Viruses and Geometry: Group, Graph and Tiling Theory Open Up Novel Avenues for Anti-Viral Therapy

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Viruses are responsible for a wide range of devastating illnesses in humans, animals and plants, yet options for treatment or prevention are limited. This is in part due to the occurrence of escape mutants that exhibit changes in the structures of the drug targets. New insights into evolutionary conserved features and constraints on virus structure are therefore key for the development of novel, more stable forms of anti-viral therapy. The area of Mathematical Virology, that focuses on the development and applications of mathematical tools to tackle open questions in virology, provides new opportunities to address this. Based on our unique interdisciplinary approach in which mathematical techniques from group, graph, tiling and lattice theory play key roles, we demonstrated that virus structure is much more constrained than previously appreciated. In particular, we identified structural constraints acting simultaneously at the level of the viral protein containers that encapsulate the viral genomes, and at the level of the packaged genomes. These new insights into virus structure have consequences for how viruses form, evolve and infect their hosts, and we developed mathematical techniques and models to quantify this. We derived new ways of characterizing genome organization within the viral capsids via graph theory, and used this to elucidate the mechanisms underpinning virus assembly. This resulted in the identification of previously unappreciated cooperative roles of single-stranded RNA genomes in virus assembly, and led to the discovery of a new anti-viral strategy against single-stranded RNA viruses, a major group of viral pathogens including HIV and Hepatitis C, in collaboration with experimental collaborators at the Universities of Leeds and Helsinki.

The following provides a summary of the key mathematical results and a discussion of their implications for our understanding of viruses and anti-viral therapy.
1. **Affine extensions of noncrystallographic Coxeter groups and virus geometry**

Protein containers encapsulating viral genomes are salient features of virus architecture. In most viruses, these containers are organized with icosahedral symmetry (cf. Fig. 1a) for reasons of genetic economy, and group theory can therefore be used to better understand virus geometry. We developed affine extensions of icosahedral symmetry to derive predictive information on the organization of viruses at different radial levels [28,32,35,45], revealing a previously unappreciated scaling principle in the overall organization of viral particles [14] (cf. Fig. 1c). Since icosahedral symmetry is non-crystallographic in three dimensions, i.e. is not compatible with periodic lattices, standard techniques for affine extensions do not apply in this case. We therefore developed a new framework for the construction of such affine extensions in the context of non-crystallographic Coxeter groups [3,8,16]. We also demonstrated that these mathematical structures, originally developed for applications in virology, can account for the atomic positions in nested fullerenes, carbon cage structures called carbon onions [7], demonstrating that they are more widely applicable in science.

2. **Viral Tiling Theory in virology and bio-nanotechnology**

The affine extended groups are by construction related to aperiodic tilings such as the Penrose tiling. We developed Viral Tiling theory ([27,36-44], see also Fighting viruses with mathematics, a case study by the Institute of Mathematics and its Applications) to model virus architecture via spherical tilings (cf. Fig. 1b). These tilings generalize the triangulations used in Caspar-Klug theory and indicate the relative positions of the proteins in viral capsids. Viral Tiling theory solved a long-standing open problem concerning the structures of the cancer-causing papillomaviridae [44], and also delivered models for tubular malformations that can arise during self-assembly of the major capsid protein of these viruses [37]. We recently further developed our tiling approach to make it applicable to broader classes of protein containers, including protein nanoparticles used in vaccine design (the SAPN system) [4]. Our approach resulted in a classification of all possible structural blueprints consistent with the self-assembly properties of the SAPN building blocks. In combination with information from mass spectrometry on the approximate numbers of constituent building blocks in each nanoparticle, this allowed us to...
identify the surface structures of the nanoparticles conclusively, which would not have been possible via experiment alone.

3. **Lattice Transitions provide insights into structural transitions important for infection**

A significant number of viruses must undergo structural rearrangements of their protein lattices in order to become infectious. Such maturation events are transient, and therefore difficult to monitor experimentally. We have developed a new mathematical framework to model such transitions based on our description of virus structure in terms of affine extended symmetry groups and surface lattices. Our approach uses the constraints encoded by the affine extensions of the icosahedral group as descriptors of capsid geometry, and exploits their relation with bases of six-dimensional crystallographic lattices to model the transition paths via projection of lattice transitions in six dimensions [5,17]. Since the descriptors derived from affine extended symmetry are by construction related to the vertex sets of tilings, this approach can also be used to model quasi-lattice transitions [18]. In order to better understand the biophysical aspects of capsid transitions, we also used an energy function to capture the interplay of different energetic contributions and describe capsid transitions in the context of a dynamical systems approach [15]. This work provides insights into the roles of asymmetric viral components in the initiation of the capsid transition, and characterizes the consequences of the resulting symmetry breaking for the expansive motion of the capsid.

4. **Novel applications of graph theory result in a paradigm shift in our understanding of virus assembly**

The assembly of viral protein containers from their component parts is a vital step in a viral life cycle. For decades, this process had been thought of as predominantly driven by protein-protein interactions, with the viral genomes at best playing minor roles via nonspecific, electrostatic interactions. Through the use of graph theory to describe the organization of the viral genomes within the capsids (see Fig. 1d), we were able to demonstrate that, by contrast, the viral genomes play important cooperative roles in virus assembly [2,12,13,20,23-26,31]. In particular, we used the concept of Hamiltonian path to formulate constraints on the organization of the packaged genomes in single-stranded RNA viruses. In combination with a novel bioinformatics approach developed by us, this revealed the existence of multiple dispersed sequence patterns/secondary structure elements in the genomes of these viruses that specifically interact with viral capsid proteins to promote capsid formation [10,12,13,22]. We subsequently extended this work to include a number of important Human and plant pathogens, including Hepatitis B, C and HIV [46]. We moreover developed models for virus assembly to elucidate the mechanism by which these multiple dispersed contacts confer efficiency and fidelity to capsid assembly [11]. For this we used the concept of Hamiltonian paths to quantify how viruses efficiently navigate the complex network of assembly intermediates, hence effectively solving a virus assembly equivalent of the Levinthals Paradox in protein folding [6].

5. **New mathematical models for virus assembly underpin the development of a novel anti-viral therapy**

Our assembly models are the first that take the cooperative roles of multiple dispersed packaging signals into account [6,11], and allow us to better understand the mechanisms underpinning packaging signal mediated assembly [1]. The models show that the importance of packaging signals in virus assembly can only be fully appreciated, if specific features of an
in vivo infection (such as the gradual build up of capsid protein, called the protein ramp) are factored into the analysis, perhaps explaining why the existence and crucial roles of the packaging signals had so long been overlooked. They also provide an in silico testing ground to probe the effects of anti-viral strategies targeting specific groups of packaging signals. The results show that such novel forms of anti-viral intervention can reduce viral load by delaying assembly and triggering misencapsidation of cellular RNAs. Moreover, an analysis of packaging signal motifs and positions across different viral strains, enabled by our novel graph theory based analysis techniques, revealed conserved features that lend themselves as drug targets for more stable anti-viral therapy [46].

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