On the use of low-frequency normal modes to enforce collective movements in refining macromolecular structural models

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As more and more structures of macromolecular complexes get solved in different conditions, it has become apparent that flexibility is an inherent part of their biological function. Normal mode analysis using simplified models of proteins such as the elastic network model has proved very effective in showing that many of the structural transitions derived from a survey of the Protein Data Bank can be explained by just a few of the lowest-frequency normal modes. In this work, normal modes are used to carry out medium- or low-resolution structural refinement, enforcing collective and large-amplitude movements that are beyond the reach of existing methods. Refinement is carried out in reciprocal space with respect to the normal mode amplitudes, by using standard conjugate-gradient minimization. Several tests on synthetic diffraction data whose mode concentration follows the one of real movements observed in the Protein Data Bank have shown that the radius of convergence is larger than the one of rigid-body refinement. Tests with experimental diffraction data for the same protein in different environments also led to refined structural models showing drastic reduction of the rms deviation with the target model. Because the structural transition is described by very few parameters, over-fitting of real experimental data is easily detected by using a cross-validation test. The method has also been applied to the refinement of atomic models into molecular envelopes and could readily be used to fit large macromolecular complex rearrangements into cryo-electron microscopy-reconstructed images as well as small-angle x-ray scattering-derived envelopes.

Understanding protein flexibility is a major challenge in structure–function relationship studies that structural molecular biology addresses routinely through both computational and experimental methods. Although normal modes analysis (NMA) has been known for years to be applicable to proteins to describe their movements (1–3), the method has recently enjoyed renewed interest because of the emergence of simplified models that proved easy to use and very effective in capturing the dynamics of proteins (4) but also because more and more structures of the same protein have been solved in different conformational states, providing more test cases. In many cases, it was observed that low-frequency normal modes are very good at explaining collective large-amplitude motions of proteins known in different conformational states (5). This result was shown for instance in a number of proteins undergoing induced fit upon ligand binding, particularly proteins existing in both a closed and an open form (6, 7). Other studies involve hemoglobin (8), but also large macromolecular complexes such as the ribosome (9) and even an entire viral particle (10).

In an extensive survey using a database of protein movements observed in protein structures deposited in the Protein Data Bank (PDB), Gerstein and colleagues (5) showed that, in most cases, a handful of the lowest-frequency modes deduced from NMA and a simplified representation of proteins is sufficient to explain the observed movements. This finding is based on the so-called elastic network model of proteins first described by Tirion (4) and then further developed independently by Hinsen (11) and Bahar and colleagues (12, 13). Normal modes calculated with this method are able to describe faithfully the B-factors (12, 14) in protein crystals. The obvious advantage of the elastic model is that NMA is very rapid and also that, thanks to the rotation–translation block (RTB) method recently developed by Sanejouand and colleagues (15), there is virtually no limit in the size of the macromolecular assembly one can study.

However, in most of these studies, NMA is essentially a “postmortem” analysis, in the sense that both structures have to be known before a tentative molecular explanation of the biologically relevant movement(s) can be offered. For the time being, the prediction of such movements from just one of the two structures remains a risky exercise.

In this article, we propose to use NMA to refine a starting structural model against experimental data such as medium- or low-resolution x-ray diffraction data, electron microscopy-reconstructed images of large macromolecular complexes or molecular envelopes derived from small-angle x-ray scattering data. We show that such a refinement procedure, using a very small number of parameters (the normal mode amplitudes of 10–20 lowest-frequency modes), has a very large radius of convergence. It is fast and robust, and its effectiveness can easily be monitored by the combination of the two reciprocal-space scores, Rwork and Rfree, that the community of crystallographers are routinely using during refinement of atomic models (16).

This method has numerous applications. The first application is in classical x-ray crystallography where it should supersede rigid body refinement, being just a generalization of the latter method. Indeed, it allows for global translations and rotations, but also for the movement of domains, without the obligation for the user to define subjective and somewhat arbitrary boundary regions of the different movable domains. In this respect, it is obvious that normal mode (NM) refinement could be used with benefit to get a better starting model after running a molecular replacement (MR) program, such as AMORE for instance (17), or even to identify the correct solution. Because the success of MR relies heavily on the ability to cope with large conformational changes in the original model, we expect NM refinement to be more useful than just rigid-body refinement, especially in the context of structural genomics.

A second application is to deform, in a plausible manner, an existing model into a known molecular envelope obtained either through electron microscopy (EM) image reconstruction techniques or small-angle x-ray scattering (SAXS) data. Indeed, EM and SAXS often offer the possibility of getting a low resolution image of the same macromolecular object in different conforma-
tional states, for instance after binding one or several ligands. In this case, NM refinement allows one to quickly gain detailed structural insight into the deformations induced upon ligand binding.

Due to the low number of parameters being refined, the method has a very favorable (no. of observations/no. of parameters) ratio that makes it very attractive when only a set of low-resolution experimental constraints is available.

Crystallographic refinement using NMA has been attempted before, especially in the pioneer papers of Kidera and Go (18) and Diamond (19). However, both works involved only the modeling of atomic (anharmonic) disorder through a better estimation of the Debye–Waller B-factors by using normal modes and made no attempt to refine the amplitudes of the normal modes to carry out positional refinement.

The application of deformations along normal mode eigenvectors has been used before by Tirion et al. to fit fiber diffraction data on F-actin (20), but it seems that this work has remained one parameter in this simple model, namely the cutoff residues in the protein, other strategies should be considered, mal modes of a protein of PAQ workstation for calculating the 100 lowest-frequency normal modes. Typically, it takes 2 min of cpu on an XP1000 COM-PAO workstation for calculating the 100 lowest-frequency normal modes, read precaution proved unnecessary in all of the cases described here. The program used to calculate the normal modes, which are the distance between two atoms a and b whereas r_{ab}^0 is their distance in the reference state. C is a constant as in the original Tirion model (4). The sum is restricted through the Heaviside function H to “interacting” atoms, i.e., if r_{ab}^0 \leq R_c. By construction, the reference conformation corresponds to the global minimum of the energy E. It would be easy to modify the model so as to incorporate two elastic constants instead of just one, one for C-alpha atoms adjacent along the chain (whose distance should always be 3.8 Å) and the other (softer) for C-alpha atoms close in space. However, this precaution proved unnecessary in all of the cases described here.

The program used to calculate the normal modes, which are obtained by diagonalizing the matrix of the second derivatives of E (the so-called Hessian matrix), was written by Y.-H. Sanejouand. Typically, it takes 2–3 min of cpu on an XP1000 COM-PAO workstation for calculating the 100 lowest-frequency normal modes of a protein of \approx 800 residues. If there are \approx 1,000 residues in the protein, other strategies should be considered, such as the use of superresidues (15).

For each normal mode \ell, a set of eigenvectors \mathbf{u}_{\ell}^{(i)} is obtained for each eigenvalue; they represent the displacement at every residue i in the protein for this mode \ell. To describe a large-amplitude conformational change of a given protein, we need only the associated amplitudes c_{\ell}, which are the parameters to be adjusted against experimental data. The first six modes (associated with the same degenerate null eigenvalue) correspond to global translations and rotations. Refining against only these six modes will amount to a simple rigid body minimization.

Crystallographic Refinement. In reciprocal space, the structure factors \mathbf{F}_{\text{calc}}(\mathbf{H}) of the modified model with amplitudes \alpha_{\ell} along the first Nmod (typically 20) lowest-frequency normal modes, read

\[ \mathbf{F}_{\text{calc}}(\mathbf{H}) = \sum_{i=1}^{\text{Natm}} \exp[2i\pi \mathbf{H}(\mathbf{r}_i^0 + \sum_{\ell=1}^{\text{Nmod}} \alpha_{\ell} \mathbf{u}_{\ell}^{(i)})] \]  

To optimize the normal mode amplitudes against experimental data, the crystallographic residual R-factor is minimized by using a conjugate-gradient algorithm (21):

\[ R = \sum_{i} \frac{|\mathbf{F}_{\text{calc}}(\mathbf{H})| - k|\mathbf{F}_{\text{calc}}(\mathbf{H})|}{\sum_{i} |\mathbf{F}_{\text{calc}}(\mathbf{H})|^2} \]

where k is a scaling factor that may also incorporate a global B-factor.

Here, we used the routine ZCGR, which is also the minimizer used in CNS (22); the derivatives \partial R/\partial \alpha_{\ell} needed by the minimizer were calculated analytically and checked numerically. Alternatively, one can minimize the score 1 – C_C(\mathbf{F}_{\text{calc}}, \mathbf{F}_{\text{calc}})

\[ \sum_{i} |\mathbf{F}_{\text{calc}}(\mathbf{H})| - k |\mathbf{F}_{\text{calc}}(\mathbf{H})| \]  

This score also reduces to the R-factor when the amplitudes are zero.

Synthetic Test Structure Factors. For the purpose of testing the program, a number of structures deformed along the normal modes directions were generated by randomly choosing the amplitudes of typically the 10–20 lowest-frequency normal modes, and the corresponding structure factors were generated in the appropriate space group.

We found it useful to monitor both the “mode concentration,” called \alpha, as defined by Gerstein and colleagues (4): \alpha = \sum_{\ell=1}^{\text{Nmod}} |c_{\ell}|^2 / \sum_{\ell=1}^{\text{Nmod}} |c_{\ell}|^2 are the normalized absolute values of the amplitudes.

At the end of the refinement, in test cases where the answer is known, the residual of the mode amplitudes R-mode = \sum_{\ell=1}^{\text{Nmod}} |c_{\ell}|^2 / \sum_{\ell=1}^{\text{Nmod}} |c_{\ell}|^2 was computed.

PDB Test Structure Factors. Two sets of structures were chosen for validating the refinement procedure, namely citrate synthase (PDB ID codes SC3C and 6SCS) and the maltodextrin binding protein (PDB ID codes 1OMP and 1ANF) in both their open and closed forms. Experimental observed structure factors are available from the PDB for the closed forms. Their space groups are respectively P2_12_12_1 and P1. Citrate synthase is a dimer of \approx 850 residues whereas the maltodextrin binding protein contains \approx 370 residues. The overlap between the displacement vectors of the lowest-frequency normal mode and the difference vectors between the two forms is 0.83 and 0.86, respectively (6). The transition between the two forms is best described as a shear movement and a hinge-bending movement, respectively (5).

For MR tests, the open and closed forms of human polymerase \beta (1BPX and 1BPY) were used (335 residues). Structure factors of 1BPY were retrieved from the PDB (space group P2_1). The transition between the two forms is well described by NMA (7).

Envelope. For the envelope test, the envelope map was calculated from the coordinates of the full atomic model. For every point
on a grid of size 100 Å × 100 Å × 100 Å and step size 5 Å the density was set to 1 if its distance to any protein atom was <1.8 Å. The density of all other points was set to 0. Structure factors of the envelope were obtained up to a 12 Å resolution by using the CCP4 package (23).

For the minimization, the phased correlation coefficient was used instead of the correlation coefficient on structure factors modulus only: in this case, Eq. 4 was replaced by

\[ CC(F_{\text{obs}}, F_{\text{calc}}) = \frac{\sum_{i} F_{\text{obs}}(H) F_{\text{calc}}^*(H)}{\left( \sum_{i} |F_{\text{obs}}(H)|^2 \sum_{i} |F_{\text{calc}}(H)|^2 \right)^{1/2}} \]

where \( F_{\text{calc}}(H)^* \) is the complex conjugate of \( F_{\text{calc}}(H) \).

Although it is possible to refine C-alpha models directly, we found slightly better results in using the set of points on the grid inside the envelope as the protein elastic network. This procedure ensures that \( CC \) is exactly 1.00 for the correct envelope, allowing maximum contrast between a good and bad fit, and does not prevent an accurate calculation of the normal modes, as recently shown by Doruker and Jernigan (24).

Most fitting methods in reconstructed envelopes of macromolecules are based on correlation of model and experimental electron densities in real space (refs. 25 and 26, but see ref. 27 for a reciprocal space approach). Here, the refinement is carried out in reciprocal space with a model that has very few parameters.

**Recovering the Final Model.** After completion of the refinement process, the weighted sum of the normal mode displacement vectors, \( \Delta r_i = \sum_{i=0}^{l} \text{Nmode}(i)^{l/2} \), was calculated and applied directly to the \( i \)th C-alpha coordinates of the initial model. For non-C-alpha atoms, interpolation was needed. We used a spherical averaging procedure with a weighting factor equal to the inverse of the mutual distance between the point of interest and points where the displacement vectors are known, i.e. the C- alphas, and located <10 Å away.

The resulting model was subjected to 250 cycles of Newton-Raphson energy minimization in CNS (22), while harmonically restraining the positions of the C- alpha s with a constant of 100 kcal/mol, so as to restore the stereochemistry of the protein chain. Superpositions of models and rms deviations (rmsd) were calculated with the Mclachlan algorithm (28), as implemented in the program PROFIT by A. C. R. Martin (www.bioinf.org.uk/software).

In the case of model refinement inside an envelope represented by a set of lattice points, the interpolation method of finding the displacement vectors for all of the atoms of the model was simply a tri-linear interpolation scheme (21).

**Results**

**Refinement Against Synthetic Diffraction Data.** The citrate synthase (ID code 5C5C) was considered first, and structure factors were calculated for deformed models of the protein obtained by varying amplitudes of the 10 lowest-frequency normal modes. These amplitudes were chosen randomly (positive or negative), and the calculated structure factors became the target structure factors against which the undistorted model was refined. The score being minimized was the correlation coefficient on the structure factors moduli (see Eq. 4). The absolute scale of the amplitudes was increased from one experiment to the other so as to restore the stereochemistry of the protein chain. Superpositions of models and rms deviations (rmsd) were calculated with the Mclachlan algorithm (28), as implemented in the program PROFIT by A. C. R. Martin (www.bioinf.org.uk/software).

In the case of model refinement inside an envelope represented by a set of lattice points, the interpolation method of finding the displacement vectors for all of the atoms of the model was simply a tri-linear interpolation scheme (21).

**Refinement Against Real Diffraction Data.** The open form of maltodextrin binding protein (ID code 1ANF) was subjected to NM refinement against the experimental structure factors of its closed form.
form (ID code 1OMP). Only C-alpha coordinates were selected, and data up to 4 Å resolution were used (~5,500 reflections). Ten percent of the data were omitted from the refinement to monitor the R-free. The rmsd between 1ANF and 1OMP C-alpha coordinates is 3.8 Å. Only the five lowest-frequency modes were included. The results are presented in Table 1.

After conjugate-gradient minimization, the correlation coefficient on structure factors moduli increased from 8.9% (9.7%) to 30.8% (28.1%) where the values in parenthesis correspond to CC-free. The rmsd between the refined model and the target model was 1.14 Å, a considerable improvement from the initial value (3.8 Å). Displacement vectors corresponding to the five modes under consideration, weighted by their respective refined amplitudes, were applied to all atoms according to the procedure described above. The rmsd of the final full atomic model with the target model was 1.48 Å, after refinement with CNS to regularize the stereochemistry. The cpu time of the whole process was <3–5 min even though no special effort was made to speed up the computation. The same experiment was repeated with the 15 lowest-frequency modes instead of only the lowest 5 ones, and the refinement converged to the same solution with a final correlation coefficient of 33% (28%).

As a control experiment, the correct model 1OMP was subjected to the same refinement protocol. The initial correlation coefficient was 47% (49%) and was not much modified after completion of the refinement, reaching 48% (49%). The amplitudes of the normal modes were very small, and the final model was only 0.18 Å away from the initial one, meaning that the correct model was not affected by the refinement, as expected (see Table 1).

The same NM refinement was done on citrate synthase (5CSC and 6CSC). The rmsd between the open and closed forms is 3.0 Å. Refinement of the control model (6CSC) against its structure factor moduli gave a refined model only 0.05 Å away from the initial one and a stable correlation coefficient of 45%. Starting from the 5CSC, however, and refining against the 6CSC experimental structure factors, the correlation coefficient increased from 19% (15%) to 40% (39%) in <20 cycles (see Table 1). In this process, the third lowest-frequency normal mode received the largest amplitude, as it should (Fig. 3).

In comparison, standard protocols of simulated annealing refinement in dihedral angle space (22) failed to reduce the $R_{free}$ factor, in the same conditions, for both test cases: $R_{free}$ remained stuck at 47.5% and 52.5% for 5CSC and 1ANF, respectively. The same was true for standard rigid body refinement using CNS.

Because the model moves a lot during the refinement, it might seem advisable to update the normal mode calculation during the refinement. To test this idea, we stopped the refinement every time the model moved by >1.5 Å away from its original position and recalculated the normal modes before reentering the minimizer. The program converged to the same solution for the maltose binding protein and 15 normal modes (data not shown). The reason for this phenomenon is that the first 15 normal modes of the target model form a very good basis set for the refinement. To test this idea, we stopped the refinement every time the model moved by >1.5 Å away from its original position and recalculated the normal modes before reentering the minimizer. The program converged to the same solution for the maltose binding protein and 15 normal modes (data not shown). The reason for this phenomenon is that the first 15 normal modes of the target model form a very good basis set for the 15 normal modes of the initial model, and vice versa.

**Table 1. Normal mode refinement against experimental diffraction data**

<table>
<thead>
<tr>
<th>Starting model</th>
<th>Target model</th>
<th>Resolution (Rnl)</th>
<th>Initial rmsd, Å</th>
<th>Final rmsd, Å</th>
<th>Initial CC (Free-CC)</th>
<th>Final CC (Free-CC)</th>
<th>Nmod</th>
<th>Natom (space group)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1ANF</td>
<td>1OMP</td>
<td>10 to 4 Å (5,500)</td>
<td>3.8</td>
<td>1.1</td>
<td>8.9 (9.7)</td>
<td>30.8 (28.1)</td>
<td>5</td>
<td>370 (1)</td>
</tr>
<tr>
<td>1OMP</td>
<td>1OMP</td>
<td>10 to 4 Å (5,500)</td>
<td>0.0</td>
<td>0.2</td>
<td>47.6 (49.6)</td>
<td>48.2 (49.4)</td>
<td>5</td>
<td>370 (1)</td>
</tr>
<tr>
<td>1ANF</td>
<td>1OMP</td>
<td>10 to 4 Å (5,500)</td>
<td>3.8</td>
<td>1.1</td>
<td>8.9 (9.7)</td>
<td>33.2 (28.5)</td>
<td>15</td>
<td>370 (1)</td>
</tr>
<tr>
<td>5CSC</td>
<td>5CSC</td>
<td>10 to 4 Å (7,860)</td>
<td>3.0</td>
<td>1.6</td>
<td>19.2 (15.2)</td>
<td>40.0 (39.7)</td>
<td>5</td>
<td>855 (19)</td>
</tr>
<tr>
<td>6CSC</td>
<td>6CSC</td>
<td>10 to 4 Å (7,860)</td>
<td>0.0</td>
<td>0.05</td>
<td>49.6 (48.2)</td>
<td>49.8 (48.5)</td>
<td>5</td>
<td>855 (19)</td>
</tr>
</tbody>
</table>

**Fig. 2.** Refinement with respect to the normal modes associated with null frequency: rigid-body refinement. The rmsd with the target model of the refined model is plotted as a function of the rmsd of the starting model. The refinement was carried out against calculated structure factors at a resolution of 8 Å. The minimized quantity is the score based on the correlation coefficient of structure factor moduli defined in Eq. 4, as in Fig. 1.

**Fig. 3.** Projection of the set of displacement vectors, for each low-frequency normal mode from 1 to 106, onto the set of difference vectors between the two forms the maltodextrin binding protein (PDB ID codes 1OMP and 1ANF). In green we show the refined amplitudes, using conjugate-gradient minimization in reciprocal space against real data.
were refined together with rigid body degrees of freedom. The correct solution, which is ranked second after rigid body refinement, is solution no. 5 in Fig. 4. NM refinement clearly identifies the correct solution by yielding a significantly higher correlation coefficient than the false positive solution no. 1. Clearly, this finding seems of real interest to enhance the range of application of molecular replacement in difficult cases.

Refinement Against Molecular Envelopes. Finally, an envelope was constructed for the closed form of citrate synthase (6CSC), and its structure factors were calculated in P1. The open form envelope was also calculated, and a set of grid points inside the envelope was used to compute normal modes, whose amplitudes were refined against the phased correlation coefficient of the closed form (see Eq. 5). The refinement was done at 12 Å resolution with \( \approx 1,200 \) reflections. A definite improvement of the fit into the envelope of the closed form was observed with only five modes whereby the correlation coefficient increased from 87% to 93% in just a few seconds of cpu time (Fig. 5). The weighted displacement vectors of the grid points were then used to move the lattice points inside the envelope as well as the open form C-alpha coordinates, by using standard linear interpolation techniques (21). The rmsd of the resulting model with the target model was reduced from 3.0 to 1.8 Å. Adding more modes did not improve significantly the final correlation coefficient. The same refinement was attempted with maltodextrin binding protein (1OMP and 1ANF), resulting in the same kind of rmsd reduction, from 3.8 to 1.8 Å.

Discussion

Normal Modes and the Elastic Model. Impressive results were found for the refinement of structural models against medium- or low-resolution experimental data in reciprocal space, by using as sole parameters the amplitudes of a small set of low-frequency normal modes derived from a simple one-parameter elastic model of the protein. It is clear that normal mode refinement has a larger radius of convergence than classical rigid-body refinement (compare Figs. 1 and 2).

This finding might seem counterintuitive at first sight. In fact, it is just a consequence of the model described by Eq. 1 where low-frequency normal modes do enforce collective large-amplitude movements. Because the model contains very few parameters and is exactly accounted for in Eqs. 1 and 2, it is perhaps not so surprising that a relatively simple but robust conjugate-gradient minimizer can find the correct solution in very little cpu time.

The real question, therefore, is how well the model encoded in Eq. 1 describes biologically relevant large-amplitude movements of macromolecules. Loop movements and refolding events are likely to be poorly described by low-frequency normal modes.

Fortunately, it is possible here to invoke the systematic study of Gerstein and colleagues (5), who showed that a large proportion of protein movements documented in the PDB are indeed well described for the same protein in different environments (crystal packing, substrates, etc.), by just a few normal modes calculated by using the model represented by Eq. 1. This finding is complemented by a number of studies on specific systems (7–10). Also, a fortunate feature of the analysis in ref. 5 is that models with a mode concentration reproducing the one observed in the PDB refine even better than expected.

Limitations of the Model. At first sight, it might seem that a severe limitation of the method comes from the use of a highly partial model of the final structure, namely only the C-alpha coordinates are used both in the normal mode calculation and in the refinement process. However, several authors have argued convincingly that NM derived from C-alphas only (and not from the full atomic model) work well (5, 6). Two separate questions remain to be addressed.

First, the score for the correct model cannot be perfect if the model is partial, so the contrast between the correct answer and the starting model is not maximum, i.e., as large as it could be if the model were complete. However, we show here that this contrast is good enough to lead to a satisfactory refinement of NM amplitudes using real experimental data (see above).

Second, how can one generate the complete atomic model from...
the refined C-alpha coordinates? Here, we use techniques derived from homology modeling. There are actually at least two possibilities. The first one is to reconstruct all main chain atoms from C-alpha only, using fragments of the PDB database (30) as, for instance, implemented in the lego_auto.mc option in O (31) and then build side chains by using, e.g., mean-field techniques (32).

The second one is to interpolate displacement vectors at every atom position and then regularize the stereochemistry of the chain by using any molecular dynamics force field. We chose the second method with excellent results.

Fitting Models into Envelopes. It might seem that the approximation used here (to represent as a set of lattice points the model to be refined into a given experimental envelope) is a very gross one. However, it was shown recently that normal modes derived from such an approximation still correlate well with the ones obtained from the starting envelope. It proved very easy to interpolate the field of displacement vectors on a regular lattice very appealing for further analysis using graphics programs such as O (31).

For small-angle x-ray scattering data, the fitting could be done either in envelopes reconstructed from the experimental data (36, 37), or directly in reciprocal space, because the method presented here expresses the constraints in reciprocal space anyway. Further work is needed to assess which of the two methods is best.

Application to the MR Method. Highly partial models such as C-alpha-only models can be used and refined in MR techniques, as documented elsewhere (38, 39). The refinement program described here was successfully used to discriminate between alternative solutions of the MR problem, in a manner reminiscent of the Patterson-correlation-refinement method (40).

On a more general level, once the correct solution of the target envelope is defined, there is a definite need to start from the best possible model. This is because it is very difficult to get rid of errors in the initial model, which propagate during the refinement through phase combination. Even though phase combination allows for a better (safer) refinement process, there is still a definite advantage in refining early on (i.e., at low resolution) the molecular form of the model in the best possible way. NM refinement allows for such a possibility in cases where large-amplitude collective movements occur and where dihedral angle dynamics fail.

NM refinement is better than rigid-body refinement because there is no need for the user to define the movable domains, which are found by NMA. For shear movements as in citrate synthase, this feature is a definite improvement on existing methods because the movable domains are very hard to define by visual inspection. And, indeed, to explore large-amplitude rearrangements of the molecule, it seems natural to explore first the “most natural movements” of the protein, which are given by a linear combination of the lowest-frequency normal modes-associated directions.

Conclusion and Perspectives

In summary, we have shown here that refinement in reciprocal space at medium or low resolution with respect to NM amplitudes is very efficient, with a very large radius of convergence that goes beyond the capacities of pure rigid-body refinement and also of dihedral angle refinement. There is no danger of overfitting the data because the number of parameters is very small; in addition, the \( R_{\text{free}} \) indicator can be used to ensure that overfitting is not taking place. We checked in test cases that, even when degrees of freedom are allowed, they refine correctly to almost zero amplitude. With real data, NM refinement proceeded correctly even though the model contains only one atom per residue.

Envelope refinement was made possible through the use of a points approximation of the envelope, from which it was possible to interpolate the deformations of C-alpha coordinates of the molecule. Alternatively, C-alpha models can also be refined directly. We expect that the extraction of the normal modes amplitudes best fitting a molecular envelope change upon ligand binding should be extremely useful in the quickly expanding field of structural studies of large macromolecular biological complexes and/or molecular motors (43).

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