Conformational shift in the closed state of GroEL induced by ATP-binding triggers a transition to the open state

Yuka Suzuki$^{1,5}$ and Kei Yura$^{2,3,4}$

$^1$Department of Biology, Ochanomizu University, Bunkyo-ku, Tokyo 112-8610, Japan
$^2$Graduate School of Humanities and Sciences, Ochanomizu University, Bunkyo-ku, Tokyo 112-8610, Japan
$^3$Center for Informational Biology, Ochanomizu University, Bunkyo-ku, Tokyo 112-8610, Japan
$^4$National Institute of Genetics, Mishima, Shizuoka 411-8540, Japan
$^5$Present address: Okinawa Institute of Science and Technology, Onna-son, Okinawa 904-0495, Japan

Received June 11, 2015; accepted January 12, 2016

We investigated the effect of ATP binding to GroEL and elucidated a role of ATP in the conformational change of GroEL. GroEL is a tetracamer chaperonin that helps protein folding by undergoing a conformational change from a closed state to an open state. This conformational change requires ATP, but does not require the hydrolysis of the ATP. The following three types of conformations are crystalized and the atomic coordinates are available; closed state without ATP, closed state with ATP and open state with ADP. We conducted simulations of the conformational change using Elastic Network Model from the closed state without ATP targeting at the open state, and from the closed state with ATP targeting at the open state. The simulations emphasizing the lowest normal mode showed that the one started with the closed state with ATP, rather than the one without ATP, reached a conformation closer to the open state. This difference was mainly caused by the changes in the positions of residues in the initial structure rather than the changes in “connectivity” of residues within the subunit. Our results suggest that ATP should behave as an insulator to induce conformation population shift in the closed state to the conformation that has a pathway leading to the open state.

Key words: adenosine triphosphate, elastic network model, insulator, population shift, protein conformational change

Adenosine triphosphate (ATP) binds to a protein and provides energy for chemical reactions by the hydrolysis of the terminal phosphate [1]. The mechanisms of energy transfer from ATP to the chemical reactions have been studied extensively [1–3]. On the other hand, numerous cases have been found where the role of ATP molecule seems to be completed before the hydrolysis reaction. The role of ATP in these cases seems to reside in the binding itself, but the mechanistic detail of ATP binding to the conformation of protein remains to be elucidated [4–6].

GroEL is a tetracamer protein found in eubacteria and it mediates folding of unfolded proteins together with a heptamer protein GroES and seven ATP molecules. The single subunit of GroEL was initially identified as a two-domain protein [7] and later as three, i.e. equatorial, intermediate and apical domains [8]. We scrutinized the architecture of GroEL is a chaperonin that undergoes a conformational change from a closed state to an open state. This change requires ATP, but does not require ATP hydrolysis. To investigate the role of ATP in the conformational change, we conducted computer simulations using Elastic Network Model. We found that an ATP molecule behaves as a blockage that induces a population shift and increases the population of conformation that can lead to the open state conformation and decreases the population of conformation that cannot reach the open state conformation.
Biophysics and Physicobiology Vol. 13

molecule, and without hydrolysis of the ATP, GroEL with GroES transformed to the open state [12]. Hence, ATP hydrolysis is not required for the change from the closed to the open conformations in GroEL and the role of ATP molecules in conformational change is obscure.

Conformational change from the closed state to the open state of GroEL has been a research target for computational biology. Ma et al. performed a targeted molecular dynamic simulation from closed to open conformations of GroEL and found highly complicated displacement of the subunits in a tetradecamer [13]. Tehver et al. applied their original elastic network model method on GroEL complex and found the critical residues within and between the subunits for positive cooperativity that drove the transition from the closed to open conformations. They also found that the lowest normal mode dominated in the conformation change form the closed to the open states [14]. Yang et al. developed an adaptive anisotropic network model and described the collective dynamics of GroEL system. They found formations of critical interactions within the subunit for the transition [15]. All these computational studies focused on the pathway toward the open state via P state of GroEL, and none of these studies addressed the role of ATP molecule in the conformational change.

The previous experiments showed that the hydrolysis of ATP is not required for the transition of GroEL from the closed state to the open state [5], hence the energy barrier of this transition is likely to be low, but this low energy barrier should be high enough to be overcome only by ATP binding.

Figure 1  The structures of GroEL single subunit in P, M and open states. A. The distance map of GroEL single subunit (4HEL_A). The cutoff distance is 30 Å. B–D. The conformations of M (B), P (C) and open (D) states. These structures are retrieved from Protein Data Bank (PDB) [16]. PDB IDs of M, P and open states are 4HEL_A, 1KP8_A and 1AON_A, respectively. Equatorial domain is colored in red, intermediate domain in green, arm domain in magenta, and head domain in cyan. The grey molecule in P state is an ATP analogue. The grey molecule in D is ADP. In P state, three residues mutated for crystallization are shown in yellow space-filling model. Images of the structures were drawn with PyMOL [26].
To elucidate the role of ATP binding, we here conducted computer simulations of closed to open state transition, starting with P state and with M state, targeting at the open state. We found that M state could not reach the open state, whereas P state could reach the open state. Based on the simulation results, we suggest that ATP binding should cause population change of GroEL in the closed conformation from M state to P state by preventing P state from transferring to other presumably dead-end conformations, and that the role of ATP molecule should be an insulator that effectively shifts the conformation rather than be an energy supplier.

Materials and Methods

Protein coordinate data

Protein structure data were retrieved from PDB [16]. For structures of the α-th normal mode, and

\[ \Delta R_{i,j} = \sum_{\alpha=1}^{N} \Gamma_{i,j} \cos(\omega_{\alpha}t + \phi_{\alpha}) \]

where \( R_{i,j} \) is a distance vector between the \( i \)-th and the \( j \)-th Ca atoms. The fluctuation in the distance vector \( R_{ij} \) is \( \Delta R_{ij} = R_{ij} - R_{ij}^0 \), where \( R_{ij}^0 \) is the distance vector in the equilibrium. \( N \) is the total number of residues. \( \Gamma_{i,j} \) is the Kirchhoff or connectivity matrix that shows whether two Ca atoms are within a cutoff distance (\( r \)). Conformational changes were computed by repeating the following four operations (Fig. 2).

1. Normal modes were computed on a structure using the Tirion potential, \( \Gamma_{ij} \) is the Kirchhoff or connectivity matrix.
2. After obtaining 3\( N \)–6 modes, five modes with the five smallest non-zero eigenvalues were selected [14,21]. The displacements were calculated by

\[ \Delta q_i = \sum_{\alpha=1}^{N} C L_{(i)}^{(\alpha)} \cos(\omega_{\alpha}t + \phi_{\alpha}) \]

where \( \Delta q_i \) is the amount of displacement on the \( i \)-th generalized coordinate. \( L_{(i)}^{(\alpha)} \) is an eigenvector of the \( i \)-th component of the \( \alpha \)-th normal mode, and \( \omega_{\alpha} \) is an eigenvalue corre-

Simulations of conformational change

We conducted six types of simulations using two different models, namely Targeted Network Model (TNM) and Constant Connectivity Model (CCM). A brief summary of the six simulations is shown in Table 1. The six simulations are named TNM-P, TNM-M, CCM-PP, CCM-PM, CCM-MP, CCM-MM. P and M after the dash are short for P and M states, respectively.

Targeted network model (TNM)

For TNM, we conducted two types of simulations of conformational change from the closed states to the open state with P and M states as an initial structure (Table 1). All heteroatoms, including ATP molecules, and all atoms except Ca atoms were removed. The simulation was executed through computing normal modes of one subunit based on the Elastic Network Model (ENM) [19], which has been applied in the previous studies by others to simulate the conformational change of various proteins [13,15,20]. In ENM, potential energy was calculated by applying Tirion potential \( V \) [19,21],

\[ V = \frac{\gamma}{2} \left( \sum_{i=1}^{3N} (\Delta R_{ii})^2 \right) \]

\[ \Gamma_{ij} = \begin{cases} -1, & \text{if } i \neq j \text{ and } | R_{ij} | \leq r_c \\ 0, & \text{if } i \neq j \text{ and } | R_{ij} | > r_c \\ \sum_{j \neq i} \Gamma_{ij}, & \text{if } i = j \end{cases} \]

where \( r_c \) is a distance between the \( i \)-th and the \( j \)-th Ca atoms. The fluctuation in the distance vector \( R_{ij} \) is \( \Delta R_{ij} = R_{ij} - R_{ij}^0 \), where \( R_{ij}^0 \) is the distance vector in the equilibrium. \( N \) is the total number of residues. \( \Gamma_{ij} \) is the Kirchhoff or connectivity matrix that shows whether two Ca atoms are within a cutoff distance (\( r \)). Conformational changes were computed by repeating the following four operations (Fig. 2).

1. Normal modes were computed on a structure using the Tirion potential,
2. After obtaining 3\( N \)–6 modes, five modes with the five smallest non-zero eigenvalues were selected [14,21]. The displacements were calculated by

\[ \Delta q_i = \sum_{\alpha=1}^{N} C L_{(i)}^{(\alpha)} \cos(\omega_{\alpha}t + \phi_{\alpha}) \]

where \( \Delta q_i \) is the amount of displacement on the \( i \)-th generalized coordinate. \( L_{(i)}^{(\alpha)} \) is an eigenvector of the \( i \)-th component of the \( \alpha \)-th normal mode, and \( \omega_{\alpha} \) is an eigenvalue corre-
Corresponding to the eigenvector $L^{(i)}$, $C$, $t$ and $\delta$ are constant values. Two structures were obtained by changing the structure toward positive and negative directions of $\Delta q_i$. Incorporating the five lowest modes in the straightforward manner may seem insufficient for searching for the next conformation and all possible combinations of modes may be examined for the next conformation. According to the previous work on GroEL by Tehver et al., however, the lowest mode dominated in the transition from the open to the closed states and all other modes played insignificant roles in the conformational change [14]. In our calculation therefore, we took advantage of this result and considered only the lowest mode.

Remaining four low modes were included as perturbation terms about the lowest mode, (3) Full atom structures except hydrogen atoms of the two conformations were reconstructed on PULCHRA [22], and (4) Heavy-atom RMSDs between each of these reconstructed structures and the open state were calculated. Out of the reconstructed two structures, the one with the smaller RMSD was chosen as a new structure.

At each cycle of (1)–(4), we assigned the secondary structure with DSSP [23], and calculated the ratio of the number of Cα atoms that assumed helices in a computed structure to that in the initial structure. If the ratio was less than 0.65, the structure was regarded as broken. In addition, if the distances between the adjacent Cα atoms on the primary structure were out of the range between 3.6 and 3.9 Å, the structure was also regarded as broken. When a structure was broken, the other structure that had not been selected previously due to its larger heavy-atom RMSD would be selected. If there is a relatively high-energy barrier on the way, the structure should break, so that the simulation halts (dead-end) or starts to find another path. Each simulation was continued until the maximum of the gradient over the latest cycle or starts to find another path. This difference in the direction of the movement seems insufficient for searching for the next conformation and all possible combinations of modes may be examined for the next conformation.

**Constant connectivity model (CCM)**

We conducted four types of simulations with CCM (Table 1). The difference between TNM and CCM lies in the connectivities employed. In TNM, connectivity or Kirchhoff matrix was updated at each cycle, whereas in CCM the connectivity was constant throughout the simulation. We applied the connectivity of either P state or M state at every cycle. The other operations are the same as those of TNM (Fig. 2).

**Determination of the parameters $r_c$ and $\gamma$**

The parameter to determine the maximum length of connectivity $r_c$ was determined in such a way that more than two but no more than six eigenvalues in the first step of the simulation were effectively zero ($|\text{eigenvalue}| < 10^{-10}$). The parameter to determine the strength of connectivity (Hooke’s constant) in the Tirion potential, $\gamma$, was determined as a maximum value in the range where non-negative eigenvalues for all the calculations were found.

**Results and Discussion**

Of the six types of simulations, the simulations over the models of TNM-M, CCM-MM, CCM-MP, CCM-PM and CCM-PP converged by 5,000th cycle, and the simulation over the model of TNM-P converged at 7,317th cycle. In the following discussion, only the converged trajectories are used and dead-end trajectories (with broken covalent bonds or broken helices) were discarded.

**TNM-P reaches the open state whereas TNM-M does not**

The final structures obtained from the simulations of TNM-P (simulation from P state to the open state) and TNM-M (simulation from M state to the open state) are shown in Figure 3. The heavy-atom RMSD between the final structure in TNM-P and the open state was about 6.4 Å, and that in TNM-M was about 8.1 Å. TNM-P evidently approached the open state better than TNM-M. To identify the differences in the final structures of TNM-P and TNM-M, we compared the two structures using a difference map, a map that shows the subtraction of a distance map of TNM-M from that of TNM-P (Fig. 4). One of the most outstanding features in the difference map was found in the arm domain. The arm domain was the only domain that has negative (blue) distances with all other domains (Fig. 4). This result indicated that the direction of the movement of the arm domain in TNM-P and TNM-M was opposite from each other. This difference in the direction of the movement seems to depend on the difference in the initial structures of P and M states, as shown in the structural comparison on the initial steps of each simulation by DynDom [24] (Fig. 5).

The arm domain and the head domain in TNM-M rotated together against the upper part of the intermediate domain. In TNM-P, however, the arm domain, by making the region connecting the arm and head domains as a pivot, shifted toward the head domain. The difference in the rotation move-
The change in distance $R_1$ during the simulations is shown in Figure 7B. In the trajectories of TNM-P and TNM-M, the distance decreased, indicating that the intermediate and equatorial domains approached each other. This movement was also noted in the molecular dynamics simulation by Ma et al [13]. The trajectory of the distance between the geometrical centers of the domains shown in Figure 7C indicates that the changes in the distance of TNM-P and TNM-M were completely different throughout the simulations. The trajectory of TNM-P went down and reached the value close to the distance found in the open state (about 22 Å), whereas the trajectory of TNM-M steadily went up.

Qualitatively speaking, the current simulations showed that TNM-P was capable of reaching the open state, whereas TNM-M was incapable of it. Therefore, the function of the ATP molecule is either to induce the conformational change from M state to P state or to shift the conformational population from M state to P state. The small RMSD between the
two closed states suggests that the two states are reachable with each other without imposing high energy. A normal mode analysis between the two closed states showed that M state exhibited a strong tendency to approach P state [15]. These results are consistent with the fact that ATP plays a physical role that affects the deformation of GroEL by its existence, rather than a chemical role that provides high energy for conformational change. If this is the case, M and P states are reachable from each other and there could be a path between the two states. In reality, though, ATP binding is required for GroEL deformation and deformation from M state to the open state through P state without ATP has not been observed. The possible explanation of this behavior, based on the results of our calculations, is that M state has at least two major paths for the conformational change, and one is directed close to the ‘open state’ and the other to P state. Without ATP, the path directed close to the ‘open state’ should be wide-open and this is the path we found in the simulation as dead end. With ATP, the path to the dead-end ‘open state’ as well as other possible states are blocked and the path to the P state remains. The path is a detour to the open state, but reachable to the open state. M and P states are likely in equilibrium, where the population of P state is much smaller than that of M state before ATP binding but the population of P state increases after ATP binding. The role of bound ATP molecule is, hence, likely to be an insulator to shift the population from M state to P state.

Potential of forming a multimer during the conformational transition

For the transition from the closed to the open states, the multimer configuration is maintained so that the conformational transition of the single subunit should not interrupt the subunit interactions. Conceptual multimers of GroEL based on the final structures of the simulations are shown in Figure 8. The figure shows a dimer in the tetradecamer structure generated by superimposing the final structures onto the structure of GroEL open state. Evidently, the dimer structure derived from the result of TNM-M simulation had a clash between the two subunits, namely the arm domain of one of the subunits overlapped the other subunit. On the contrary, no serious clash was observed in the dimer structure of TNM-P. Note that the dislocation of the arm domain in TNM-M was suggested by the difference map and DynDom analyses on TNM-P and TNM-M. The direction of the rota-
location of Cα atoms and the connectivities (interactions) of systems, the differences between the M and P states are the possible steric effect of ATP on the conformation change in GroEL

Possible steric effect of ATP on the conformation change in GroEL

In our simulations, ATP molecule was removed. In the structure of P state, an ATP molecule bound to the cleft formed between the intermediate and equatorial domains of GroEL (Fig. 1). Simulation over the model after removing the ATP molecule on the P state may have resulted in the collapse of the space in the cleft and relocated the domains to the positions that were close-by. Considering this effect, there was a possibility that the movement of the intermediate domain in TNM-P had been an artifact. In order to reject the possibility of the artifact, we tested whether the final structure was capable of binding an ATP molecule. We superimposed the equatorial domain of the initial and the final structures of TNM-P and transferred the coordinates of the ATP analogue bound to the initial structure to the final structure (Fig. 9). In the final structure, ATP analogue was located in a pocket (cyan) that corresponds to the pocket of ATP analogue in the initial structure. The intermediate domain (pink) in the final structure did not overlap the ATP molecule. Hence the domain relocations we found in the simulation without ATP molecule was less likely to be the effect of the artificial removal of ATP.

Analyses of the effect of ATP on intra-subunit interaction

So far, the simulations suggested that a direct path from M state to the open state should be a dead-end and that P state should have a path to the open state. In these simulation systems, the differences between the M and P states are the location of Cα atoms and the connectivities (interactions) of Ca atoms. Hence either of them or both of them should be the factors to cause the difference in the final structures. These factors cannot be separated in the real world experiments, but can be separated in a computer simulation. By employing connectivity of M state to P state and conduct a simulation, one can test whether the initial locations of atoms or the initial intra-subunit interactions is the main factor for paving the path to the open state. We, therefore, conducted four types of simulations with CCM. The simulations are summarized in Table 1.

In order to compare the features of the computed conformational changes, we measured the same angles and distances as we did on the results with TNM (Fig. 6). The trajectories of angle θ and distance R1 in CCM simulations had a reasonable similarity to the ones in TNM simulations in the time evolution (Fig. 7A, B). However, the trajectories of distance R2 were divided into two groups (Fig. 7C). The computed structures in CCM-MM and CCM-MP had the similar features to the result of TNM-M, while the computed structures in CCM-PM and CCM-PP had features similar to the result of TNM-P. The simulations in the former group all started with the conformation in M state, whereas those in the latter group started with the conformation in P state. These results strongly suggest that the time evolution of conformational change should depend on the types of initial structures. This result indicated that the difference of connectivity between M state and P state had a minor effect on the conformational change and that the initial location of atoms dominates the time evolution of the conformation.

Role of ATP in GroEL conformational change

In the present simulations, we obtained the following results. (1) M state hardly reaches the open state, (2) P state likely reaches the open state, (3) M state with connectivity of P state hardly reaches the open state, and (4) P state with connectivity of M state likely reaches the open state. These results clearly demonstrates that the structural changes in the closed state are the major factors for the transition of GroEL from the closed to the open state, and that the changes in the interacting residues through the transition from the closed to the open states are non-essential.

The two conformations of GroEL in the closed state are likely to be reachable with each other without poring energy into its conformational change, because the difference is subtle and a normal mode analysis suggested the path between the two [15]. An ATP molecule is known to keep its γ-phosphate during the transition from the closed to the open states [5]. Hence the essence of ATP-binding is the shift in the populations of two closed conformations toward the one that is in a path leading to the open state. The ATP molecule is suggested to regulate the conformational change from closed to open states by shifting GroEL closed state to another stable state rather than by changing interactions within or outside of GroEL. This role of ATP is likely implemented by blocking paths of the conformational change of...
GroEL to M state and other possible states.

The role of ATP molecules is generally discussed over the hydrolysis of the γ-phosphate and the release of the high energy. In the binding of ATP to GroEL, hydrolysis of the molecules has long been known to have no direct relation to the conformational transition from the closed to open state [5], but without ATP, the transition will not be realized [7]. Based on the present simulation, the role of the ATP molecule is suggested as an insulator that prevents GroEL from shifting to the conformational pathway that does not reach to the open state. The current results further suggest that other molecules that resemble ATP in their structure but not in their chemical property can function as insulators for GroEL conformational change.

Acknowledgement

This work was supported by the Platform Project for Supporting in Drug Discovery and Life Science Research (Platform for Drug Discovery, Informatics, and Structural Life Science) from Japan Agency for Medical Research and Development (AMED).

Conflicts of Interest

Y. S. and K. Y. declare that they have no conflict of interest.

Author Contribution

Y. S. and K. Y. directed the entire project and co-wrote the manuscript. Y. S. performed all the simulations and analyses. K. Y. analyzed the computed structures.

References

[1] Westheimer, F. H. Why nature chose phosphates. Science 235, 1173–1178 (1987).
[2] Miller, D. L. & Westheimer, F. H. The hydrolysis of γ-phenylpropyl di- and triphosphates. J. Am. Chem. Soc. 88, 1507–1511 (1966).
[3] Kobayashi, E., Yura, K. & Nagai, Y. Distinct conformation of ATP molecule in solution and on protein. Biophysics 9, 1–12 (2013).
[4] Wellhauser, L., Luna-Chavez, C., D’Antonio, C., Tainer, J. & Bear, C. E. ATP induces conformational changes in the carboxyl-terminal region of CIC-5. J. Biol. Chem. 286, 6733–6741 (2011).
[5] Taguchi, H., Tsukuda, K., Motojima, F., Koike-Takeshita, A. & Yoshida, M. BeF3 stops the chaperonin cycle of GroEL-GroES and generates a complex with double folding chambers. J. Biol. Chem. 279, 45737–45743 (2004).
[6] Gunderson, K. L. & Kopito, R. R. Conformational states of CFTR associated with channel gating: the role of ATP binding and hydrolysis. Cell 82, 231–239 (1995).
[7] Saibil, H. R., Zheng, D., Roseman, A. M., Hunter, A. S., Watson, G. M., Chen, S., et al. ATP induces large quaternary rearrangements in a cage-like chaperonin structure. Curr. Biol. 3, 265–273 (1993).
[8] Braig, K., Otwinowski, Z., Hegde, R., Boisvert, D. C., Joachimiak, A., Horwich, A. L., et al. The crystal structure of the bacterial chaperonin GroEL at 2.8 Å. Nature 371, 578–586 (1994).
[9] Goloubinoff, P., Christeller, J. T., Gatenby, A. A. & Lorimer, G. H. Reconstitution of active dimeric ribulose bisphosphate carboxylase from an unfolded state depends on two chaperonin proteins and Mg-ATP. Nature 342, 884–889 (1989).
[10] Martin, J., Langer, T., Boteva, R., Schramel, A., Horwich, A. L. & Hartl, F. U. Chaperonin-mediated protein folding at the surface of groEL through a “molten globule”-like intermediate. Nature 352, 36–42 (1991).
[11] Jackson, G. S., Stanforth, R. A., Halsall, D. J., Atkinson, T., Holbrook J. J., Clarke, A. R., et al. Binding and hydrolysis of nucleotides in the chaperonin catalytic cycle: implications for the mechanism of assisted protein folding. Biochemistry 32, 2554–2563 (1993).
[12] Hartl, F. U. & Hayer-Hartl, M. Molecular chaperones in the cytosol: from nascent chain to folded protein. Science 295, 1852–1858 (2002).
[13] Ma, J., Sigler, P. B., Xu, Z. & Karplus, M. A dynamic model for the allosteric mechanism of GroEL. J. Mol. Biol. 302, 303–313 (2000).
[14] Teller, R., Chen, J. & Thirunamalai, D. Allotropy wiring diagrams in the transitions that drive the GroEL reaction cycle. J. Mol. Biol. 387, 390–406 (2009).
[15] Yang, Z., Májek, P. & Bahar, I. Allosteric transitions of supramolecular systems explored by network models: application to chaperonin GroEL. PLoS Comput. Biol. 4, e1000360 (2009).
[16] Berman, H. M., Westbrook, J., Feng, Z., Gilliland, G., Bhat, T. N., Weissig, H., et al. The protein data bank. Nucleic Acids Res. 28, 235–242 (2000).
[17] Wang, J. & Boisvert, D. C. Structural basis for GroEL-assisted protein folding from the crystal structure of (GroEL·K·MgATP)14 at 2.0 Å resolution. J. Mol. Biol. 327, 843–855 (2003).
[18] Xu, Z., Horwich, A. L. & Sigler, P. B. The crystal structure of the asymmetric GroEL-GroES-(ADP), chaperonin complex. Nature 388, 741–750 (1997).
[19] Tirion, M. M. Large amplitude elastic motions in proteins from a single-parameter, atomic analysis. Phys. Rev. Lett. 77, 1905–1908 (1996).
[20] Matsumoto, A., Kamata, T., Takagi, J., Iwasaki, K. & Yura, K. Key interactions in integrin ectodomain responsible for global conformational change detected by elastic network normal-mode analysis. Biophys. J. 95, 2895–2908 (2008).
[21] Rader, A. J., Chennubhotla, C., Yang, L. W. & Bahar, I. The Gaussian network model: Theory and applications. in Normal mode analysis: theory and applications to biological and chemical systems. (Cui, Q. & Bahar, I. eds.), pp. 41–64 (Chapman & Hall CRC Press, Chicago, IL, 2006).
[22] Rotkiewicz, P. & Skolnick, J. Fast procedure for reconstruction of full-atom protein models from reduced representations. J. Comput. Chem. 29, 1460–1465 (2008).
[23] Kabsch, W. & Sander, C. Dictionary of protein secondary structure: pattern recognition of hydrogen-bonded and geometrical features. Biopolymers 22, 2577–2637 (1983).
[24] Hayward, S. & Berendsen, H. J. Systematic analysis of domain motions in proteins from conformational change: new results on citrate synthase and T4 lysozyme. Proteins 30, 144–154 (1998).
[25] Ranson, N. A., Farr, G. W., Roseman, A. M., Gowen, B., Fenton, W. A., Horwich, A. L., et al. ATP-bound states of GroEL captured by cryo-electron microscopy. Cell 107, 869–879 (2001).
[26] The PyMOL molecular graphics system, version 1.2r.pre. Schrödinger, LLC.