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Markov Methods for Hierarchical Coarse-Graining of Large Protein Dynamics

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ABSTRACT

Elastic network models (ENMs) and, in particular, the Gaussian Network Model (GNM) have been widely used in recent years to gain insights into the machinery of proteins. The extension of ENMs to supramolecular assemblies presents computational challenges, because of the difficulty in retaining atomic details in mode decomposition of large protein dynamics. Here, we present a novel approach to address this problem. We rely on the premise that, all the residues of the protein machinery (network) must communicate with each other and operate in a coordinated manner to perform their function successfully. To gain insight into the mechanism of information transfer between residues, we study a Markov model of network communication. Using the Markov chain perspective, we map the full-atom network representation into a hierarchy of ENMs of decreasing resolution, perform analysis of dominant communication (or dynamic) patterns in reduced space(s) and reconstruct the detailed models with minimal loss of information. The communication properties at different levels of the hierarchy are intrinsically defined by the network topology. This new representation has several features, including: soft clustering of the protein structure into stochastically coherent regions thus providing a useful assessment of elements serving as hubs and/or transmitters in propagating information/interaction; automatic computation of the contact matrices for ENMs at each level of the hierarchy to facilitate computation of both Gaussian and anisotropic fluctuation dynamics. We illustrate the utility of the hierarchical decomposition in providing an insightful description of the supramolecular machinery by applying the methodology to the chaperonin GroEL-GroES.

Key words: information propagation, Markov process, network model.

1. INTRODUCTION

WITH ADVANCES IN SEQUENCE and structure genomics, an emerging view is that to understand and control the mechanisms of biomolecular function, knowledge of sequence and structure is insufficient. Additional knowledge in the form of *dynamics* is needed. In fact, proteins do not function as static entities or in isolation; they are engaged in functional motions, and interactions, both within and between molecules. The resulting motions can range from single amino acid side chain reorientations (*local*) to

concerted domain-domain motions (*global*). The motions on a local scale can be explored to a good approximation by conventional molecular dynamics (MD) simulations, but the motions at a global scale are usually beyond the range of such simulations. Elastic network models (ENM), based on polymer mechanics, succeed in providing access to global motions (Bahar and Rader, 2005; Rader et al., 2005; Ma, 2005).

A prime example of an EN is the Gaussian Network Model (GNM) (Bahar et al., 1997; Haliloglu et al., 1997). In graph-theoretic terms, each protein is modeled by an undirected graph \mathcal{G} , given by $\mathcal{G} = (\mathcal{V}, \mathcal{E})$, with residues $\mathcal{V} = \{v_i \mid i = 1, ..., n\}$ defining the nodes of the network, and edges $\mathcal{E} = \{e_{ij}\}$ representing interactions between residues v_i and v_j . The set of all pairwise interactions is described by a non-negative, symmetric affinity matrix $A = \{a_{ij}\}$, with elements $a_{ij} = a_{ji}$. GNM chooses a simple interaction model, which is to set the affinity $a_{ij} = a_{ji} = 1$, for a pair of residues v_i and v_j whose C^{α} atoms are within a cut-off distance of r_c . The interactions represent both bonded and non-bonded contacts in the native configuration of the protein. The cutoff distance represents the radius of the first coordination shell around residues observed in Protein Data Bank (PDB) (Berman et al., 2000) structures and is set to be 7 Å (Bahar and Jernigan, 1997; Miyazawa and Jernigan, 1985).

The motions accessible under native state conditions are obtained from the Kirchhoff matrix Γ , defined in terms of the affinity and degree matrices as $\Gamma = D - A$. Here D is a diagonal matrix: $D = \operatorname{diag}(d_1, \ldots, d_n)$ and d_j represents the degree of a vertex v_j : $d_j = \sum_{i=1}^n a_{ij} = \sum_{i=1}^n a_{ji}$. Γ is referred to as the combinatorial Laplacian in graph theory (Chung, 1997). The Kirchhoff matrix multiplied by a force constant γ that is uniform over all springs defines the stiffness matrix of an equivalent mass-spring system. The eigenvalue decomposition of Γ yields the shape and frequency dispersion of equilibrium fluctuations. In most applications, it is of interest to extract the contribution of the most cooperative modes, i.e., the low frequency modes that have been shown in several systems to be involved in functional mechanisms (Bahar and Rader, 2005; Ma, 2005). Also, of interest is the inverse of Γ , which specifies the covariance matrix for the Boltzmann distribution over equilibrium fluctuations.

GNM is a linear model, and as such it cannot describe the transition between configurations separated by an energy barrier (or any other non-linear effect), so it only applies to fluctuations in the neighborhood of a single energy minimum. The energy well is approximated by a harmonic potential, which limits the magnitude of the predicted motion. The topology of inter-residue contacts in the equilibrium structure is captured by the Kirchhoff matrix Γ . Also, there is no information on the 'directions' of motions in different vibrational modes, but on their sizes only. The fluctuations are assumed to be isotropic and Gaussian, but for anisotropic extension of GNM called ANM see (Atilgan et al., 2001; Doruker et al., 2000) or equivalent EN-based normal mode analyses (NMA) (Hinsen, 1998; Tama and Sanejouand, 2001). Despite this simplicity, many studies now demonstrate the utility of GNM and other EN models in deducing the machinery and conformational dynamics of large structures and assemblies (Bahar and Rader, 2005).

The application and extension of residue-based ENMs to more complex processes, or larger systems, is computationally expensive, both in terms of memory and time, as the eigen decomposition scales on the order of $O(n^3)$, where n is the number of nodes in the graph. Given that the Kirchhoff matrix is sparse, there are a plethora of efficient sparse eigensolvers that one can use (Barnard and Simon, 1994; Fowlkes et al., 2004; Lehoucq et al., 1996; Simon and Zhou, 2000), including eigensolvers designed specifically for decomposing graph Laplacians (Koren et al., 2003).

Another way to reduce complexity is to adopt coarser-grained models. For example, in the hierarchical coarse-graining (HCG) approach, sequences of m consecutive amino acids are condensed into unified nodes, which reduces the computing time and memory by factors of m^3 and m^2 , respectively (Doruker et al., 2002); or a mixed coarse-graining has been proposed in which the substructures of interest are modeled at single-residue-per-node level and the surrounding structural units at a lower resolution of m-residues-per node (Kurkcuoglu et al., 2004); another common representation of the structure is to adopt rigidly translating and rotating blocks (RTB) (Marques, 1995; Tama et al., 2000), or the so-called block normal mode analysis (BNM) (Li and Cui, 2002).

While these methods have been useful in tackling larger systems, the choice and implementation of optimal model parameters to retain physically significant interactions at the residue-, or even atomic level, has been a challenge. The level of HCG has been arbitrarily chosen in the former group of studies, requiring *ad-hoc* readjustments to spring constants or cutoff distances of interaction. In the case of RTB or BNM approaches, all atomic, or residue level information is lost, and substructures that may contain internal degrees of freedom—some of which being functional—are assumed to move as a rigid block. Overall, information is lost on local interactions as structures are coarse-grained. Clearly, the challenge is to map

a high resolution model to a low resolution, with a minimal loss of information. In this paper, we present a novel approach to address this problem.

Our approach is to model structures as networks of interacting residues and study the Markov propagation of "information" across the network. We rely on the premise that, the components (residues) of a protein machinery (network) communicate with each other and operate in a coordinated manner to perform their function successfully. Using the Markov chain perspective, we map the full atom network representation into a hierarchy of intermediate ENMs, while retaining the Markovian stochastic characteristics, i.e., transition probabilities and stationary distribution, of the original network. The communication properties at different levels of the hierarchy are intrinsically defined by the network topology. This new representation has several features, including: soft clustering of the protein structure into stochastically coherent regions thus providing a useful assessment of elements serving as hubs and/or transmitters in propagating information/interaction; automatic computation of the contact matrices for ENMs at each level of the hierarchy to facilitate computation of both Gaussian and anisotropic fluctuation dynamics; and a fast eigensolver for NMA. We illustrate the utility of the hierarchical decomposition by presenting its application to the bacterial chaperonin GroEL–GroES.

2. A MORKOV MODEL FOR NETWORK COMMUNICATION

We model each protein as a weighted, undirected graph \mathcal{G} given by $\mathcal{G} = (\mathcal{V}, \mathcal{E})$, with residues $\mathcal{V} = \{v_i \mid i = 1, ..., n\}$ defining the nodes of the network, and edges $\mathcal{E} = \{e_{ij}\}$ representing interactions between residues v_i and v_j . The set of all pairwise interactions is described by a non-negative, symmetric affinity matrix $A = \{a_{ij}\}$, with elements $a_{ij} = a_{ji}$ and where a_{ij} is defined as

$$a_{ij} = \frac{N_{ij}}{\sqrt{N_i N_j}}. (1)$$

Here N_{ij} is the is the total number of atom-atom contacts made within a cutoff distance of $r_c=4$ Å between residues v_i and v_j and (N_i, N_j) are the total number of heavy atoms in the individual residues (v_i, v_j) . The self-contact a_{ii} is similarly defined. This representation captures to a first approximation the strong (weak) interactions expected to take place between residue pairs with large (small) numbers of atom-atom contacts, and removes biases due to size effects. The degree of a vertex v_j is defined as $d_j = \sum_{i=1}^n a_{ij} = \sum_{i=1}^n a_{ji}$, which are organized in a diagonal matrix of the form $\mathbf{D} = \operatorname{diag}(d_1, \ldots, d_n)$. A discrete-time, discrete-state Markov process of network communication is defined by setting the communication (or signalling) probability m_{ij} from residue v_j to residue v_i in one time-step to be proportional to the affinity between nodes, $a_{i,j}$. In matrix notation, this conditional probability matrix $\mathbf{M} = \{m_{ij}\}$, also called the Markov transition matrix, given by

$$\boldsymbol{M} = \boldsymbol{A} \boldsymbol{D}^{-1}, \tag{2}$$

defines the stochastics of a random walk on the protein graph \mathcal{G} . Note, $m_{ij} = d_j^{-1}a_{ij}$ where d_j gives a measure of local packing density near residue v_j and serves as a normalizing factor to ensure $\sum_{i=1}^n m_{ij} = 1$. Alternatively, m_{ij} can be viewed as the conditional probability of interacting with residue v_i , that is transmitting information to residue v_i , given that the signal (or perturbation) is initially positioned, or originates from, v_j . Suppose this initial probability is p_j^0 . Then, the probability of reaching residue v_i using link e_{ij} is $m_{ij} p_j^0$. In matrix notation, the probability of ending up on any of the residues $\mathbf{v} = [v_1, v_2, \dots, v_n]$ after one time step is given by the distribution $\mathbf{p}^1 = \mathbf{M} \mathbf{p}^0$, where $\mathbf{p}^k = [p_1^k, \dots, p_n^k]$. Clearly this process can be iterated, so that after β steps we have

$$p^{\beta} = M^{\beta} p^{0}. \tag{3}$$

Assume the graph is connected, i.e. there is a path connecting every pair of residues in the graph. Then, as $\beta \to \infty$ the Markov chain p^{β} approaches a unique *stationary* distribution π , the elements of which are given by: $\pi_i = d_i / \sum_{k=1}^n d_k$. While the evolution of the random walk is a function of the starting

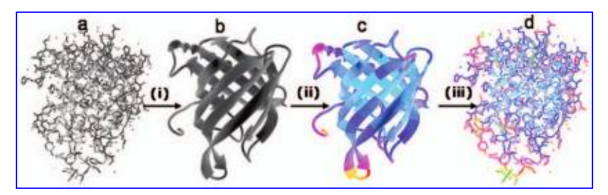


FIG. 1. Hierarchical network decomposition overview. Step (i) map the structure (a) to its optimal reduced level representation (illustrated here for retinol-binding protein mapped from full atomic scale to intermediate-chain representation). This step may involve several intermediate levels of resolution (b) (Fig. 2); step (ii) perform structural analysis (e.g. GNM) at a coarse-grained scale (c); and step (iii) reconstruct the detailed structure-dynamics (d). The communication/coupling of residues at a given level are assumed to obey a Markov process controlled by atom-atom contact topology. The steps (i) and (iii) are achieved by two operators, R for model reduction, and R for model reconstruction. R and R ensure that similar stochastic characteristics (transition probabilities and stationary distributions) are retained between successive levels of the hierarchy.

distribution, the stationary distribution is invariant to the precise details of how the random walk is initiated.

The main goal in undertaking random walks is to reveal the communication patterns *inherent* to the network because of its architecture. However, a naive random walk on a large protein, as will be presented below for the GroEL-GroES complex, is computationally challenging. We address this problem by building a hierarchy of intermediate resolution network models, performing the analysis in the reduced space and mapping the results back to the high resolution representation as illustrated in Figure 1.

3. NETWORK HIERARCHY TO REDUCE COMMUNICATION COMPLEXITY

The objective in designing a network hierarchy is to map the Markov process operating at the highest resolution onto successively lower resolution network models, while maintaining its stochastic characteristics (Chennubhotla and Jepson, 2005). In particular, using the stationary distribution π and the Markov transition matrix M, we build a coarse-scale Markov propagation matrix \widetilde{M} (size: $m \times m$, where $m \ll n$) and its stationary distribution δ . The random walk initiated on the coarse-scale network $\widetilde{\mathcal{G}}(m)$, and reaching distribution δ , is equivalent to the random walk on the full resolution network $\mathcal{G}(n)$ with stationary distribution π . To build a hierarchy of intermediate resolution networks we devise two sets of new operators at each level of the hierarchy: R for model reduction, and K for model expansion/reconstruction.

3.1. Deriving stationary distribution in the reduced model

We begin by expressing the stationary distribution $\pi = [\pi_1, \pi_2, \dots, \pi_n]$ as a probabilistic mixture of *latent* distributions,

$$\pi = K\delta, \tag{4}$$

where $\delta = [\delta_1, \delta_2, \dots, \delta_m]$ is an unknown stationary distribution in a reduced (*m*-dimensional) representation of the structure; $K = \{K_{ij}\}$ is an $n \times m$ non-negative *kernel* matrix with elements K_{ij} and columns K_j being latent probability distributions that each sum to 1, and $m \ll n$. The kernel matrix acts as an *expansion* operator, mapping the low-dimensional distribution δ to a high-dimensional distribution π .

We derive a maximum likelihood approximation for δ using an expectation-maximization (EM) type algorithm (McLachlan and Basford, 1988). To this aim we minimize the *Kullback-Liebler* distance measure (Kullback, 1959; Kullback and Leibler, 1951) between the two probability distributions π and $K\delta$, subject

to the constraint that $\sum_{j=1}^{m} \delta_j = 1$ and ensured by the Lagrange multiplier λ in the equation below:

$$E = -\sum_{i=1}^{n} \pi_i \ln \sum_{j=1}^{m} K_{ij} \delta_j + \lambda \left(\sum_{j=1}^{m} \delta_j - 1 \right).$$
 (5)

Setting the derivative of E with respect to δ_i to be zero we obtain

$$\sum_{i=1}^{n} \frac{\pi_i K_{ij} \delta_j}{\sum_{k=1}^{m} K_{ik} \delta_k} = \lambda \delta_j.$$
 (6)

The contribution made by kernel j to a node i (or its stationary probability π_i) is given by K_{ij} (or the product $K_{ij}\delta_j$), and hence we can define an *ownership* of node i in the high resolution representation by a node j in the low resolution representation as

$$R_{ij} = \frac{K_{ij}\delta_j}{\sum_{k=1}^{m} K_{ik}\delta_k}.$$
 (7)

 R_{ij} is also referred to as the responsibility of node j in the low resolution representation, for node i in the high resolution. We note that the mapping between the two resolutions is not deterministic, but probabilistic in the sense that $\sum_{j=1}^{m} R_{ij} = 1$.

Using this relation, and the equalities $\sum_{j=1}^{m} \delta_j = 1$ and $\sum_{i=1}^{n} \pi_i = 1$, summing over j in Equation (6) gives $\lambda = 1$. This further leads to the stationary distribution δ at the coarse scale

$$\delta_j = \sum_{i=1}^n \pi_i R_{ij}. \tag{8}$$

The matrix R therefore maps the high dimensional distribution π to its low-dimensional counterpart δ and hence the name *reduction* operator. Following Bayes theorem, K_{ij} can be related to the *updated* δ values as

$$K_{ij} = \frac{R_{ij}\pi_i}{\delta_i}. (9)$$

In summary, the operators K and R and stationary distribution δ are computed using the following EM type procedure: (1) select an initial estimate for K and δ (see Section 3.2); (2) E-step: compute ownership maps R using Equation (7); (3) M-step: estimate δ and update K using Equations (8) and (9) respectively; and finally, (4) repeat E- and M- steps until convergence.

3.2. Kernel selection details

As an initial estimate for δ , a uniform distribution is adopted. The kernel matrix K is conveniently constructed by diffusing M to a small number of iterations β to give M^{β} and selecting a small number of columns. In picking the columns of M^{β} , a greedy decision is made. In particular, column i in M^{β} corresponds to information diffusion from residue v_i . The first kernel K_i that is picked corresponds to the residue v_i with the highest stationary probability π_i . Following the selection of K_i , all other residues j (and the corresponding columns K_j in M^{β}) that fall within the half-height of the peak value of the probability distribution in K_i are eliminated from further consideration. This approach generates kernels that are spatially disjoint. The selection of kernels continues until every residue in the protein is within a half-height of the peak value of at least one kernel. While other kernel selection procedures are conceivable, we chose the greedy method for computational speed. In practice, we observed the EM algorithm generates results of biological interest that are insensitive to the initial estimates of K and δ .

3.3. Transition and affinity matrices in the reduced model

The Markov chain propagation at the reduced representation obeys the equation $q^{k+1} = \widetilde{M} q^k$, where q^k is the coarse scale m-dimensional probability distribution after k steps of the random walk. We expand q^k into the fine scale using $p^k = K q^k$, and reduce p^k back to the coarse scale by using the ownership value $R_{i,j}$ as in $q_j^{k+1} = \sum_{i=1}^n p_i^k R_{i,j}$. Substituting Equation (7) for ownerships, followed by the expression for p^k , in the equation for q_j^{k+1} , we obtain

$$\widetilde{\boldsymbol{M}} = \operatorname{diag}(\boldsymbol{\delta}) \, \boldsymbol{K}^{\mathsf{T}} \operatorname{diag}(\boldsymbol{K}\boldsymbol{\delta})^{-1} \, \boldsymbol{K}. \tag{10}$$

Using the definition of \widetilde{M} , and the corresponding stationary distribution δ , we generate a *symmetric* affinity matrix \widetilde{A} that describes the node-node interaction strength in the low resolution network

$$\widetilde{A} = \widetilde{M}\operatorname{diag}(\delta). \tag{11}$$

To summarize, we use the stationary distribution π and Markov transition matrix M at the fine-scale to derive the operator K and associated reduced stationary distribution δ , using the EM algorithm described in the previous section. K and δ are then used in Equations (10) and (11) to derive the respective transition \widetilde{M} and affinity \widetilde{A} matrices in the coarse-grained representation. Clearly, this procedure can be repeated recursively to build a hierarchy of lower resolution network models.

4. HIERARCHICAL DECOMPOSITION OF THE CHAPERONIN GroEL-GroES

We examine the structure and dynamics of the bacterial chaperonin complex GroEL-GroES-(ADP)₇ (Xu et al., 1997), from the perspective of a Markov propagation of information/interactions. GroEL is a cylindrical structure, 150 Å long and 140 Å wide, consisting of 14 identical chains organized in two back-to-back stacked rings (*cis* and *trans*) of seven subunits each. The GroES co-chaperonin, also heptameric, binds to the apical domain of GroEL and closes off one end of the cylinder. During the allosteric cycle that mediates protein folding, the *cis* and *trans* rings alternate between open (upon binding of ATP and GroES) and closed (unliganded) forms, providing access to, or release from, the central cylindrical cavity, where the folding of an encapsulated (partially folded or misfolded) protein/peptide is assisted.

First, the inter-residue affinity matrix A based on all atom-atom contacts is constructed (Fig. 2a), from which the fine-scale Markov transition matrix M is derived using Equation (2). The kernel selection algorithm applied to M^{β} ($\beta=4$) yields 1316 (reduced level 1) kernels. Using these kernels as an initialization, a recursive application of the EM procedure derives stationary distributions δ (Equation (8)), updated expansion matrices K (Equation (9)), reduced level probability transition matrices M (Equation (10)) and the corresponding residue interaction matrices M (Equation (11)). The respective dimensions of M

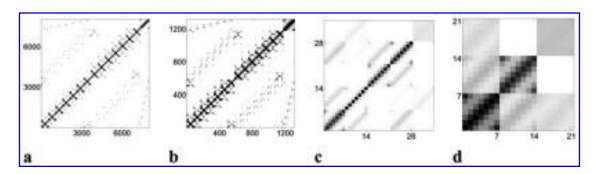


FIG. 2. Affinity matrix hierarchy for the protein GroEL/GroES (PDB: **1AON**). The respective sizes of the reduced models, and the associated affinity matrices, across the hierarchy are n = 8015 (fine-scale, **a**) and m = 1316 (coarse-scale 1, **b**), 483 (coarse-scale 2), 133 (coarse-scale 3), 35 (coarse-scale 4, **c**) and 21 (coarse-scale 5, **d**). The affinity matrices are real-valued but are shown here as *dot* plots (**a,b**), to highlight the similarity in the matrix structure across the hierarchy. The affinity matrices for the two lowest resolution models (**c,d**) are shown as images, where the affinity value is *inversely* proportional to the brightness of a pixel.

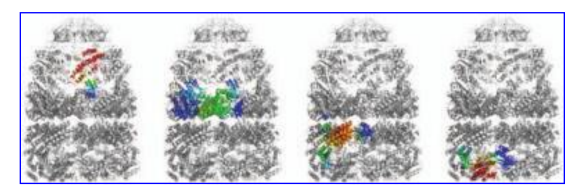


FIG. 3. Four different soft clusters located on GroEL.

turn out to be 483 (reduced level 2), 133 (reduced level 3), 35 (reduced level 4, Fig. 2c) and 21 (reduced level 5, Fig. 2d). We note that the individual subunits of the GroEL/GroES are distinguished by their strong intra-subunit interactions, and a number of inter-subunit contacts are maintained at all levels, which presumably establish the communication across the protein at all levels. The dimension m of the reduced model is automatically defined during the kernel selection at each level of the hierarchy. The method thus avoids the arbitrary choices of sampling density and interaction cutoff distances at different hierarchical levels.

In contrast to the *deterministic* assignment of one-node-per-residue in the original ENM, the Markovchain-based representation adopts a stochastic description in the sense that each node probabilistically "owns," or "is responsible for" a subset of residues. To see this, consider the ownership matrix $R^{(l,l+1)}$ $\{R_{ij}^{(l,l+1)}\}$ that relates information between two adjacent levels l and l+1 of the hierarchy. Likewise, the matrix $\mathbf{R}^{(0,L)} = \prod_{l=0}^{L-1} \mathbf{R}^{(l,l+1)}$ ensures the passage from the original high resolution representation 0 to the top level L of the hierarchy. In particular, the ijth element $\mathbf{R}_{ij}^{(0,L)}$ describes the probabilistic participation of residue v_i (at level 0) in the cluster j (at level L), and $\sum_j \mathbf{R}_{ij}^{(0,L)} = 1$. Hence, the nodes at level L perform a soft partitioning of the structure. This type of soft distribution of residues among the m nodes, or their partial/probabilistic participation in neighboring clusters, establishes the communication between the clusters, and is one of the key outcomes of the present analysis. Of interest is to examine the ownership of clusters at a reduced representation. We select the coarse-scale 4, for example, which maps the structure into a graph of 35 clusters (Fig. 2c). Figure 3 demonstrates the ownership of the individual clusters at this level. Essentially there are five sets of seven clusters each, centered near the apical and equatorial domains of the cis and trans rings, and at the individual GroES chains. The intermediate domains are being shared between the clusters at the apical and equatorial domains. As such, they play a key role in establishing intra-subunit communication. The color-coded ribbon diagrams in Figure 3 display the loci of representative clusters from each of these four distinct types (excluding the GroES clusters). The color code from red-to-blue refers to the higher-to-lower involvement (or responsibility) of the individual residues in the indicated clusters. Evidently, the regions colored red serve as hubs for broadcasting the information within clusters, and those colored blue play the key role of establishing the communication, or transferring information between clusters. Detailed examination of the ownership of these clusters reveal several interesting features, correlating with the experiments and summarized in Section 6.

Next, we benchmark the utility and robustness of the presently introduced methodology in so far as the equilibrium dynamics of the examined structure is concerned. Mainly, we compare the collective modes of motion predicted for the GroEL-GroES complex using a full-residue (8015 nodes) ENM (Keskin et al., 2002), with those captured by the hierarchy of reduced models. The newly introduced representation hierarchy will be shown below to successfully map structure-dynamics information between successive levels with minimal loss in accuracy.¹

¹The ownership matrix can also be used to propagate the location information of the residues from one level of the hierarchy to another. This in turns help perform anisotropic fluctuation modeling, a procedure that we plan to elaborate in future study.

5. HIERARCHICAL GAUSSIAN NETWORK MODEL

Here we present a methodology for generating GNM modes at different levels of coarse-graining the information on contact topology inherent in \mathcal{G} , and reconstructing the detailed mode behavior by projecting the eigenvectors and eigenvalues generated at low levels of resolution back to their fine scale counterparts using the Markov chain propagation formalism, a method shortly referred to as hierarchical GNM (hGNM).

For hGNM, assume that the dimensions of the Kirchhoff matrices at the coarse, intermediate and fine scales are e, m, and n, respectively, where $e \le m \ll n$. The affinity and Kirchhoff matrices at the coarsest level are not likely to be sparse, however a full eigen decomposition of the coarsest Kirchhoff matrix (size: $e \times e$) will be computationally the least expensive step.

To reconstruct the eigen information at the fine-scale, assume we have access to the leading eigenvectors \widehat{U} (size: $m \times e$) for $\widehat{\Gamma}$ (size: $m \times m$). Using this we generate the leading eigenvectors \widetilde{U} (size: $n \times e$), and the leading eigenvalues $\widetilde{\mathbf{\Lambda}} = [\lambda_1, \lambda_2 \cdots \lambda_e]$ (size: $e \times 1$) of the fine-scale Kirchhoff matrix Γ (size: $n \times n$). Let $\{U, \Lambda\}$ denote the eigenvectors and eigenvalues obtained from a direct decomposition of Γ . There are several steps to the eigen reconstruction process. (i) The coarse-scale eigenvectors \widehat{U} can be transformed using the kernel matrix K as $\widetilde{U} = K \widehat{U}$ to generate \widetilde{U} as an approximation to U. (ii) This transformation alone is unlikely to set the directions of \widetilde{U} exactly aligned with U. So, we update the directions in \widetilde{U} by repeated application of the following iteration (called *power* iterations (Watkins, 2002)): $\widetilde{U} \Leftarrow \Gamma_g \widetilde{U}$ Note, here instead of using Γ we use an adjusted matrix Γ_g given by $\Gamma_g = \nu I - \Gamma$, where ν is a constant and Iis an identity matrix. The power iterations will direct the eigenvectors to directions with large eigenvalues. But for fluctuation dynamics, we are interested in the slow eigen modes with small eigenvalues, hence the adjustment Γ_g is made. In particular, because of Gerschgorin disk theorem (Watkins, 2002) the eigenvalues of Γ are bound to lie in a disk centered around the origin with a radius ν that is no more than twice the largest element on the diagonal of Γ . (iii) Steps i and ii need not preserve orthogonality of the eigenvectors in U. We fix this by a Gram-Schmidt orthogonalization procedure (Watkins, 2002). Finally, the eigenvalues are obtained from $\widetilde{\Lambda} = \operatorname{diag}(\widetilde{U}^{\mathsf{T}} \Gamma \widetilde{U})$. In Chennubhotla and Jepson (2005), we present more details of this coarse to fine eigen mapping procedure, including a discussion on the number of power iterations to use; setting the thresholds for convergence and a comparison of the speed ups obtained over a standard sparse eigensolver for large matrices.

5.1. Collective dynamics in the reduced space: benchmarking against GNM

As discussed earlier, the eigenvalue decomposition of Γ yields the shape and frequency dispersion of equilibrium fluctuations. The shape of mode k refers to the normalized distribution of residue displacements along the principal axis k, given by the elements $\boldsymbol{u}_i^{(k)}$ ($1 \le i \le n$) of the kth eigenvector $\boldsymbol{u}^{(k)}$, and the associated eigenvalue λ_k scales with the frequency of the kth mode. In most applications, it is of interest to extract the contribution of the most cooperative modes, i.e., the low frequency modes that have been shown in several systems to be involved in functional mechanisms. To this end, we used the Markov-chain based hierarchy to build reduced Kirchhoff matrices $\widetilde{\Gamma}$ at increasingly lower levels of resolution. We then performed their mode decompositions and propagated the information back over successive levels of the hierarchy, so as to generate the eigenvectors and eigenvalues for the fine-scale Kirchhoff matrix Γ . We now show that hGNM maps the structure-dynamics information between successive levels of the hierarchy with minimal loss in accuracy.

First, our previous study identified ten slowest modes of interest, including the counter-rotation of the two rings around the cylindrical axis (non-zero mode 1) and other collective deformations proposed to be involved in chaperonin function (Keskin et al., 2002). Results presented in Figure 4 show the mechanism of the dominant mode, mainly a global twisting of the structure where the cis and trans undergo counter rotation about the cylindrical axis (mode 1). The most important point is that these results corroborate previous findings (Keskin et al., 2002; Ma, 2005) and are reproduced here by adopting a reduced representation down to m = 21 nodes and mapped back to full-residue level.

Second, Figure 5a compares the frequencies obtained by the full-residue-level representation, with those obtained by hGNM, upon propagation of the topology information from reduced level 4 (Fig. 2c). An excellent agreement is observed between the reconstructed eigenvalues $\widetilde{\lambda}$ (red curve) and their original values λ (open circles). In Figure 5b, we display the correlation cosine between the eigenvectors $\boldsymbol{u}^{(k)}$ and

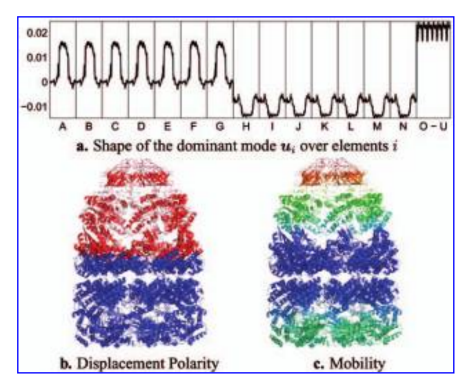


FIG. 4. Dominant mode shape and mobility. (a) The labels on the abscissa indicate the chain identities, A-G belong to the *cis* ring, H-N come from the *trans* ring and O-U are from the GroES cap. The black curve gives the shape of the *slowest* eigen mode. The ordinate value is the normalized distribution of residue displacements along the dominant mode coordinate. (b) Ribbon diagram illustrating the polarity of the displacement, color coded to be red for positive and blue for negative, indicating the anticorrelated motions of the two halves of the complex. (c) Ribbon diagram color-coded after residue mobilities in mode 1. The mobility of residue v_i given by the squared displacement: $(u_i^{(1)})^2$, with a color code that is red for high and blue for low.

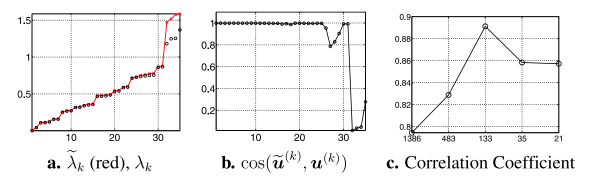


FIG. 5. (a) hGNM results comparing eigenvalues λ (circles) from a direct decomposition of the Γ with multi-scale eigensolver spectrum $\widetilde{\lambda}$ (red line). For the direct eigen decomposition, we use the Matlab program svds.m which invokes the compiled ARPACKC routine (Lehoucq et al., 1996), with a default convergence tolerance of 1e-10. (b) Mode shape correlation: diag($|\widetilde{U}^T U|$), between the matrix of eigenvectors \widetilde{U} derived by hGNM and U from direct decomposition. (c) Correlation coefficient between the theoretical B-factors (derived at each level of the hierarchy) versus experiment. The abscissa labels indicate the size m of the network at successive levels of the hierarchy.

 $\widetilde{u^{(k)}}$ obtained by the full-residue representation and the reconstruction from reduced level 4 respectively. Notably, the reduced representation contains only 35 nodes. Yet, the correlation cosine with the detailed representation containing 8015 nodes is almost unity throughout all the leading 25 modes, and above 0.8 for all modes, except the terminal four modes. The contribution of the latter to the overall dynamics is negligibly small compared to the large group of slow modes.

Finally, in order to assess the effect of coarse-graining on fluctuation dynamics, we compared in Figure 5c the mean-square fluctuations obtained from different levels of the hierarchy with the experimental B-factor values. The theoretical B-factor for each residue v_i is computed using (Kundu et al., 2002)

$$B_{i} = \frac{8\pi^{2}k_{B}T}{\gamma} \sum_{k=2}^{n} \lambda_{k}^{-1} \left(\mathbf{u}_{i}^{(k)} \right)^{2}, \tag{12}$$

where the summation is performed over all n-1 modes in the GNM, or over all the m-1 reduced eigenvectors and eigenvalues reconstructed from different levels of the hierarchy in hGNM. Because experimental B-factors correspond to each atom and our representation at the fine-scale is a summary of atom-atom contact information for each residue, we average the experimental B-factors over all atoms for each residue. As shown in Figure 5c, a correlation coefficient value of 0.86 is achieved between the experimental and theoretical B-factors after mapping the structure of 8015 residues into a representative network of 21 nodes. Thus, the fluctuation behavior of individual residues is accurately maintained despite a drastic reduction in the complexity of the examined network. Interestingly, a maximum in correlation coefficient is obtained at an intermediate level of resolution, m=133, which may be attributed to an optimal elimination of noise in line with the level of accuracy of experimental data at this level of representation.

6. DISCUSSION

A new method is introduced in the present study, which permits us to use structural information at atomic level in building network representations of different complexity, which lend themselves to efficient analysis of collective dynamics and information propagation stochastics. The approach is particularly useful for analyzing large structures and assemblies, or cooperative/allosteric processes that are usually beyond the range of conventional molecular simulations.

We illustrated the utility of the methodology by way of application to the chaperonin GroEL-GroES, a widely studied structure composed of n=8015 residues. Notably, we start with the full-atomic representation of the complex, which involves a total of $\approx 10^6$ atom-atom contacts (based on an interaction range of 4 Å). Interatomic contacts define the affinities of pairs of residues, which are, in turn, used to define the weights of the connectors between residues (nodes) in the graph/network representation of the structure. The affinities also define the conditional probabilities of information transfer across residues following a Markovian process. The original network of n nodes is mapped into lower dimensional representations, down to m=21 nodes, by an EM algorithm that maintains two basic properties of the original stochastic process: its Markovian conditional probabilities and stationary distribution (i.e. communication probability/potential) of individual residues. Two sets of operators, ensuring model reduction and reconstruction at different hierarchical levels permit us to perform the analysis at reduced scales but reconstructing the behavior.

A major utility of the present study is to identify the nodes (clusters of residues) that control the dynamics of the structure-reduced levels. First, these are soft nodes: they do not necessarily contain/own well-defined residues, but exhibit a probabilistic ownership, i.e. some residues exhibit a high responsibility in these clusters, while others contribute only partially, being shared by neighboring clusters. The identity of residues lying in the two groups is crucially important for assessing the residues that can serve as hubs and messengers of information, respectively. Careful examination of the ownership of clusters at level 4 (Fig. 3) revealed for example several interesting features such as (Chennubhotla and Bahar, 2006): (i) the incorporation of the GroES mobile loop residues Glu18-Ala33 into the adjoining clusters centered at the apical domains of *cis* ring subunits, thus providing evidence for the critical role of this loop in establishing the communication between the chaperone and co-chaperonin, consistent with several experimental studies (Hohfeld and Hartl, 1994; Kovalenko et al., 1994; Landry et al., 1993; Richardson and Georgopoulos, 1999;

Richardson et al., 1999, 2001; Shewmaker et al., 2001), (ii) the identity of amino acids that play a key role in establishing the intra-ring, inter-subunit (positive) cooperativity in the *cis* ring, mainly Val38-Ile49 on the equatorial domains, and Asp179-Leu183 and Val381-Lys392 on the intermediate domains of the subunits, (iii) the stronger inter-subunit coupling between the apical domain residues of the subunits in the *trans* ring, compared to the *cis* ring, and in particular the involvement of the previously pointed out (Braig et al., 1994; Ma et al., 2000) inter-subunit salt bridge Glu386-Arg197 in this stabilizing effect evidenced by the participation of these residues in the same cluster, (iv) the tendency of the *cis* and *trans* ring subunits to incorporate into their clusters, their clockwise and counter-clockwise neighbors, respectively, when viewed from the cap. The last property is particularly interesting in view of the known negative cooperativity between the two rings (Yifrach and Horovitz, 1995) and the tendency of the two rings to rotate in opposite directions (Fig. 4) in the most cooperative GNM mode, a feature that is also suggested by comparing the structures of the complex at different states of the chaperonin cycle (Saibil and Ranson, 2002).

The correlations observed between the present results and those from the previous experimental studies, as well as more detailed computations, show promise in the utility of the present methodology as a tool for exploring both the collective dynamics and the communication stochastics of complex structures, including those derived from cryo-EM.

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REFERENCES

Atilgan, A.R., Durell, S.R., Jernigan, R.L., et al. 2001. Anisotropy of fluctuation dynamics of proteins with an elastic network model. *Biophys. J.* 80, 505.

Bahar, I., Atilgan, A.R., and Erman, B. 1997. Direct evaluation of thermal fluctuations in protein using a single parameter harmonic potential. *Folding Design* 2, 173–181.

Bahar, I., and Jernigan, R.L. 1997. Inter-residue potentials in globular proteins and the dominance of highly specific hydrophilic interactions at close separation. *J. Mol. Biol.* 266, 195.

Bahar, I., and Rader, A.J. 2005. Coarse-grained normal mode analysis in structural biology, *Curr. Opin. Struct. Biol.* 15, 586–592.

Barnard, S., and Simon H. 1994. A fast multi-level implementation of recursive spectral bisection for partitioning unstructured grid. *Concurrency Pract. Exper.* 6, 101–117.

Berman, H.M., Westbrook, J., Feng, Z., et al. 2000. The protein data bank. Nucleic Acids Res. 28, 235-242.

Braig, K., Otwinowski, Z., Hegde, R., et al. 1994. The crystal structure of the bacterial chaperonin GroEL at 2.8 Å. *Nature* 371, 578–586.

Chennubhotla, C., and Jepson, A. 2005. Hierarchical eigensolver for transition matrices in spectral methods. NIPS 17, 273–280.

Chennubhotla, C., and Bahar, I. 2006. Markov propagation of allosteric effects in biomolecular systems. *Mol. Syst. Biol.* 2:36.

Chung, F.R.K. 1997. Spectral Graph Theory. CBMS Lectures, AMS.

Doruker, P., Atilgan, A.R., and Bahar, I. 2000. Dynamics of proteins predicted by molecular dynamics simulations and analytical approaches: application to α-amylase inhibitor. *Proteins* 40, 512–524.

Doruker, P., Jernigan, R.L., and Bahar, I. 2002. Dynamics of large proteins through hierarchical levels of coarse-grained structures, *J. Comput. Chem.* 23, 119.

Fowlkes, C., Belongie, S., Chung, F., et al. 2004. Spectral grouping using the Nyström method. IEEE PAMI 26, 2.

Haliloglu, T., Bahar, I., and Erman, B. 1997. Gaussian dynamics of folded proteins. *Phys. Rev. Lett.* 79, 3090–3093.

Hinsen, K. 1998. Analysis of domain motions by approximate normal mode calculations. Proteins 33, 417.

Hohfeld, J., and Hartl, F.U. 1994. Role of the chaperonin cofactor hsp10 in protein folding and sorting in yeast mitochondria. *J. Cell Biol.* 126, 305–315.

Keskin, O., Bahar, I., Flatow, D., et al. 2002. Molecular mechanisms of chaperonin groel-groes function. *Biochemistry* 41, 491–501.

- Koren, Y., Carmel, L., and Harel, D. 2003. Drawing huge graphs by algebraic multigrid optimization, multiscale modeling and simulation. *SIAM* 1, 645–673.
- Kullback, S. 1959. Information Theory and Statistics. Dover Publications, New York.
- Kullback, S., and Leibler, R.A. 1951. On information and sufficiency. Ann. Math. Statist. 22, 79-86.
- Kundu, S., Melton, J.S., Sorensen, D.C., et al. 2002. Dynamics of proteins in crystals: comparison of experiment with imple models. *Biophys. J.* 83, 723–732.
- Kovalenko, O., Yifrach, O., and Horovitz, A. 1994. Residue lysine-34 in groes modulates allosteric transitions in groel. Biochemistry 33, 14974–14978.
- Kurkcuoglu, O., Jernigan, R.L., and Doruker, P. 2004. Mixed levels of coarse-graining of large proteins using elastic network model methods in extracting the slowest motions. *Polymers* 45, 649–657.
- Landry, S.J., Zeilstra-Ryalls, J., Fayet, O., et al. 1993. Characterization of a functionally important mobile domain of groes. *Nature* 364, 255–258.
- Lehoucq, R.B., Sorensen, D.C., and Yang, C. 1996. ARPACK user guide: solution of large scale eigenvalue problems by omplicitly restarted Arnoldi methods [Technical Report]. Department of CAM, Rice University.
- Li, G.H., and Cui, Q. 2002. A coarse-grained normal mode approach for macromolecules: an efficient implementation and application to Ca²⁺-ATPase. *Biophys. J.* 83, 2457.
- Ma, J., Sigler, P.B., Xu, Z.H., et al. 2000. A dynamic model for the allosteric mechanism of groel. *J. Mol. Biol.* 302, 303–313.
- Ma, J. 2005. Usefulness and limitations of normal mode analysis in modeling dynamics of biomolecular systems. *Structure* 13, 373–380.
- Marques O. 1995. BLZPACK: description and user's guide [Technical Report PA/95/30]. CERFACS, Toulouse, France. McLachlan, G.J., and Basford, K.E. 1988. *Mixture Models: Inference and Applications to Clustering*. Marcel Dekker, New York.
- Miyazawa, S., and Jernigan, R.L. 1985. Estimation of effective inter-residue contact energies from protein crystal structures: quasi-chemical approximation. *Macromolecules* 18, 534.
- Rader, A.J., Chennubhotla, C., Yang, L.-W., et al. 2005. The Gaussian network model: theory and applications. In: Cui, Q. and Bahar, I., eds. Normal Mode Analysis: Theory and Applications to Biological and Chemical Systems. Chapman & Hall/CRC Mathematical and Computational Biology Series, CRC Press, Taylor and Francis Group, 41–64.
- Richardson, A., and Georgopoulos, C. 1999. Genetic analysis of the bacteriophage t4-encoded cochaperonin gp31. *Genetics* 152, 1449–1457.
- Richardson, A., van der Vies, S.M., Keppel, F., et al. 1999. Compensatory changes in groel/gp31 affinity as a mechanism for allele-specific genetic interaction. *J. Biol. Chem.* 274, 52–58.
- Richardson, A., Schwager, F., Landry, S.J., et al. 2001. The importance of a mobile loop in regulating chaperonin/co-chaperonin interaction: humans versus escherichia coli. *J. Biol. Chem.* 276, 4981–4987.
- Shewmaker, F., Maskos, K., Simmerling, C., et al. 2001. The disordered mobile loop of groes folds into a defined beta-hairpin upon binding groel. *J. Biol. Chem.* 276, 31257–31264.
- Saibil, H.R., and Ranson, N.R. 2002. The chaperonin folding machine. Trends Biochem. Sci. 27, 627-632.
- Simon, H., and Zha, H. 2000. Low-rank matrix approximation using the Lanczos bidiagonalization process with applications. SIAM J. Sci. Comput. 21, 2257–2274.
- Tama, F., and Sanejouand, Y.H. 2001. Conformational change of proteins arising from normal mode calculations. *Protein Eng.* 14, 1–6.
- Tama, F., Gadea, F.X., Marques, O., et al. 2000. Building-block approach for determining low-frequency normal modes of macromolecules. *Proteins* 41, 1–7.
- Watkins, D.S. 2002. Fundamentals of Matrix Computations. Wiley-Interscience, New York.
- Xu, Z.H., Horwich, A.L., and Sigler, P.B. 1997. The crystal structure of the asymmetric groel-groes(adp)₇ chaperonin complex. *Nature* 388, 741–750.
- Yifrach, O., and Horovitz, A. 1995. Nested cooperativity in the atpase activity of the oligomeric chaperonin groel. *Biochemistry* 34, 5303–5308.

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