Functional Modes of Proteins Are among the Most Robust

S. Nicolay and Y.-H. Sanejouand

Laboratoire de Physique, Ecole Normale Superieure, 46 allées d’Italie, 69364 Lyon Cedex 07, France

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It is shown that a small subset of modes which are likely to be involved in protein functional motions of large amplitude can be determined by retaining the most robust normal modes obtained using different protein models. This result should prove helpful in the context of several applications proposed recently, like for solving difficult molecular replacement problems or for fitting atomic structures into low-resolution electron density maps. It may also pave the way for the development of methods allowing us to predict such motions accurately.

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For two-domain proteins, it is well known that a few low-frequency normal modes can provide a fair description of their large amplitude motion upon ligand binding [1–3]. Recently, it has been shown that this is also true for proteins with complex architectures [4–8], as long as their functional motion is a collective one, i.e., if it concerns large parts of the structure [9–11]. For instance, a single mode of the T form of hemoglobin is enough to describe accurately its conformational change upon oxygen binding [5].

This result has been successfully applied for exploiting fiber diffraction data [12,13], solving difficult molecular replacement problems [14,15], or fitting atomic structures into low-resolution electron density maps [15–17]. The principle of these applications is to perturb a known structure along low-frequency modes so as to get a deformed structure that is consistent with low-resolution biophysical data, which are obtained after the protein has undergone some large amplitude conformational change. It was also shown that when variations of a few key distances are known, through spectroscopic measurements, for instance, it is possible, using linear response theory, to identify which modes are the most involved in the conformational change [18,19]. However, if such experimental data are missing, it is difficult to guess which low-frequency modes are the functional ones. Hereafter, we show that they are among the most robust ones, i.e., among the most conserved ones when different models are considered. The robustness of the functional modes was recognized when it was shown that they can be obtained [9–11] with simple protein descriptions, like elastic network (EN) models [20–23]. Herein, this property is used so as to identify them.

First, standard normal modes were calculated for a set of five proteins of different sizes and architectures after preliminary energy minimization. The CHARMM program [24] was used, with the EEF1.1 implicit solvent model [25], as done in recent studies performed at this level of detail [26]. Then, for each energy-minimized structure, low-frequency normal modes were calculated with the all-atom EN model proposed by Tirion [21], where the many-parameters empirical energy function $E_p$ used in programs like CHARMM is replaced by:

$$E_p = \sum_{d_{ij} < R_c} C(d_{ij} - d_{ij}^0)^2,$$

where $d_{ij}$ is the distance between atoms $i$ and $j$, $d_{ij}^0$ being their distance in the studied structure. The strength of the potential $C$ is a constant assumed to be the same for all interacting pairs. It is required only in order to define units. As done in previous studies [14], $R_c$, the cutoff parameter, is set to 5 Å.

In order to compare both sets of normal modes, $n_i^{\text{eff}}$, the effective number of EN modes involved in the description of standard mode $i$, is calculated as follows [27]:

$$n_i^{\text{eff}} = \exp[-\sum_n \alpha I_{ij}^2 \ln(\alpha I_{ij}^2)],$$

where $n$ is the number of EN modes taken into account ($n = 100$ herein), $I_{ij}$ being the scalar product between standard mode $i$ and EN mode $j$. The normalization factor $\alpha$ is such that: $\sum \alpha I_{ij}^2 = 1$. Thus, $n_i^{\text{eff}}$ gives the effective number of nonzero $I_{ij}^2$. It ranges from 1 to $n$. As shown in Fig. 1, for each protein considered, a few standard modes, with low ranking, can be described accurately with less than 5–6 EN modes. For a given protein, the set of such modes defines a subspace which is robust, that is, well conserved when modes are calculated with very different models.

Next, such robust modes were sought for, using this time two different EN models. In both cases, as often done [9–11,20,22,23,28], only $C_{\alpha}$ atoms are kept. In the first model, pairs of interacting neighbors are determined according to a distance-cutoff criterion. Setting $R_c$ to a value lower than 8–10 Å splits the elastic network into several independent ones and the number of zero-frequency modes becomes larger than 6. To avoid this artifact, values of 10–15 Å have been used [9,19,28]. For adenylate kinase, with $R_c = 12$ Å, $n_c$, the average number of interacting neighbors per $C_{\alpha}$ atom is 25 ± 7, ranging from 10 to 42, as a function
of the degree of burial of the amino acid in the protein interior.

We designed the second EN model so as to keep $n_c$ as constant as possible from one amino acid to the other. To do so, the following algorithm was used. First, all pairs of $C_a$ atoms are sorted, according to their distance. Then, starting from the atom pair separated by the largest distance, they are removed one after the other, unless one atom of the pair has already $n_c$ neighbors. With this algorithm, setting $n_c = 10$, the average distance between pairs of interacting neighbors is $6.2 \pm 1.8 \text{ \AA}$, close to typical cutoff values used for determining contacts between amino acids in proteins [29]. In the case of adenylate kinase, $n_c$ can be set to a value as low as 7 (see Fig. 2) without splitting the network into independent ones.

As done above, normal modes obtained with both EN models were compared, seeking for robust ones, for a set of 22 proteins considered in previous studies performed with the distance-cutoff criterion [9,10,14]. Like in the case of all-atom models, modes are considered to be robust whenever $n^\text{eff}_i \leq 6$. Statistics of the number of robust modes found for all studied proteins are shown in Fig. 3. In most cases, the number of robust modes is four or less. In only three cases, it is larger than 7. Not surprisingly, they correspond to proteins with large sizes (more than 500 amino acids) and complex architecture, like the DNA polymerase of bacteriophage RB69 (pdb code 1ih7), which is the protein of our data set with the largest number of robust modes (eleven). In four cases, no robust mode is found. Interestingly, the known conformational change of these proteins, namely, tyrosine phosphatase, triose phosphate isomerase, che Y, and HIV-1 protease (pdb codes 1yts, 3tim, 3chy, 1hlp, respectively), is a small amplitude one, with a $C_a$ root-mean-square displacement (rmsd) of 1.5 \text{ \AA} at most.

Then, it was checked that robust modes yield accurate descriptions of protein functional motions. To do so, $Q_d$, the quality of the motion description is calculated as follows [5,10]:

$$Q_d = 100 \sum_{i=1}^{n} I_{id}^2,$$

where $n$ is the number of modes taken into account in the description and $I_{id}$ is the scalar product between mode $i$ and the mode $d$. This quantity ranges from 0 (worst case) to 100 (best case).


FIG. 2. The open form of adenylate kinase (pdb code 4ake). Right: the new $N$-neighbors elastic network model introduced in this study. Most $C_a$ atoms are linked to seven neighbors ($N = 7$). Drawn with Molscript [31].
and the direction of the conformational change observed by crystallographers. Note that \( Q_d = 100\% \) when all modes are included in the description, since they form a complete basis set.

As shown in Fig. 4, the conformational change of lactoferrin upon ligand (iron) binding can be described accurately (\( Q_d \) over 85\%) as a linear combination of the seven lowest-frequency modes of the open form (pdb code 1cb6). Interestingly, all seven modes are found to be robust. In Fig. 5, \( Q_d \) is given as a function of the amplitude of the functional motion of each protein considered when \( n = 100 \) normal modes or when only the robust ones are taken into account in the description. For most proteins with small amplitude motions, i.e., of less than 2–3 Å of rmsd, robust modes fail to capture any information about the nature of the known conformational change, while in several cases some information is indeed present in the normal modes.

On the other hand, when considering proteins with large amplitude motions, the description of the conformational change with robust modes is almost as accurate (\( Q_d \) over 75\%) as when \( n = 100 \) normal modes are taken into account. The only counter example is adenylyl kinase, whose rmsd upon ligand binding is 5.3 Å. As a matter of fact, when standard normal modes of adenylyl kinase are compared to all-atom EN ones, only a single robust mode is found (see Fig. 1), and it is not involved in the conformational change (\( Q_d = 4\% \)). However, using the pair of \( C_\alpha \)-EN models, six robust modes are found and they allow for an almost perfect description of the conformational change (\( Q_d = 91\% \)).

Of course, when using all-atom models, more robust modes can be obtained by raising the robustness criterion. In the case of adenylyl kinase, if a given mode is said robust whenever \( \eta_{eff} \leq 10 \), then five robust modes are found. However, it is still not enough (\( Q_d = 73\% \)) for describing its conformational change as well as with robust modes obtained using \( C_\alpha \)-EN models. Raising the robustness criterion so as to obtain six robust modes does not change significantly the quality of the description (\( Q_d = 77\% \)). As a matter of fact, robust modes obtained using all-atom models always yield poorer description of protein functional motions than when using simpler models, in which only \( C_\alpha \) atoms are kept (open circles are below open squares in Fig. 5). This is likely due to the fact that standard normal mode analysis requires a preliminary energy minimization, during which the structure is significantly distorted, while normal mode analysis of EN models does not, as illustrated by the case of DNA polymerase β. For this protein, when the \( C_\alpha \)-EN models are built using the crystal structure (pdb code 1bpx), seven robust modes are found, which are able to describe accurately (\( Q_d = 84\% \)) the conformational change upon nucleotide binding. However, when they are built using the energy-minimized structure, only three robust modes are found, which are not able to describe the conformational change (\( Q_d = 21\% \)) much better than the three ones obtained using all-atom models.
models ($Q_d = 16\%$). In that case, the distortion due to energy minimization is large (rmsd = 2.5 Å), as a consequence of the removal of the ligand, a 16 base pair DNA, prior to the calculation. However, protein distortions upon energy minimization are usually not that large (rmsd = 1–2 Å) and further work is required in order to fully clarify the origin of this counterintuitive, slight but systematic, effect.

In the present study, modes obtained with different protein models were compared. For most protein cases, several robust modes are found, confirming results obtained previously [9,20–23,30], namely, that the lowest-frequency modes are little sensitive to details in the protein description. Since in the case of current EN models atom–atom interactions are defined with a distance-cutoff criterion, this can be explained in two different ways. First, robust modes may capture information about the protein mass distribution in space. Second, they may capture information about the rigidity of the protein in the vicinity of each amino-acid residue. Indeed, with a distance-cutoff criterion, amino acids in the protein interior are more rigid (more neighbors) than those on the surface (less neighbors). So, we designed a novel $C_\alpha$-EN model whose main raison d’être was to decide between these two possibilities. In this model, each $C_\alpha$ atom has a given number of interacting neighbors and rigidity is fairly constant from one point of a protein to another. When modes obtained with this model are compared to those obtained with a $C_\alpha$-EN model based on the distance-cutoff criterion, robust modes are also found. This means that they are also not sensitive to the distribution of rigidity in the protein.

Moreover, we have shown that these robust modes are likely to be involved in protein functional motions, at least when the functional motion is a large amplitude one (rmsd $\geq$ 2–3 Å). This result should prove helpful in the context of applications like those mentioned in the introduction, since they all concern large amplitude conformational changes [14–17].

This result could also pave the way for the development of methods allowing to predict such motions accurately, i.e., to predict their amplitude, since exploring a subspace of small dimensionality (three or four in most cases considered) should be enough for finding conformations close to functional ones. Interestingly, seeking for robust modes could also indicate whether a given protein can exhibit a large amplitude functional motion or not. Indeed, the functional motions of the proteins found to have no robust mode are small amplitude ones.