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## Packing and trimer-to-dimer protein reconstruction in icosahedral viral shells with a single type of symmetrical structural unit

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Understanding the principles of protein packing and the mechanisms driving morphological transformations in virus shells (capsids) during their maturation can be pivotal for the development of new antiviral strategies. Here, we study how these principles and mechanisms manifest themselves in icosahedral viral capsids assembled from identical symmetric structural units (capsomeres). To rationalize such shells, we model capsomeres as symmetrical groups of identical particles interacting with a short-range potential typical of the classic Tammes problem. The capsomere particles are assumed to retain their relative positions on the vertices of planar polygons placed on the spherical shell and to interact only with the particles from other capsomeres. Minimization of the interaction energy enforces equal distances between the nearest particles belonging to neighboring capsomeres and minimizes the number of different local environments. Thus, our model implements the Caspar and Klug quasi-equivalence principle and leads to packings strikingly similar to real capsids. We then study a reconstruction of protein trimers into dimers in a *Flavivirus* shell during its maturation, connecting the relevant structural changes with the modifications of the electrostatic charges of proteins, wrought by the oxidative switch in the bathing solution that is essential for the process. We highlight the key role of pr peptides in the shell reconstruction and show that the highly ordered arrangement of these subunits in the dimeric state is energetically favored at a low pH level. We also discuss the electrostatic mechanisms controlling the release of pr peptides in the last irreversible step of the maturation process.

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## 1. Introduction

Viruses span an intermediate region between living and non-living matter, requiring at certain stages of their life cycle a host cell to replicate their genome and produce the genome-enclosing protein shell (capsid).<sup>1</sup> Self-assembly of new viruses is a multi-stage process requiring many steps controlled exclusively by physical forces and chemical equilibria.<sup>2</sup> For many viruses, the final stage of the assembly process, referred to as maturation, involves structural transformations of the

proteinaceous capsid that convert a noninfectious precursor particle into an infectious virion, able to proliferate in host cells.<sup>3</sup> Studying the principles of self-assembly and the structural transformations involved in the capsid maturation process is therefore crucial for the understanding of virus infectivity as well as developing appropriate antiviral strategies.

Being encoded by a relatively simple and by necessity, relatively short viral genome,<sup>4</sup> the capsid proteins cannot be too complex and can be characterized by a small number of conformational states and types of bond with their nearest neighbors. While an unlimited number of crystallographically equivalent positions would exist in an ideal periodic protein lattice, any closed discrete shell such as the proteinaceous capsid, by necessity, supports only a limited number of crystallographically equivalent positions. For identical asymmetric proteins in the case of capsids with icosahedral symmetry, this number equals 60, which is also the order of the capsid symmetry group I, implying furthermore that when the number of proteins exceeds 60, their environments cannot be strictly identical. Therefore, most proteinaceous capsids exhibit a hidden, *i.e.*, approximate symmetry<sup>5–8</sup> characterized by a local periodic order, allowing the proteins to occupy quasi-equivalent positions, as the next best thing to the geometrically forbidden

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strictly equivalent crystallographic positions. This principle of quasi-equivalence, introduced by Caspar and Klug (CK), permits the description of the structure of icosahedral capsids as consisting of pentamers and hexamers within their well-known viral shell model.<sup>9</sup>

While the CK model seems to be applicable to a plethora of viral capsids, it is not universal, and anomalous viral shells that cannot be rationalized within it, consisting of identical symmetric capsomeres, namely, dimers,<sup>10–14</sup> trimers,<sup>13,15,16</sup> or more rarely pentamers<sup>17,18</sup> and decamers,<sup>10,11,14</sup> abound in nature. These symmetrical structural units pre-assemble in a solution from identical proteins or symmetrically arranged protein domains before the capsid assembly, with intra-capsomere protein bonds clearly differing from the bonds between proteins of neighboring capsomeres. This situation begs the question of how do the viral proteins minimize the number of their local environments in this type of capsid. Some large viral shells assembled from trimers exhibit a locally periodic order of SU environments similar to the CK model, while some small anomalous viral capsids exhibit a hidden locally periodic order of SU environments, quite unlike the CK model.<sup>5</sup> In order to get a clearer grasp of the nature of local environments and packings of symmetrical structural units in these anomalous proteinaceous shells, we will show in what follows how their tendency to form dense packings on spherical shells also minimizes the number of different local environments.

The simplest dense particle packing on a spherical surface can be formed by disks, and the well-known Tammes problem is devoted to finding the densest packings for  $N$  identical disks embedded on a spherical surface.<sup>19</sup> The equilibrium dense packing of disks can be obtained by minimizing the sum of the pair interaction energies of equivalent point particles located at the disk centers. If all disks are identical and touch each other, then the short-range potential of such pair interaction decreases with increasing distance  $l$  between their centers as  $l^{-n}$ , where  $n \rightarrow \infty$ .<sup>20,21</sup> Going beyond the disk-like structural units, we now generalize the Tammes problem and consider equilibrium configurations of identical symmetrical groups of particles distributed on the spherical surface. In the proposed model, within the SU that corresponds to a capsomere, the distances between particles are kept fixed while particles from different SUs interact with each other in the same way as in the Tammes problem. As we show, the resulting equilibrium packings on a spherical surface turn out to be similar to the structures of certain actually existing viral shells. An analysis of the structural data<sup>22</sup> for such shells reveals that the separations between nearest proteins belonging to neighboring symmetrical structural units are similar. This allows us to use the short-range potential to reproduce the general features of the considered dense protein packings and clarify how the CK quasi-equivalence principle works in the considered case.

As part of the generalization of the Tammes problem and as an illuminating illustration of our theory, along with other examples of capsid self-assembly from a single type of symmetrical SU, we consider specifically the packaging of proteins in the immature and mature outer shells of the Dengue virus and similar viruses from the *Flavivirus* genus in the family

*Flaviviridae*. In viruses of this genus, the immature viral shell first self-assembles from trimers and then reorganizes itself into a shell composed of dimers during the maturation process. This structural change occurs with an increase in the acidity of the bathing environment,<sup>23</sup> and we show how the change in the electrostatic energy of the system induced by a change in the pH level controls this intriguing dimer-to-trimer transformation. Moreover, we also show that the pH change finally assists in the subsequent irreversible stage of the capsid maturation yielding the final form of the virion.

## 2. Results and discussion

### 2.1. Modelling dense packing of identical structural units

To construct the model energy function that would describe the capsomere packing, we consider regular identical rigid polygons placed on the spherical shell, with interactions operating only between their vertices. We assume that these polygons can change the position and orientation, while their centers remain confined to the spherical surface, and their plane is constrained to coincide with the tangential plane to the surface. Because the polygons are rigid, only the interactions between different polygons need to be taken into account, and we therefore postulate that the interaction energy is of the form:

$$E = \sum_{i>j,k,l} \frac{N^2}{|\mathbf{R}_i - \mathbf{R}_j + \mathbf{r}_{i,k} - \mathbf{r}_{j,l}|^n}, \quad (1)$$

where  $\mathbf{R}_i$  and  $\mathbf{R}_j$  are the centers of  $i$ -th and  $j$ -th polygons, while  $\mathbf{r}_{i,k}$  and  $\mathbf{r}_{j,l}$  are the vectors connecting these centers and the  $i,k$ -th and  $j,l$ -th interacting vertices.  $N$  is the number of vertices or equivalently the number of interacting particles in each polygon,  $k = 1, 2, \dots, N$ ,  $l = 1, 2, \dots, N$ . The interaction magnitude of all interacting vertices is identical and taken symbolically as 1. In the limit of long-range electrostatic interactions,  $n = 1$ , while for short range contact interactions  $n \rightarrow \infty$ , as in the Tammes problem.<sup>20,21</sup> In addition to the value of  $n$ , the second main parameter in the model is the ratio  $r/R$  between the radius of the polygons and the radius of the embedding sphere.

Note that the interaction energy (1) contains only terms corresponding to repulsive interactions. While at first glance it may seem that a model disregarding the attractive interaction between distant particles is an oversimplification, in actuality when self-assembly is modeled on the surface of a finite manifold, the confinement on the surface is equivalent to an effective attraction of SUs, and energy minimization leads to a relatively uniform distribution of SUs over the surface.

Eqn (1) can be expanded into a series in terms of the ratios  $r/R_{ij}$  (where  $R_{ij}$  is the distance between the centers of  $i$ -th and  $j$ -th polygons) by using the standard theory of multipole decomposition, but the ensuing representation of the pair interaction energy involving convolutions of various multipole tensors is quite cumbersome and will not be pursued further. It is easier to use the symmetry of the problem from the very beginning, in particular the fact that the pair interaction energy in (1) is invariant under the  $2\pi/N$ -rotation of the regular polygon with  $N$  sides. Consequently, the interaction energy must be expanded



in a series of  $\cos(2\pi m\phi/N)$  and  $\sin(2\pi m\phi/N)$  terms, where  $m$  is an integer, and  $\phi$  is a rotation of the polygon around its  $N$ -fold axis. It is then possible to expand the interaction energy of two polygons to the lowest order in terms of the dependence on  $\phi$ . In particular, for two dimers ( $N = 2$ ) of radius  $r$ , whose centers are separated by a distance  $R$ , the expansion up to order  $O((r/R)^4)$  yields:

$$E^{(4)} = \frac{4}{R^n} + \frac{n(n+2)r^2}{R^{n+2}} + \frac{n^2r^2[\cos(2\phi_1) + \cos(2\phi_2)]}{R^{n+2}}, \quad (2)$$

where the angles  $\phi_1$  and  $\phi_2$  are the angles between the vector  $\mathbf{R}$  and the direction from the corresponding dimer center to its vertex. We emphasize that expansion (2) depends neither on the rotation of the dimers around the direction of the vector  $\mathbf{R}$  nor on their relative orientation. As is also clear from expansion (2), the minimum interaction energy configuration for dimers that are far from each other should correspond to orientation perpendicular to the vector  $\mathbf{R}$ . As in the case of regular dimers, the expansions for the pair interaction energy of regular trimers, tetramers and pentamers, in terms of the separation between a pair of them, are also independent of the rotation around the direction of the vector  $\mathbf{R}$ , as well as of the relative orientation of the corresponding polygons relative to each other. However, in the case of  $N > 2$ , there emerges an additional dependence on the angle between the normal to the polygon and the direction of the vector  $\mathbf{R}$ . In this case, the dependence of the pair interaction energy on  $r$  remains in the term scaling as  $R^{-n-2}$ , but the dependence on  $\phi$  appears only with terms decreasing as  $R^{-n-N}$ .

The features of the multipole expansions can be very fruitful for constructing various model interaction potentials for different nano-objects,<sup>23</sup> including interactions between capsomeres in viruses, that are sufficiently far apart. However, when considering specifically the problem of dense (contact) packing of capsomeres, one should stick with the complete expression of the interaction energy (1). Comparison of the spherical packings of viral shell capsomeres based on the interaction energy (1) is carried out in the next section.

## 2.2. Model structures of viral capsids

**2.2.1. Capsids self-assembled from dimers.** As is known, icosahedral viral capsids assembled from a single type of capsomere can consist of dimers,<sup>10–14</sup> trimers,<sup>13,15,16</sup> pentamers<sup>17,18</sup> and decamers.<sup>10,11,14</sup> We selected such shells from the Protein Data Bank (PDB)<sup>22</sup> to analyze their protein packings in the framework of our model. For numerical minimization of energy (1), we used a value  $n = 200$ . In the limiting case  $n \rightarrow \infty$ , the distances between all pairs of nearest interacting points should tend to be the same; for the value of  $n$  used, the scattering of such distances for all model structures considered in our work does not exceed  $\pm 0.25\%$ . The transition to the limit  $n \rightarrow \infty$  will be considered in more detail below.

Equilibrium packings arising from energy minimization by the rotation method (see the Methods), as in other similar methods such as the gradient descent method, generally depend on initial coordinates and orientations of the SUs. We

used PDB data to define the initial state of the system before minimization. First, we defined centers of mass (CoMs) of individual proteins. Then, these coordinates were used to calculate coordinates of the centers and orientations of symmetrical SUs. Since CoMs only approximately characterize positions of the proteins in the structure, we performed multiple minimizations of the energy (1) by slightly varying the initial conditions to make sure that a model structure similar to a real capsid would not be accidentally missed.

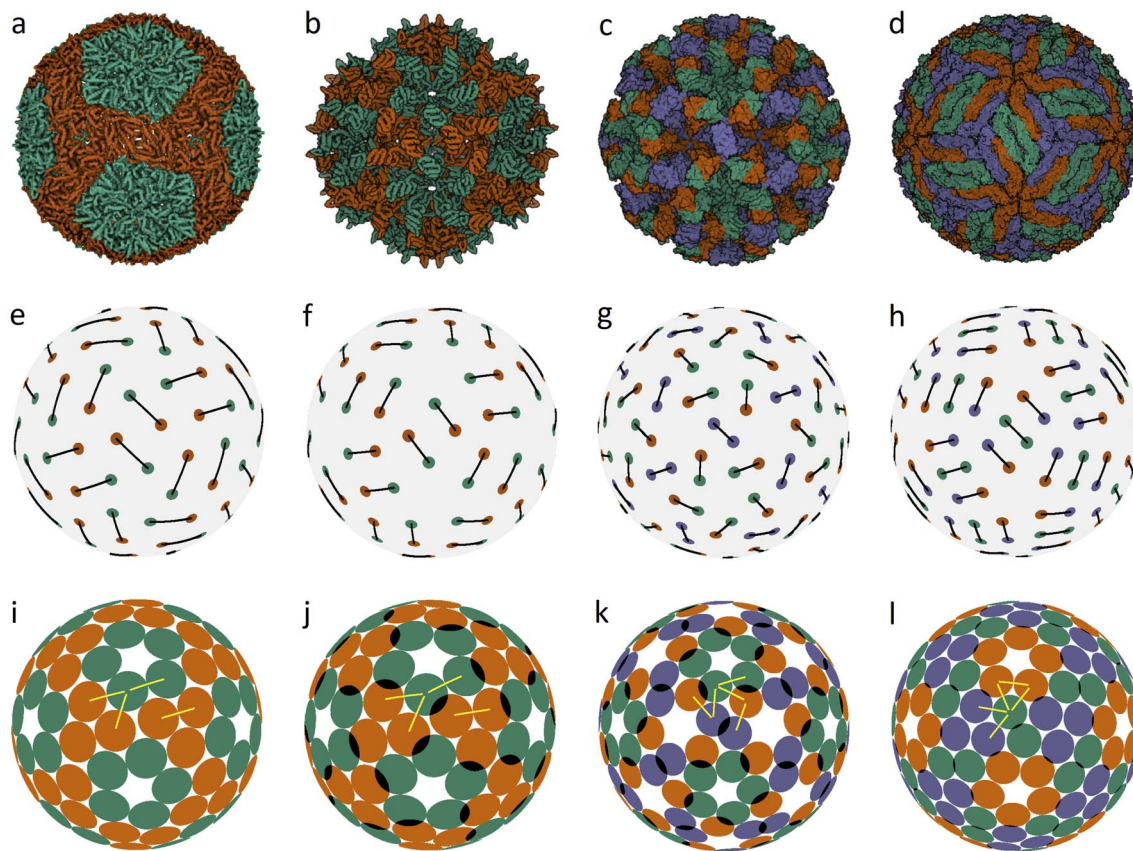
In an icosahedral capsid, the dimer centers can lie on the twofold axis and/or occupy general positions, so that such a capsid can be assembled from  $30T$  dimers, where  $T$  is an integer equal to the number of crystallographic orbits of proteins in the capsid. Fig. 1 shows examples of capsids self-assembling from dimers with  $T = 2, 3$  and models of these shells obtained by minimization of energy (1). Since the capsids can significantly change their shape and rearrange themselves during viral maturation, we explicitly indicate the PDB identifiers of the structures under consideration. In this work, all images of viral capsids were taken directly from the PDB website or obtained with UCSF Chimera.<sup>25</sup>

Fig. 1a shows the mature capsid of bacteriophage  $\phi 6$ , which acquires an approximately spherical shape after genome packaging. In the mature state, the mass centers of proteins, despite their grouping into dimers, correspond to a conventional packing of disks on a sphere: the distance between the mass centers of proteins in a dimer and that between the proteins from neighboring dimers practically coincide.

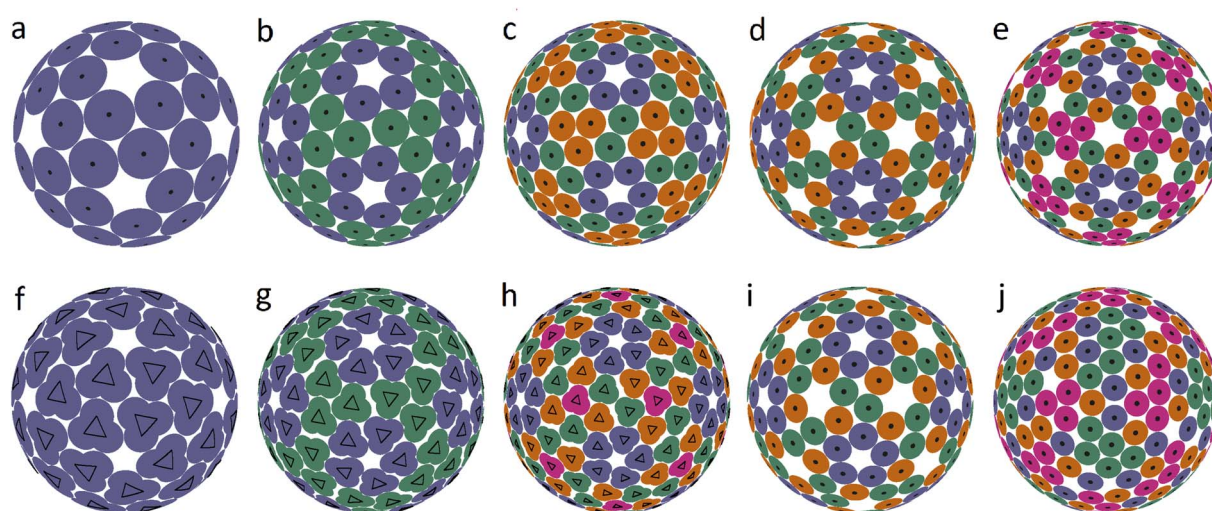
To assess the stability of the model structures, we randomly shifted the centers of the dimers as well as randomly rotated them. Under this perturbation, the displacement of the dimer center and the shift of the dimer vertices (mass centers of individual proteins), due to dimer rotation in the local coordinate system, were both limited by a certain value  $V$ . After this perturbation, we minimized the energy (1) again and observed that the model structure shown in Fig. 1i is stable within  $V/R \approx 0.05$ . Importantly, we could not find any structures with lower energies and thus assumed that probably this packing of dimers corresponds to the global minimum of energy (1). The structure shown in Fig. 1j is stable at slightly larger random distortions  $V/R \lesssim 0.06$ ; however, we found an asymmetric packing with a slightly lower energy. The model structure from the 3rd column (unlike the other structures presented in this figure) is stable only if the icosahedral symmetry is imposed during the energy minimization. Otherwise, the packing loses its symmetry after small perturbations, turning into a more energetically favorable structure with slightly shifted dimers, as in the case of the icosahedral packing of non-overlapping disks (Fig. 2d) that undergoes a similar transformation into a more energetically favorable structure (Fig. 2i).

The mature shell of Dengue virus from the *Flaviridae* family consists of dimers (see the last column of Fig. 1). The immature shell initially self-assembles from trimers and then radically rearranges its structure, the situation which is considered in more detail in the next sections. The structural model of the virus (Fig. 1l) is stable for  $V/R \approx 0.04$  and probably corresponds to the global minimum of energy (1).





**Fig. 1** Capsid structures from PDB: (a) bacteriophage  $\phi$ 6 (4BTQ) (Cystoviridae) ( $T = 2$ ); (b) penicillium stoloniferum virus S (3IYM) (Partitiviridae) ( $T = 2$ ); (c) Norwalk virus (1IHM) (Caliciviridae) ( $T = 3$ ); (d) dengue virus (3J05) (Flaviridae) ( $T = 3$ ); (e–h): position of the CoMs according to PDB with the corresponding arrangement of dimers. The ratio of the dimer length to the sphere radius  $r/R$  for cases (a–d) is: 0.172, 0.1355, 0.104, and 0.13; (i–l) optimized icosahedral model structures with the same  $r/R$  ratio. In each structure, all disks are identical in size, and a pair of overlapping disks corresponds to a dimer. In (a), distances between the disks in a dimer and disks between neighboring dimers practically coincide. The yellow segments correspond to quasi-equivalent bonds that are not symmetry equivalent. The lengths of these segments in the limit  $n \rightarrow \infty$  become exactly equal. The corresponding distances between the protein CoMs in real structures (see (e–h)) are close, but not equal.



**Fig. 2** Dense packings of disks and model trimers. The equilibrium spherical structures obtained by minimizing the interaction energy of point particles (represented by small black circles), which interact with each other with the  $l^{-200}$  pair potential. (a–e) Icosahedral packings of disks. Different colors denote different 60-fold orbits of the symmetry group I. (f–h) Icosahedral packings of particles rigidly bound into symmetrical trimers. Ratios  $r/R$  for these cases are: 0.12, 0.08, and 0.004, while the packings consist of 60, 120 and 180 trimers, respectively. (i and j) Packings arising from (d and e) when the requirement of icosahedral symmetry is removed.



Concluding the discussion of dimer shells, we note that minimization of energy (1) of an arbitrary initial arrangement of dimers with imposed icosahedral symmetry always leads to the model structures shown in the lower line of Fig. 1. Naturally, the obtained packings depend on the ratio  $r/R$  and the number of dimers (60 or 90).

It is interesting that the model structures presented in the last line of Fig. 1 can be additionally symmetrized if all particles interact identically with a pair potential  $l^{-200}$ , *i.e.*, particles are no longer grouped into dimers. The resulting packings are shown in Fig. 2b–d. This figure shows the first 5 spherical structures (panels a–e) with the icosahedral symmetry and location of SUs in general crystallographic positions. Since proteins cannot occupy positions with nontrivial symmetry, such icosahedral dense packings are not exhibited in Fig. 2. The dense packing makes the local environment of symmetry-nonequivalent disks quasi-equivalent, and therefore in the structure (b), both types of disk (blue and green) have 5 nearest neighbors, while in the structures (c–e), the blue disks that form pentamers around the 5-fold axes have 5 nearest neighbors, and the others have 4 nearest neighbors.

Structures 2(a–c) are stable for  $V/R \lesssim 0.12$ , 0.1, 0.09 and, according to our calculations, (b and c) are potential solutions to the Tammes problem for 120 and 180 disks. Recall that so far, only solutions to the Tammes problem for 100 or fewer disks have been found.<sup>26,27</sup> Packings (d and e) are stable only for such distortions of the SU positions that preserve icosahedral symmetry, while small random distortions force these structures to lose their icosahedral symmetry. Packing (d) is transformed into a slightly distorted, more energetically favorable packing (k), and packing (e) is converted into various strongly distorted structures like (l). With stronger perturbations, the packing (k) subsequently rearranges itself into a more energetically favorable structure. A complete analysis of the energy landscape of the proposed model is computationally cumbersome, as is well known from the studies on other similar systems<sup>28</sup> and clearly goes beyond the goals of this work.

While packings 2a–e are a somewhat rougher approximation to the structures of the relevant capsids, they are nevertheless still useful, since they can be helpful in rationalizing other types of spherical viral shell. It is straightforward to see that the close-packed structures shown in Fig. 2a–e are based on so-called icosahedral spherical lattices (SLs), in which the nodes lying on the symmetry axes of the icosahedron are excluded. The relationship between these SLs and the arrangement of proteins in small spherical viruses was analyzed in ref. 5. Recall that regular SLs (without exclusion of positions) are the basis of the CK model of viral capsids,<sup>9</sup> in which twelve 5-valent nodes of SLs are occupied by pentamers, and the remaining nodes with 6 neighbors are occupied by hexamers. SLs can be constructed as the mapping of the nodes of a simple hexagonal lattice onto the surface of the icosahedron. Due to geometric constraints, the edge of the icosahedron must be a translation of the hexagonal lattice, and translation indices are used to characterize the resulting SL.

**2.2.2. Capsids self-assembled from trimers, pentamers and decamers.** The organization of icosahedral viral capsids can be

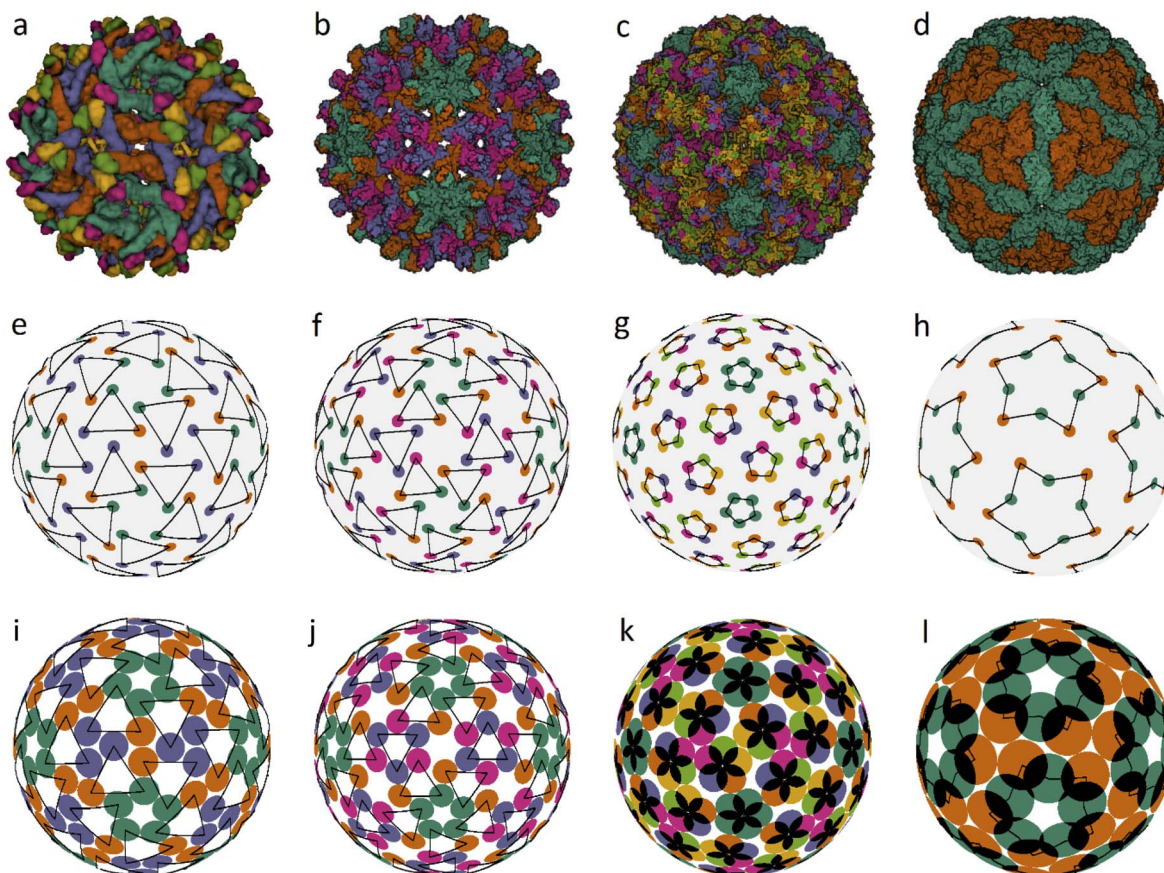
quite variable, but it is almost always based on icosahedral SLs. For example, in large viral capsids<sup>29–33</sup> self-assembled from trimers, their centers are located at all those nodes of the corresponding SLs that do not lie on the 2-fold and 5-fold icosahedral axes.<sup>6</sup> Similar model packings shown in Fig. 2f–h are obtained by minimizing the energy (1) of structures based on SLs (1, 2), (1, 3) and (1, 4), where the small trimers occupy the lattice nodes. The resulting structures are stable at  $V/R \lesssim 0.07$ , 0.05, 0.01. The structures shown in Fig. 2f and g probably correspond to the global energy minima. Note that for trimers that are sufficiently far apart, it is energetically favorable to line up parallel to each other. On the triangular faces of a spherical icosahedron, trimers form regions of local translational order, which are called symmetrons in the case of large viral capsids.<sup>33,34</sup> Most large viral shells consisting of trimers are actually arranged in this way.<sup>33,35,36</sup> Similar to the edges of large capsids that form linear structural defects, referred to as cleavage lines,<sup>33,34,37,38</sup> in our model, the translational order is likewise violated on the edges of spherical icosahedra.

We also found several examples of small icosahedral shells assembled from 60 and 80 trimers. Two of them are shown in Fig. 3. They are those of the immature *Zika* virus (6LNU) (*Flaviviridae*) and *Hepatitis B* virus (6W0K). The corresponding model structures are stable under random distortions within  $V/R \approx 0.03$  and 0.01, respectively. While the first structure probably corresponds to the global minimum of energy (1), it is strongly reconstructed during the maturation of *Zika* virus. The self-assembly of *Hepatitis B* virus is also very interesting. After the initial stage of self-assembly, where dimers are formed in solution, they are subsequently reassembled into trimers, and the resulting structure is often treated as consisting of pseudo-hexamers.

Assembly of some viruses also involves formation of intermediate complex SUs from dimers. For example, a trimer of dimers is the basic building block of HIV<sup>39</sup> and some other large viruses (from families *Phycodnaviridae* and *Iridoviridae*).<sup>33,35,36</sup> Self-assembly of  $T = 2$  capsids of dsRNA viruses from the *Reoviridae*<sup>44</sup> and *Totiviridae*<sup>40</sup> families starts with the formation of pentamers of dimers in the bathing solution. These transient SUs are also called pseudo-decamers,<sup>10</sup> and an example of such a capsid is shown in Fig. 3d. Our analysis of PDB structural data on *Reoviridae* and *Totiviridae* reveals that CoMs of individual proteins in viruses from these two families practically coincide. The CoM positions, shown in Fig. 3h, make individual decamers clearly distinguishable. For comparison, this is not the case in the  $T = 2$  structures shown in Fig. 1e and f. Indeed, during self-assembly of  $T = 2$  capsids from the families *Partitiviridae* and *Cystoviridae*, no decamers are formed.<sup>14</sup> The arrangement of CoMs of proteins in the decamer (Fig. 3h) has  $C_5$  symmetry, and the CoMs form two regular pentagons rotated relative to each other. Within the framework of energy (1), the packing of decamers into an icosahedral capsid corresponds to the global energy minimum. Note that in the considered case, the system energy has only two minima.

Fig. 3c shows another very interesting example of *Human Papillomavirus* capsid (panel c) and similar capsids assembled from 72 pentamers and belonging to the same family





**Fig. 3** Viral capsids self-assembled from trimers and pentamers, as well as their corresponding model structures. Capsid structures according to PDB: (a) immature Zika virus (6LNU), (b) Hepatitis B virus (6W0K), (c) Human Papillomavirus (5KEP), (d) Leishmania RNA virus (6H83), (e–h) positions of the CoMs of individual proteins. The location of trimers, pentamers and decamers is shown with triangles, pentagons and five-pointed stars, respectively, drawn with black lines. The ratios of the capsomere radius to the sphere radius  $r/R$  for cases (e–g) are: 0.254, 0.187, and 0.12; the decamer in (h) is given by two pentagons rotated relative to each other at  $\sim 30^\circ$  with  $r/R = 0.32$  and 0.48, respectively. (i–l) Optimized model structures with the same ratio  $r/R$ .

*Papillomaviridae*. In such capsids, the pentamer centers are localized at all SL (2, 1) nodes. Fig. 3g and k show positions of protein CoMs and the model structure corresponding to the global energy minimum at the considered ratio  $r/R = 0.12$ .

An analysis of the real structures shown in Fig. 1 and 3, as well as other similar structures self-assembling from a single type of capsomere, reveals that the number of different distances between CoMs of the neighboring proteins and, accordingly, the number of corresponding bond types, are minimal. Bonds of the first type connect CoMs of nearest proteins within symmetrical SUs, whereas bonds of the second type connect CoMs of nearest proteins belonging to neighboring SUs. Bonds of the second type are similar in length, which is, however, not determined by the symmetry of the system.

In the proposed model, minimization of the total energy for short-range interaction potential with  $n = 200$  furthermore equalizes the second-type bond lengths up to a spread of  $\pm 0.25\%$ . In the case  $n \rightarrow \infty$ , all these symmetry-nonequivalent lengths become exactly the same. It is easy to determine the total number of such lengths for an arbitrary packing of the

considered type. In the general case, in order to uniquely specify the position of the capsomere on the sphere, all three coordinates must be determined. Two of them determine the position of its center, and the third one describes the capsomere orientation. If the capsomere position coincides with a symmetry axis of the icosahedron, then only one coordinate remains, specifying the rotation of the capsomere. To determine  $M$  coordinates, it is necessary to solve the system of equations stemming from the condition that  $M + 1$  symmetry-nonequivalent distances are identical. Thus, there should be 4 such distances (see the corresponding yellow lines in Fig. 1) for structures (k and l) and 5 for structures (m and n). However, finding actual solutions of the corresponding system of equations leads to coordinate corrections, which are so small that they are indistinguishable on the scale of Fig. 1–3. It is also interesting to note that our model compares favorably with real structures not only in the case of distances just discussed (that become exactly equal in the  $n \rightarrow \infty$  limit), but also for other structural properties. For example, in the vicinity of the 3-fold axis (see Fig. 1l), the distance between the centers of SUs in the model structure turns out to be somewhat larger than the



minimal one, resulting in a small gap between the disks, which tallies with the situation in real capsids (see Fig. 1d), where the corresponding distances between protein CoMs (see Fig. 1h) are also similar in length. In all cases presented in Fig. 1 and 3, it is easy to notice other similar examples by paying attention to the corresponding model structure and finding disks separated by a small gap. Thus, both real structures and model packings are in good agreement with the CK principle of quasi-equivalence of symmetry-nonequivalent bonds between nearest proteins.

### 2.3. Morphological changes in *Flavivirus* viral shells during their maturation

In the previous sections, we proposed a simple packing model for immature and mature viral shells of *Flaviviruses*. Recall that the immature shell self-assembles from trimers, and then during the maturation process, the trimers reorganize into dimers. Let us discuss this intriguing transformation that has two intermediate stages<sup>23</sup> in more detail using the PDB structural data on the dengue virus shell. The radical reconstruction occurs at the first maturation stage when the trimeric structure 3C6D turns into the dimeric one 3C6R (see Fig. 4a and b). Both states consist of 180 complex proteins, referred to as the heterodimers.<sup>41</sup> Each heterodimer includes the main M–E part and the pr peptide. The main part is the largest and has the shape of a short, bent rod that is thicker at one end (head). The small pr peptide is connected to the thinner end (tail) of the main SU.

In the 3C6D state, the immature shell consists of 60 trimers, each of which is formed by three heterodimers connected close to their heads. In Fig. 3e, such a structural organization is represented by a motif of triangles with vertices being CoMs of individual heterodimers belonging to the same trimer. In Fig. 4a a pair of such trimers, that is symmetric with respect to the 2-fold axis of the immature shell, is highlighted in color. Like their heads, the tails of heterodimers are also connected in triplets located above the centers of neighboring trimers. So, in Fig. 3e, one can see that the triangle vertices also form the vertices of a weakly deformed hexagon, which allows classifying the 3C6D structure as a pseudo-hexagonal one.

By establishing the correspondence between protein positions in trimeric state 3C6D and dimeric state 3C6R states, one can fully define the structural mechanism of the transition. For this aim, we superimpose the heterodimer CoMs and assume that the CoM displacements are small (see Fig. 4c). Then, considering these reasonable displacements of CoMs in the vicinity of 3-fold axes, one can see that there are two possible rearrangements of this heterodimer orbit as the corresponding CoM displacements are close in lengths. Fig. 4a and b show the rearrangement in which heterodimers rotate less. Let us note that details of the considered 3C6D → 3C6R transition are still unclear. For example, rearrangement that leads to a larger rotation of heterodimers has been recently considered in ref. 41. In contrast, earlier work<sup>42</sup> focused on a different type of rearrangement characterized by smaller rotations of heterodimers, but involving longer shifts of their CoM positions.

Independent of the rearrangement mechanism, formation of the first intermediate dimeric state 3C6R requires that

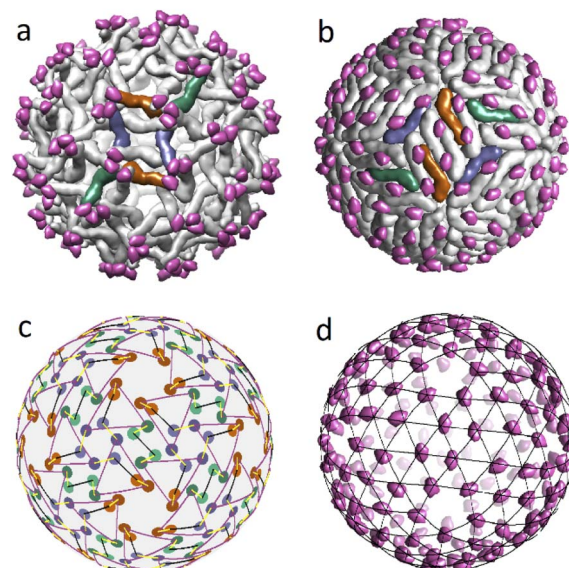


Fig. 4 Structural transformation occurring during the Dengue virus shell maturation. (a) Trimeric structure of the viral shell 3C6D. M–E subunits of six heterodimers forming two neighbouring trimers are colored in dark orange, blue and green. (b) Dimeric structure of the viral shell 3C6R. The same subunits M–E as in panel (a) are shown in color to demonstrate correspondence between two structures. (c) Superimposed mass centers of heterodimers in trimeric and dimeric structures. CoM positions in the dimeric structure are represented by lighter circles. The displacement field of the mass centers that defines 3C6D ↔ 3C6R correspondence is shown with yellow lines. (d) Positions of the pr peptides in 3C6R with superimposed icosahedral SL (3, 2). In (a, b, d), pr peptides are shown in magenta.

heterodimer tails cease to combine into triplets. In this context, it is interesting to note that in the dimeric state, the pr peptides (attached to heterodimer tails) occupy almost equidistant positions and form a relatively regular structure (see Fig. 4d). Note also that the second intermediate state 3YIA involves irreversible changes, as the connection between the pr peptide and the M part of the heterodimer is terminated. Nevertheless, in this state, cleaved pr peptides remain associated with the shell. Finally, at the end of the maturation process, when the shell is reintroduced into the neutral environment, pr peptides are released from the viral surface.

Unlike the 3C6R → 3YIA transition, the first intermediate stage of the maturation, transforming the trimeric structure 3C6D into the dimeric one 3C6R, is reversible and can be activated *in vitro* by decreasing the pH of the solution to 5.5.<sup>23</sup> This experimental finding, combined with the fact that generally the net charge of a protein is controlled by the pH of the buffer solution, suggests that the electrostatic interactions between shell proteins could play a significant role in the 3C6D → 3C6R transition. To test this hypothesis, we examined how electrostatic energies (see the Methods for details) of the two shell states depend on the acidity of the solution.

First, we calculated the degree of dissociation and partial charge of each amino acid in the heterodimer at different pH values within the framework of ref. 24 and 43 using the Henderson–Hasselbalch equation (see the Methods). Total charges



of the whole heterodimer and its pr and M-E subunits, as a function of the pH value, are shown in Fig. 5a. All three plots increase monotonously with the decrease of the bathing solution pH. Both subunits are negatively charged at neutral pH. With the decrease of pH, the pr subunit remains negatively charged, whereas the charge of the M-E subunit changes its sign. Accordingly, the heterodimer as a whole has an isoelectric point at around pH = 6.

Let us note that generally speaking, amino acids can be ionized only when they are in contact with the solvent. 3C6D  $\rightarrow$  3C6R transition should modify contacts between heterodimers changing the solvent accessibility as quantified by the solvent accessible surfaces (SASs).<sup>44,45</sup> Because of the low resolution of structural data provided in PDB (positions of amino acids are defined by a single carbon atom), we were unable to consider the effects associated with the variation of SASs, so that in our model, the charges of SUs have the same pH dependencies in both structures.

To our knowledge, after ref. 46, structures of the intermediate states in viral shells of the *Flaviviruses* have not been experimentally studied; however, as we show later, our simplified model is sufficient for obtaining results that are in good agreement with experimental data and provides valuable insight into the maturation process itself.

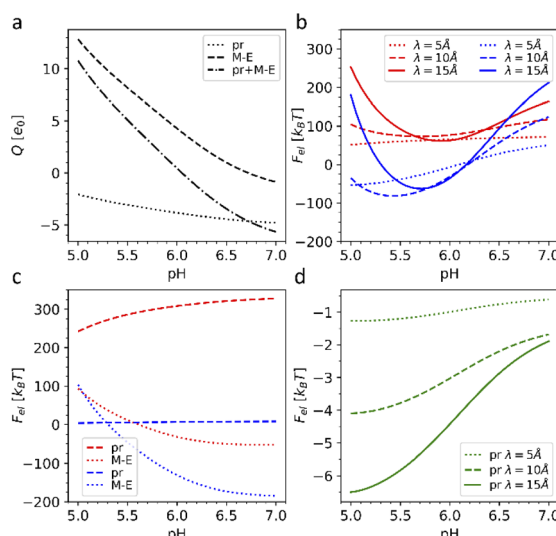


Fig. 5 Electrostatic effects associated with dengue shell maturation. (a) Charges of the single heterodimer and its M-E and pr parts separately, expressed in the elementary charge units  $e_0$  as a function of the pH level. (b) Effective electrostatic energies of the trimeric 3C6D (shown in red) and dimeric 3C6R (shown in blue) states expressed in  $k_B T$  units as a function of pH for three different screening constant values. (c) Energy of the effective electrostatic interaction between M-E (dashed line) and pr (dotted line) subunits separately (interactions between pr and M-E subunits are ignored) as a function of pH for three different electrostatic screening constant values  $\lambda$ . The plots corresponding to the trimeric and dimeric states are color-coded as in (b). (d) Effective energy of attractive electrostatic interaction between a single pr peptide and the rest of the 3IYA shell as a function of pH for three different electrostatic screening lengths  $\lambda$ . The value  $\lambda = 10 \text{ \AA}$  corresponds to the physiological salt concentration and temperature  $T = 300 \text{ K}$ .

A typical average value of the electrostatic screening length  $\lambda$  at the physiological salt concentration can be estimated to be  $\lambda = 9.74 \text{ \AA}$ ; however, since the local salt concentration can vary, we calculated energies (3) for the three typical screening lengths, namely: 5, 10 and 15  $\text{\AA}$  (see more details in the Methods). Fig. 5b shows that the electrostatic energy behaves qualitatively the same for all three  $\lambda$  values considered. For pH  $< 6.6$ , the electrostatic energy of the dimeric state becomes lower than that of the trimeric one, and the energy gap between the two states increases up to pH  $\sim 5.7\text{--}6.0$ , which are also the pH values reported for the maturation transformation.<sup>23,46</sup> Importantly, the 3C6R structure was studied experimentally at pH  $\sim 5.5$ ,<sup>46</sup> where, according to our calculations, the energy of the dimeric structure is substantially lower. The screening of electrostatic interactions implied by the energy (3) is an essential ingredient of the model, since, when  $\lambda$  exceeds the characteristic size of the capsid, the gain in the electrostatic energy of the dimeric state disappears.

To better understand why the transition from the trimeric to the dimeric structure lowers the electrostatic energy of the shell, we also considered the interaction energies of all M-E subunits and all pr subunits separately. In Fig. 5c, one can see that the energy associated with electrostatic interactions between pr peptides (dotted line) drops after the 3C6D  $\rightarrow$  3C6R maturation transition, strongly decreasing the total electrostatic energy of the whole shell. The pr subunits in the trimeric structure repel each other at all considered pH levels, substantially increasing electrostatic energy of the trimeric state. Even though this contribution slightly decreases with the decrease of pH, it is the structural transition that drastically reduces the electrostatic energy by forcing pr peptides into the positions corresponding to the icosahedral SL (3, 2) (Fig. 4d), and since the asymmetric pr peptides cannot occupy positions corresponding to the 5-fold axes, they are left vacant. Such a quasi-equidistant arrangement combined with the screening of the electrostatic interactions by the solution ions leads to the interaction energy between pr peptides that is close to zero. Thus, pr subunits play a key role in the considered reversible structural transition.

The details of the above scenario can be developed further by noting that the trimeric to dimeric structure transition could be engendered by another likely driving force. In fact, the proteinaceous outer shell self-assembles on a lipid membrane,<sup>47</sup> and it is well known that an important fraction of biological membrane lipids in general are anionic.<sup>48,49</sup> In our opinion, the membrane charge can play a role in virus surface protein rearrangement during Dengue maturation, where the membrane composition contains about 10% of anionic lipids.<sup>50</sup> Fig. 5a shows that at pH  $< 6$ , heterodimers as a whole are positively charged, and therefore, they are highly likely attracted to the negatively charged lipid membrane. Since the positive charge of heterodimers increases with the oxidation of the bathing solution, the attraction should also increase. Considering that in the dimeric state, heterodimers are located closer to the lipid membrane, we conclude that the membrane-to-shell electrostatic attraction can probably also contribute to the trimeric to dimeric structure transformation. We will analyze more details of this phenomenology in our future work.



Obviously, the proposed framework considers only electrostatic interactions between fixed protein units of outer shells and does not allow to take into account specific chemical changes during the irreversible transition to the second intermediate state 3C6R  $\rightarrow$  3IYA. Nevertheless, 3YIA structural data can be used to study the release of the pr peptides, which completes the maturation process and occurs when the viral shell returns to the neutral (pH = 7) environment. To explain this phenomenon, we analyzed how the energy of the electrostatic interaction between a single pr peptide and the rest of the 3YIA shell depends on the pH. Fig. 5d shows that with increasing pH, the attraction of the pr subunits to the viral surface significantly decreases, and an increase in the screening length leads to a steeper dependence. On a qualitative level, such behavior agrees with the way the subunit charges depend on pH (see Fig. 5a).

Our electrostatic model is based on several simplifications: we disregarded possible effects associated with the contribution of other long-range interactions,<sup>51</sup> such as van der Waals interactions, and we did not include the effects of various short-range solvent mediated and contact couplings, which certainly affect the packing of trimers and dimers and thus also the properties of the maturation transitions. Nonetheless, our analysis highlights the crucial role of the electrostatic interactions between the viral shell subunits and their connection with the variation in the pH bathing environment in both irreversible and reversible morphological transitions during the *Flavivirus* maturation process, rationalizing the existing experimental data.

### 3. Conclusions

Here, we studied the principles controlling the structural organization of capsids assembled from identical structural units (capsomers), which can contain two, three, five, or ten proteins. We demonstrated that the main morphological features of such viral shells are well reproduced by our new model, related to the Tammes problem,<sup>19</sup> that considers the densest spherical packings of identical discs. As was shown before,<sup>20,21</sup> these packings correspond to the global minimum of the short-range pair interaction energy scaling as  $l^{-n}$ , where  $l$  is the distance between disc centers, and  $n \rightarrow \infty$ . In our approach, interacting particles (located at the protein mass centers) are divided into identical SUs (which can be dimers, trimers, pentamers, or decamers), and the analogous short-range potential describes the interactions between the particles from different SUs. Within each SU, the relative coordinates of particles remain unchanged, forming a regular polygon with a fixed radius. Note also that an explicit attractive component of the interaction potential between SUs is not considered directly since an effective attraction is indirectly provided by the SU confinement on a spherical surface.

Thus, our approach takes explicitly into account only the effective size  $r/R$  and the  $n$ -fold symmetry of proteinaceous capsomers, which we believe are the two crucial properties characterizing the packing of identical capsomers on the spherical surface. The interaction model proposed in this work

appears to be the simplest meaningful implementation of the finite size and symmetry of interacting SUs confined to a spherical surface. While point-like models or models with point-like symmetry of interactions between capsomeres are certainly well known in the literature,<sup>52–57</sup> our model proposes a minimal generalization of these approaches that retains some features of the point-like interaction models but also takes into account the effective size  $r/R$  and the  $n$ -fold symmetry of the interacting SUs.

The short-range potential used in our model evens out the distances between nearest interacting particles and minimizes the number of local environments in the resulting packings, conforming fully with the Caspar and Klug quasi-equivalence principle. By comparing our results with PDB structural data, we have demonstrated that the proposed model is universal and suitably describes a large number of viral shells.

Understanding of principles controlling spherical dense packings of symmetrical capsomeres and corresponding contact interactions is essential for rationalizing structures and functions of virus shells. Concurrently, electrostatic interactions, being important in protein physics in general,<sup>58</sup> are also seen to play a key role in the virus life cycle. Proteins have electrical charge that can reach tens of elementary charges  $e_0$  per molecule depending on the pH level.<sup>59</sup> During their life cycle, viruses often travel through areas of the cell with varying pH and sometimes can even control it.<sup>60,61</sup> The electrostatic forces are therefore essential at practically all stages of virus development including capsid self-assembly,<sup>62–65</sup> genome packing,<sup>66,67</sup> and capsid modification during virus maturation.<sup>68</sup> The model describing contact interactions between identical symmetrical capsomeres, which has been developed in this paper, can be readily modified in the future to study electrostatic interactions between identical symmetrical groups of electrostatic charges corresponding to such capsomeres. To our knowledge, this problem has not been considered before, even though interactions of neutral multipoles in different systems including capsids are actively studied.<sup>2</sup>

In this work, we considered the role of electrostatic interactions in the pH controlled morphological changes of the proteinaceous outer shells of the *Flavivirus* genus. These changes are observed at the maturation and start from reversible transition from the trimeric to dimeric arrangement of proteins. As we have shown, the phenomenon can be explained by the changes in the charges of heterodimer subunits induced by the pH decrease and the tendency of the system to minimize its electrostatic energy *via* structural rearrangement. Pr peptides play a key role in this process, and the highly ordered arrangement of these subunits in the dimeric state makes this packing more energetically favorable than the initial trimeric one. We have also considered the irreversible transition of the outer virus shell that finalizes the maturation process and demonstrated that prior to the release of the pr peptides, the energy of their attractive electrostatic interaction with the viral shell significantly decreases due to the increase in the pH of the bathing solution. These results might be useful for the development of new immunogens, as some human antibodies (EDE2 A11, EDE2 B7, EDE1 C8 and EDE1 C10), neutralizing dengue



virus serotypes of *Flaviridae*, bind to E proteins specifically in the former positions of pr peptide attachment.<sup>69</sup> Thus, similar to pr peptides in the immature shell, arrangement of these antibodies on the surface of the mature capsid could be highly ordered, and their binding could be also controlled by the acidity of the surrounding medium, which we plan to examine in our future work.

## 4. Methods

### 4.1. Energy minimization

To minimize energy (1), we developed the method of finite rotations, that is a modification of the ordinary gradient descent. An arbitrary polygon motion (including a local rotation) over the sphere surface can be considered as a rotation of the polygon around a certain axis that passes through the sphere center. This approach to the considered constrain minimization problem ensures that the polygon remains embedded on the sphere surface. At each minimization step, first, we calculate moments of forces  $\mathbf{M}_i$  applied to the  $i$ -th polygon:

$$\mathbf{M}_i = \sum_{j=1}^N \left[ (\mathbf{R}_i + \mathbf{r}_{ij}) \times \mathbf{F}_{ij} \right],$$

where  $\mathbf{F}_{ij}$  is the force, applied to the  $j$ -th vertex of  $i$ -th polygon. The latter force is calculated directly by differentiating eqn (1). Then, we perform the corresponding small rotation of the  $i$ -th polygon through an angle  $\alpha_i$  that is proportional to the absolute value  $|\mathbf{M}_i|$  around the normal  $\mathbf{n}_i \parallel \mathbf{M}_i$ . The coefficient of proportionality between  $\alpha_i$  and  $|\mathbf{M}_i|$  is chosen so that the rotation decreases energy (4.1). The minimization of energy (1) is stopped when all moments  $\mathbf{M}_i$  vanish. In general, vectors  $\mathbf{M}_i$  and  $\mathbf{R}_i$  are non-collinear, which means that the rotation of the polygon around  $\mathbf{n}_i$  also involves its movement over the sphere surface. Let us recall that a rotation through angle  $\alpha$  around  $\mathbf{n}$  relates initial vector  $\mathbf{A}$  and rotated one  $\mathbf{A}'$  as

$$\mathbf{A}' = \mathbf{A} \cos \alpha + (1 - \cos \alpha)(\mathbf{n} \cdot \mathbf{A})\mathbf{n} + (\mathbf{n} \times \mathbf{A}) \sin \alpha.$$

Minimization of energy (1) with an accuracy of 10–15 digits may require about  $10^4$ – $10^5$  steps of the algorithm.

### 4.2. Calculating amino acid charges and debye screening length

To calculate partial charges of the amino acids, total charges of heterodimer subunits, and energies of their electrostatic interactions, we first extracted data on all ionizable amino acids from PDB structure files 3CHD, 3CHR, and 3IYA corresponding to the immature Zika virion, the first and the second intermediate states accordingly. In these files, each amino acid is represented by a single carbon atom. According to ref. 70, only six amino acids can carry electric charge. To calculate their charges at a given pH value, we used the Henderson–Hasselbach equation:

$$q_i = \frac{\pm 1}{1 + e^{\pm(pH - pK_a^i) \ln 10}}$$

where the plus sign is used when calculating charges of basic amino acids (arginine, histidine, and lysine), and the minus sign corresponds to acidic amino acids (aspartic acid, glutamic acid, and tyrosine). Values of the acid-based dissociation constant  $pK_a$  for these amino acids were taken from ref. 71. They are 3.71, 4.15, 10.10, 12.10, 6.04, and 10.67 for aspartic acid, glutamic acid, tyrosine, arginine, histidine, and lysine accordingly. When calculating electrostatic energy (3), we assume that the coordinates of amino acid charges are equal to those of the abovementioned carbon atoms and use the following expression for the screening length:

$$\lambda = \sqrt{\frac{\epsilon \epsilon_0 k_B T}{2 e_0^2 c_0}},$$

where  $k_B$  is the Boltzmann constant,  $\epsilon_0$  is the electric constant, and  $e_0$  is the elementary charge. The parameters we used to estimate  $\lambda$  are: temperature  $T = 300$  K, dielectric constant of water  $\epsilon = 80$  and monovalent salt concentration  $c_0 = 100$  mM, which are typical of experimental settings. These parameters correspond to the value of  $\lambda \approx 9.74$  Å.

### 4.3. Calculating the electrostatic energy

Knowing the charges  $q_i$  of the amino acids within each structural unit, we then calculated the normalized electrostatic energy of interactions between heterodimers forming the shell using the basic Debye–Hückel approximation:<sup>43,72</sup>

$$F_{el} = \frac{1}{4\pi\epsilon\epsilon_0 k_B T} \sum_{i>j,k,m} \frac{q_k^i q_m^j}{r_{km}^{ij}} \exp\left(-\frac{r_{km}^{ij}}{\lambda}\right) \quad (3)$$

where  $i$  and  $j$  enumerate heterodimers,  $k$  and  $m$  enumerate ionizable amino acids within a single heterodimer,  $r_{km}^{ij}$  are distances between the charges of amino acids,  $k_B$  is the Boltzmann constant,  $T$  is the temperature,  $\epsilon_0$  is the electric constant, and  $\epsilon$  is the dielectric constant of the solution. Thus, when calculating electrostatic energy of the shell, we considered interactions only between amino acids belonging to different heterodimers.

## Author contributions

S. R. and R. P. proposed main approaches. O. K. found examples of appropriate viral shells and drew the figures. I. G., O. K., and D. R., with the help of A. B. carried out numerical simulations. S. R. with the essential help of R. P. wrote the manuscript. All authors reviewed and edited the text. S. R. managed the project.

## Conflicts of interest

There are no conflicts to declare.

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