 12.

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Conflicts of Interest

The authors declare no conflicts of interest regarding the publication of this paper.

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