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Generalized flexibility-rigidity index

Duc Duy Nguyen,1 Kelin Xia,1 and Guo-Wei Wei1,2,3,a)

1Department of Mathematics, Michigan State University, East Lansing, Michigan 48824, USA
2Department of Biochemistry and Molecular Biology, Michigan State University, East Lansing, Michigan 48824, USA
3Department of Electrical and Computer Engineering, Michigan State University, East Lansing, Michigan 48824, USA

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Flexibility-rigidity index (FRI) has been developed as a robust, accurate, and efficient method for macromolecular thermal fluctuation analysis and B-factor prediction. The performance of FRI depends on its formulations of rigidity index and flexibility index. In this work, we introduce alternative rigidity and flexibility formulations. The structure of the classic Gaussian surface is utilized to construct a new type of rigidity index, which leads to a new class of rigidity densities with the classic Gaussian surface as a special case. Additionally, we introduce a new type of flexibility index based on the domain indicator property of normalized rigidity density. These generalized FRI (gFRI) methods have been extensively validated by the B-factor predictions of 364 proteins. Significantly outperforming the classic Gaussian network model, gFRI is a new generation of methodologies for accurate, robust, and efficient analysis of protein flexibility and fluctuation. Finally, gFRI based molecular surface generation and flexibility visualization are demonstrated. Published by AIP Publishing. [http://dx.doi.org/10.1063/1.4953851]

In living organisms, proteins carry out a vast variety of basic functions, such as structure support, catalyzing chemical reactions, and allosteric regulation, through synergistic interactions or correlations. Protein functions and interactions are determined by protein structure and flexibility.10 The importance of protein structure needs no introduction, while the importance of protein flexibility is often overlooked. Protein flexibility is an intrinsic property of proteins and can be measured by experimental means, including X-ray crystallography, nuclear magnetic resonance (NMR), and single-molecule force spectroscopy, e.g., magnetic tweezer, optical trapping, and atomic force microscopy.9 Flexibility analysis offers a unique channel for theoretical modeling to meet with experimental observations. A variety of theoretical methods, such as normal mode analysis (NMA),6,11,17,19,25 graph theory,15 rotation translation blocks (RTB) method,8,23 and elastic network model (ENM),2,4,14,18,24 including Gaussian network model (GNM)3,4 and anisotropic network model (ANM),2 have been proposed. Among them, GNM is often favored due to its accuracy and efficiency.34 These time-independent methods have been widely used not only for protein fluctuation analysis but also for entropy estimation. However, they typically suffer from two major drawbacks: (1) \(O(N^3)\) scaling in computational complexity with \(N\) being the number of elements in the involved matrix and (2) insufficient accuracy in protein B-factor predictions. The above scaling in computational complexity is due to the matrix diagonalization and makes large biomolecules inaccessible to aforementioned methods. Recently, Park et al. have shown that for three sets of structures of small-sized, medium-sized, and large-sized, the mean correlation coefficients (MCCs) for NMA and GNM B-factor predictions are, respectively, below 0.5 and 0.6.22 These researchers found that both NMA and GNM fail to work for many structures and deliver negative correlation coefficients.22 These problems call for the development of accurate, efficient, and reliable approaches for the flexibility analysis and entropy calculation of macromolecules.

One strategy to tackle the above-mentioned challenges is to develop matrix-diagonalization-free methods for flexibility analysis. To this end, we have introduced molecular nonlinear dynamics,32 stochastic dynamics,31 and flexibility-rigidity index (FRI).20,29 Our approaches make use of protein network connectivity and centrality to describe protein flexibility and rigidity. In our FRI method, we assume that protein interactions, including those with its environment, fully determine its structure in the given environment. In contrast, protein flexibility and rigidity are fully determined by the structure of the protein and its environment. Therefore, to analyze protein flexibility and rigidity, it is unnecessary to resort to the protein interaction Hamiltonian whenever an accurate protein structure is already available. As a result, FRI bypasses the \(O(N^3)\) matrix diagonalization. Our earlier FRI has the computational complexity of \(O(N^2)\) and our fast FRI (fFRI)30 based on a cell lists algorithm4 is of \(O(N)\). Anisotropic FRI (aFRI)30 and multiscale FRI (mFRI)31 have also been proposed. FRI correlation kernels are utilized to develop generalized GNM (gGNM) and generalized ANM (gANM) methods as well as multiscale GNM (mGNM) and multiscale ANM (mANM) methods,30 which significantly improves their accuracy. In the past two years, we have extensively validated FRI, fFRI, aFRI, and mFRI by a set of 364 proteins for accuracy, reliability, and efficiency. Our mFRI is about 20% more accurate than GNM on the 364 protein test set.21 Our fFRI is orders of magnitude faster than GNM on a set of 44 proteins, including one of the largest proteins in the

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a)Author to whom correspondence should be addressed. Electronic mail: wei@math.msu.edu

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Protein Data Bank (PDB), namely, an HIV virus capsid (1E6J) having 313 236 residues. Moreover, to predict the B-factors of the HIV capsid, it will take our fRI only 30 s to complete all calculations, while it would take GNMs more than 120 years to accomplish the same job had the computer memory not been a problem.  

Although various forms of FRI correlation kernels have been introduced, the general mathematical structure of FRI has not been studied. For example, only one rigidity formula and one flexibility formula were proposed. It is interesting to know whether there exists alternative FRI formulations. If so, how do they perform against other existing methods? In the present work, we systematically compare with a number of other methods. The objective of the present work is to shed lights on these issues. Motivated by the success of the Gaussian surface in explicit solvent models, we propose a new class of surface densities equipped with a wide variety of FRI correlation kernels ($\Phi (|| \mathbf{r} - \mathbf{r}_j ||; \eta_j)$).

$$\mu^2(\mathbf{r}) = 1 - \prod_{j=1}^{n} \left[ 1 - w_j \Phi (|| \mathbf{r} - \mathbf{r}_j ||; \eta_j) \right].$$  

(7)

Since both rigidity densities $\mu^2(\mathbf{r})$, $\alpha = 1, 2$ represent molecular density at position $\mathbf{r}$, it is convenient to normalize these densities by their maximal values

$$\bar{\mu}^\alpha(\mathbf{r}) = \frac{\mu^\alpha(\mathbf{r})}{\max_{\mathbf{r} \in \mathcal{R}} \mu^\alpha(\mathbf{r})}, \quad \alpha = 1, 2.$$  

(8)

In this form, the behaviors of two types of rigidity based molecular surfaces can be easily compared. Additionally, normalized rigidity densities in Eq. (8) can be used as solute domain indicators in implicit solvent models. Obviously, $\max \mu^\alpha(\mathbf{r})$ occurs at an atomic position. Therefore, we can define normalized atomic rigidity indexes,

$$\bar{\mu}^\alpha_i = \bar{\mu}^\alpha(\mathbf{r}_i), \quad \alpha = 1, 2.$$  

(9)

With atomic rigidity indexes, $\bar{\mu}^\alpha_i$, we denote the flexibility indexes proposed in our earlier work as atomic flexibility indexes of type I,

$$f^{\alpha1}_i = \frac{1}{\bar{\mu}^\alpha_i}, \quad \alpha = 1, 2.$$  

(10)

The definition of atomic flexibility indexes is not unique. One of the present objectives is to explore other forms of atomic flexibility indexes. Since the normalized atomic rigidity density can be interpreted as a solvent domain indicator, then $1 - \bar{\mu}^\alpha(\mathbf{r})$ can be regarded as a solvent domain indicator. This motivates us to propose a new form of atomic flexibility indexes,

$$f^{\alpha2}_i = 1 - \bar{\mu}^\alpha_i, \quad \alpha = 1, 2.$$  

(11)

We denote $f^{\alpha2}_i$ as atomic flexibility indexes of type II.

In the rest of this paper, we focus on the exploration of gFRI models associated $f^{\alpha\beta}_i$, i.e., $f^{11}_i$, $f^{12}_i$, $f^{21}_i$, and $f^{22}_i$, and denote these models as gFRI$^{\alpha\beta}$. We also study the performance of gFRI for various kernel implementations, namely, generalized exponential and generalized Lorentz correlation kernels.

Due to the proportionality between the atomic flexibility index and the temperature factor at each atom, the theoretical B-factor at $i$th atom, $B_i^{\alpha\beta}$, can be expressed as a linear form,

$$B_i^{\alpha\beta} = a_{\alpha\beta} f^{\alpha\beta}_i + b_{\alpha\beta}, \quad \alpha = 1, 2, \beta = 1, 2.$$  

(12)

where constants $a_{\alpha\beta}$ and $b_{\alpha\beta}$ are independent of index $i$ and can be estimated by the following minimization process:

$$\min_{a_{\alpha\beta}, b_{\alpha\beta}} \left\{ \sum_{i=1}^{N} \left( B_i^{\alpha\beta} - f^{\alpha\beta}_i \right)^2 \right\}.$$  

(13)

where $B_i^\alpha$ is the experimental B-factor for the $i$th atom. To quantitatively assess the performance of the proposed gFRI model, we calculate the correlation coefficient $R$ between the experimental and calculated B-factors.
models for the B-factor prediction, we consider correlation coefficient (CC)

\[
CC = \frac{\sum_{\alpha=1}^{N} (B_{i\alpha} - \hat{B}_{i\alpha}^\alpha)^2}{\sum_{\alpha=1}^{N} (B_{i\alpha} - \hat{B}_{i\alpha}^\alpha)^2 \sum_{\alpha=1}^{N} (B_{i\alpha}^\alpha - \hat{B}_{i\alpha}^\alpha)^2}^{1/2},
\]

where \( \hat{B}_{i\alpha}^\alpha \) and \( \hat{B}_{i\alpha}^\epsilon \) are, respectively, the statistical averages of theoretical and experimental B-factors.

We consider a set of 364 proteins used in our earlier work\(^{30}\) and coarse-grained \( C_\alpha \) atoms in each protein. Therefore, we set \( w_j = 1 \) and use a uniform characteristic distance \( \eta_j = \eta \) in all of our computations.

We first analyze the best parameter set for the B-factor prediction of each rigidity and flexibility type over a range of parameters. Table I reveals the optimal parameters and the best MCCs for gFRI\(^{1} \), with \( \alpha = 1, 2 \) and \( \beta = 1, 2 \). For the sake of visualization, Fig. 1 plots behavior of parameters for exponential and Lorentz kernels in B-factor predictions. It can be seen from Fig. 1 that gFRI\(^{21} \) and gFRI\(^{22} \) models are more sensitive to parameter \( \eta \) than their gFRI\(^{1} \) counterparts. Despite having a fewer choices of fitting parameters, gFRI\(^{21} \) and gFRI\(^{22} \) models are still able to deliver B-factor predictions as accurate as those of gFRI\(^{1} \) models.

To further demonstrate the accuracy of each type of B-factor predictions for different correlations kernels, we plot predicted B-factors against the experimental ones for protein 1DF4 in Fig. 2. In general, all B-factor prediction approaches produce a similar accuracy, especially when a Lorentz kernel is employed. Moreover, the utilization of exponential type of functions for gFRI\(^{12} \) and gFRI\(^{21} \) B-factor prediction types likely performs a little bit worse than the rest.

For an extended comparison, Table I also lists B-factor prediction performances of our earlier FRI\(^{20} \) method, generalized GNM (gGNM),\(^{30} \) generalized ANM (gANM),\(^{30} \) and the classic GNM\(^{34} \) approaches. The earlier FRI algorithm is the same as gFRI\(^{1} \) in the present work while omits the normalization process \((8)\). By employing the same correlation kernel parameters as of gFRI\(^{1} \), the earlier FRI method gives B-factor predictions similar to those of gFRI\(^{1} \). Specifically, MCCs produced by the previous FRI algorithm for exponential kernel and Lorentz kernel, are, respectively, 0.623 and 0.626. It is noted that the earlier FRI predicted B-factors over 365 proteins,\(^{20} \) while current methods employ the same data set with one left out, 1AGN, due to the unrealistic experimental B-factors.

The gGNM method\(^{30} \) is an FRI kernel generalization of GNM.\(^{34} \) In this approach, the \( i \)th B-factor of a biomolecule can be defined as

\[
B_{i}^{\text{gGNM}} = a_{\text{gGNM}} (\Gamma^{-1})_{ij}, \quad \forall i = 1, 2, \ldots, N,
\]

where \( a_{\text{gGNM}} \) is a fitting parameter and \( (\Gamma^{-1})_{ij} \) is the \( ij \)th diagonal element of the matrix inverse of the generalized Kirchhoff matrix\(^{30} \)

\[
\Gamma_{ij} (\Phi) = \begin{cases} 
-\Phi (||r_i - r_j||; \eta_j), & i \neq j, \\
\sum_{j, i = 1}^{N} \Gamma_{ij} (\Phi), & i = j.
\end{cases}
\]

Similarly, the gANM method\(^{30} \) is an FRI kernel generalization of the classic ANM method.\(^{2} \) In this approach, the generalized local \( 3 \times 3 \) Hessian matrix \( H_{ij} \) is written as

\[
H_{ij} = -\frac{\Phi (||r_i - r_j||; \eta_j)}{||r_i - r_j||^2} \left[ \begin{array}{ccc} (x_j - x_i)(x_j - x_i) & (x_j - x_i)(y_j - y_i) & (x_j - x_i)(z_j - z_i) \\
(y_j - y_i)(x_j - x_i) & (y_j - y_i)(y_j - y_i) & (y_j - y_i)(z_j - z_i) \\
(z_j - z_i)(x_j - x_i) & (z_j - z_i)(y_j - y_i) & (z_j - z_i)(z_j - z_i) \end{array} \right], \quad \forall i \neq j.
\]

We define the diagonal parts as \( H_{ii} = -\sum_{j \neq i} H_{ij}, \forall i = 1, 2, \ldots, N \). Therefore, the B-factor of \( i \)th \( C_\alpha \) in a biomolecule is expressed as

\[
B_{i}^{\text{gANM}} = a_{\text{gANM}} \text{Tr}(H^{-1})_{ii}, \quad \forall i = 1, 2, \ldots, N,
\]

where \( a_{\text{gANM}} \) is a fitting parameter and \( \text{Tr}(H^{-1})_{ii} \) is the summation of the \( i \)th diagonal element of the matrix inversion of a matrix formed by generalized local Hessian matrices.
accuracy. Since the gANM prediction with Lorentz type of functions is nowhere close to the acceptable level in terms of the accuracy, the result for that case was not reported.\textsuperscript{30} Finally, with a cutoff distance of 7 Å, the prediction of classic GNM method delivers an MCC of 0.565, which is better than that obtained by gANM, but not as good as any other methods in comparison. Since GNM is one of the most popular and the most accurate methods for B-factor predictions,\textsuperscript{34} the current comparison shown in Table I indicates that FRI and gFRI are a new generation of more accurate and robust methods for protein B-factor prediction.

To further compare the B-factor performances between different methods, we introduce two types of continuous flexibility functions,
FIG. 2. Experimental B-factors (gray) vs predicted B-factors (red) of 1DF4 using the exponential (left) and Lorentz (right) correlation kernels. The optimal parameters for each type of B-factor prediction are described in Table I. (a) gFRI\textsubscript{11}. (b) gFRI\textsubscript{12}. (c) gFRI\textsubscript{21}. (d) gFRI\textsubscript{22}.

\[ F_{\alpha 1}(r) = \frac{a_{\alpha 1}}{\bar{\mu}^1(r)} + b_{\alpha 1} \]  \hspace{1cm} (19)

or

\[ F_{\alpha 2}(r) = a_{\alpha 2} (1 - \bar{\mu}^1(r)) + b_{\alpha 2}. \]  \hspace{1cm} (20)

where \( a_{\alpha \beta} \) and \( b_{\alpha \beta} \) are determined by minimization (13). These flexibility functions are volumetric and can be projected onto a molecular surface for flexibility visualization. As mentioned earlier, rigidity densities provide excellent molecular surface representations and one can employ either rigidity density \( \bar{\mu}^1(r) \) or \( \bar{\mu}^2(r) \) for surface generation. However, for the purpose of surface visualization, rigidity density \( \bar{\mu}^2(r) \) is not suitable. The reason is that the rigidity formula (7) in the continuous form has to avoid all the grid points near each atomic center. Otherwise rigidity densities would be mostly 1 and the corresponding flexibility functions would be mostly a constant on the molecular surface. This hindrance can be remedied by using interpolation approach as discussed in our previous work.\textsuperscript{29} In the present work, we only consider \( \bar{\mu}^1(r) \) for surface representation and employ \( F_{11}^1(r) \) and \( F_{12}^2(r) \) for flexibility visualization. The parameters for \( F_{11}^1(r) \) and \( F_{12}^2(r) \) follow those of gFRI\textsubscript{11} and gFRI\textsubscript{12} as shown in Eqs. (19) and (20), respectively.

Figure 3 depicts the projection of flexibility functions \( F_{11}^1(r) \) onto the isosurface of rigidity density \( \bar{\mu}^1(r) = 0.05 \) of protein 1DF4. Parameters \( k = 1, w_j = 1, \) and \( \eta_j = 0.5 \) Å for all \( j = 1, \ldots, N \) are used in \( \bar{\mu}^1(r) \) for surface generation. Even though both gFRI\textsubscript{11} and gFRI\textsubscript{12} deliver similar B-
factor predictions, with MCCs being 0.888 for gFRI\textsuperscript{11} and is 0.889 for gFRI\textsuperscript{12}, their flexibility functions ($F_i^\beta(r)$) behave differently. It can be seen from Fig. 3 that outer region of $F_i^{11}(r)$ projection contains higher values than its counterpart $F_i^{12}(r)$, while the inner region of both $F_i^{11}(r)$ stays almost the same for both methods. This behavior is likely due to the fact that $F_i^{11}(r)$ is constructed by Eq. (19), which dramatically amplifies small rigidity densities far away from the center of mass of a molecule. In contrast, $F_i^{12}(r)$ is bound and well defined everywhere.

To predict the amplitudes and directions of atomic fluctuation, ANM\textsuperscript{2} is commonly used. Another tool for such a purpose is the anisotropic FRI (aFRI) proposed in our previous work.\textsuperscript{20} It is interesting to know whether the present flexibility formulation (11) leads to a new algorithm for protein anisotropic motion analysis. To this end, we present a brief review of aFRI theory, which establishes notions for new formulation. In ANM, the Hessian matrix is a global matrix containing $3N\times 3N$ elements with $N$ being a number of atoms. In our aFRI model, depending on one’s interest, the size of the Hessian matrix can vary from $3\times 3$ for a completely local aFRI to $3N\times 3N$ for a completely global aFRI. To construct such a Hessian matrix, we partition all $N$ atoms in a molecule into a total of $M$ clusters $\{c_1, c_2, \ldots, c_M\}$. Each cluster $c_k$ with $k = 1, \ldots, M$ has $N_k$ atoms so that $N = \sum_{k=1}^{M} N_k$. For convenience, we denote

$$
\Phi_{ij}^{uv} = \frac{\partial \Phi_{ij}}{\partial u_i \partial v_j}, \\
u, v = x, y, z; i, j = 1, 2, \ldots, N.
$$

Note that for each given $ij$, we define $\Phi_{ij} = (\Phi_{ij}^{uv})$ as a local anisotropic matrix

$$
\Phi_{ij} = \begin{pmatrix}
\Phi_{ij}^{xx} & \Phi_{ij}^{xy} & \Phi_{ij}^{xz} \\
\Phi_{ij}^{yx} & \Phi_{ij}^{yy} & \Phi_{ij}^{yz} \\
\Phi_{ij}^{zx} & \Phi_{ij}^{zy} & \Phi_{ij}^{zz}
\end{pmatrix}.
$$

In the anisotropic flexibility approach, a flexibility Hessian matrix $\Phi_i^a(c_k)$ for cluster $c_k$ is defined by

$$
\Phi_i^a(c_k) = \frac{1}{u_i} (\Phi_{ij})^{-1}, \quad i, j \in c_k; i \neq j; u, v = x, y, z,
$$

$$
\Phi_i^a(c_k) = \sum_{j=1}^{N} \frac{1}{w_j} (\Phi_{ij})^{-1}, \quad i \in c_k; u, v = x, y, z,
$$

$$
\Phi_i^a(c_k) = 0, \quad i, j \notin c_k; u, v = x, y, z,
$$

where $(\Phi_{ij})^{-1}$ denotes the unscaled inverse of matrix $\Phi_{ij}$ such that $\Phi_{ij}(\Phi_{ij})^{-1} = (\Phi_{ij})$.

Motivated by the new form of atomic flexibility indexes (11), we propose another presentation for the flexibility Hessian matrix $\Phi_i^a(c_k)$ as follows:

$$
\Phi_i^a(c_k) = \frac{1}{w_j} (\Phi_{ij})(J_3 - \Phi_{ij}), \quad i, j \in c_k; i \neq j; u, v = x, y, z,
$$

$$
\Phi_i^a(c_k) = \sum_{j=1}^{N} \frac{1}{w_j} (\Phi_{ij})(J_3 - \Phi_{ij}), \quad i \in c_k; u, v = x, y, z,
$$

$$
\Phi_i^a(c_k) = 0, \quad i, j \notin c_k; u, v = x, y, z,
$$

where $J_3$ is a $3 \times 3$ matrix with every element being one.

We can achieve $3N_k$ eigenvectors for $N_k$ atoms in cluster $c_k$ by diagonalizing $\Phi_{ij}^a(c_k)$, $\alpha = 1, 2$. Note that the diagonal part $\Phi_{ii}^a(c_k)$, $\alpha = 1, 2$, has inherent information of all atoms in the system. As a result, we can predict B-factors by employing Eq. (13) for a set of flexibility indexes collected from the diagonal parts

$$
f_i^{AF\alpha} = \text{Tr}(\Phi_{ii}^a(c_k))^\alpha \\
= (\Phi_{ii}^a(c_k))_{xx}^{ii} + (\Phi_{ii}^a(c_k))_{yy}^{ii} + (\Phi_{ii}^a(c_k))_{zz}^{ii},
$$

$$
\alpha = 1, 2.
$$

In this work, we compare the protein anisotropic motion predictions by using completely global aFRI models based on the anisotropic flexibility associated with $f_i^{AF\alpha}$ and denote these models as aFRI\textsuperscript{a}, $\alpha = 1, 2$. Note that model aFRI\textsuperscript{a} is already discussed in our previous work\textsuperscript{20} and used for a comparison. Figure 4 depicts the first three nontrivial isotropic modes of aFRI\textsuperscript{1}, aFRI\textsuperscript{2}, and ANM for protein PDBID: 2XHF. The Lorentz kernel is used for both aFRI algorithms with
FIG. 4. Comparison of modes for protein PDB ID: 2XHF. The top row is generated by using the completely global aFRI\textsuperscript{1} with $\nu = 2$ and $\eta = 30$ Å. The middle row is generated by using the completely global aFRI\textsuperscript{2} with $\nu = 2$ and $\eta = 25$ Å. The bottom row is generated by using ANM with Prody v1.8\textsuperscript{5} using default settings. (a) aFRI\textsuperscript{1}, (b) aFRI\textsuperscript{2}, (c) ANM.

$w_j = 1, \nu = 2$, and $\eta_j = 30$ Å, $\forall j = 1, \ldots, N$ for aFRI\textsuperscript{1}, and $w_j = 1, \nu = 2$, and $\eta_j = 25$ Å, $\forall j = 1, \ldots, N$ for aFRI\textsuperscript{2}. To obtain ANM prediction, we use Prody v1.8\textsuperscript{5} with default settings. It is interesting to see that each algorithm has its own set of collective protein motions. Since there is no exact answer to these fluctuation modes, we cannot conclude which motion prediction is right or wrong.

It needs to point out that the proposed gFRI can be readily incorporated into our fFRI and mFRI methodologies. Additionally, infinitely many possible atomic flexibility
indexes of a general functional form \( f(\vec{\mu}_i) \) can be designed. For example, one can choose \( f(\vec{\mu}_i) = \Phi(\vec{\mu}_i; \eta_0) \) with \( \eta_0 \) being a constant. However, it is not obvious how to design another distinct rigidity density formula. A systematic analysis of these aspects is beyond the scope of the present work.

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