On the use of low-resolution data for translation search in molecular replacement
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Low-resolution reflections (approximately 15 Å and lower) are very useful for the translation search in molecular replacement because they are less sensitive to model errors compared with the traditionally used reflections of resolution 4–10 Å. At low resolution, however, the contribution from the bulk solvent is quite significant and corresponding structure factors calculated from a macromolecular model cannot be compared with experimental values if this contribution is neglected. The proposed method provides a way of fast translation searches where low-resolution reflections are taken into account. Test calculations using several experimental data sets show a dramatic improvement in the signal after the bulk-solvent correction and low-resolution reflections were included in the calculation; this improvement allowed unambiguous identification of the solution.

1. Introduction

Molecular replacement (MR hereinafter; Rossmann, 1972) is one of the key methods for macromolecular structure determination which becomes even more important with the development of structural genomics projects. Unfortunately, the quality of MR models is sometimes quite poor and the solution cannot be found because it does not correspond to the optimum of usual search functions (for example, to the maximum of the correlation of experimental magnitudes of structure factors with corresponding values calculated from the model placed differently in the unit cell). As a consequence, even the simultaneous rotation and translation search does not help; on the contrary, a separate translation search does not help; on the contrary, a separate translation search much more efficient. A fast way for such a search is suggested.

2. Methods and test calculations

It may be noted that the basic assumption of the exponential solvent model, namely the proportionality of the molecular and solvent structure factors, is not usually fulfilled at a resolution of about 10–20 Å (Urzhumtsev & Podjarny, 1995a) and that the flat mask approach (Jiang & Brünger, 1994) is of a superior quality (Kostrewa, 1997).

The principal steps of this latter procedure are as follows:

(a) the molecular envelope (a binary function) is determined from the atomic model;

(b) structure factors $F_{\text{env}}$ are calculated as the Fourier coefficients of this function;

(c) scale parameters $k_{\text{sol}}$ and $B_{\text{sol}}$ are estimated by minimizing the residue

$$
\sum_s \left[ (F_{\text{model}}(s) - k_{\text{sol}} \exp(-B_{\text{sol}}|s|^2/4)F_{\text{env}}(s))^2 - F_{\text{obs}}(s))^2 \right];
$$

(d) the complex numbers $F_{\text{obs}} = -k_{\text{sol}} \exp(-B_{\text{sol}}|s|^2/4)F_{\text{env}}(s)$ calculated with the optimal parameter values are taken as the solvent structure factors.

Here, $F_{\text{model}}(s)$ is the structure factor calculated from an available atomic model and $F_{\text{obs}}(s)$ is an experimental structure-factor magnitude for the reflection $s$; for simplification, all values are supposed to be known on the absolute scale.

For the translation search, such a solvent correction can eventually be performed at every position of the search model (while the obtained $k_{\text{sol}}$ and $B_{\text{sol}}$ can be completely unreasonable for incorrect positions). However, such a method of solvent correction cannot be included in fast translation algorithms (Navaza, 1994; Navaza & Vernoslova, 1995), making its practical application inefficient. In
order to realize a fast search, we suggest a technique based on the following observations:

(a) Our statistical analysis of the structures deposited in the Protein Data Bank (PDB) (Bernstein et al., 1977) shows that a large majority of the models have a $k_{\text{sol}}$ value between 0.30 and 0.40 e Å$^{-3}$ and a $B_{\text{sol}}$ value between 30 and 70 Å$^2$, and that some existing extreme values are rather unjustified (Fokine & Urzhumtsev, 2002); the much smaller dispersion of these values in comparison with the dispersion of the parameters for the exponential model (Glykos & Kokkinidis, 2000) is due to a more physical meaning of the parameters of the flat mask model; the use of the mean values of $k_{\text{sol}}$ and $B_{\text{sol}}$ are used for all model positions, this allows one to calculate the solvent correction for an isolated model, add it to the model structure factors of this model and perform a search with the known FFT-based procedures (see, for example, Navaza & Vernoslova, 1995) using bulk-solvent-corrected structure factors instead of those calculated directly from the atomic model.

(b) For the positions in the unit cell where the search model does not overlap with its symmetrically related images, the mask of the region occupied by all molecules can be calculated as a junction of masks of individual molecules related by symmetries; as a consequence, the structure factors of such a total molecular envelope can be rapidly recalculated from the structure factors of the envelope of a single model; if the standard mean values of the parameters $k_{\text{sol}}$ and $B_{\text{sol}}$ are used for all model positions, this allows one to calculate the solvent correction for an isolated model, add it to the model structure factors of this model and perform a search with the known FFT-based procedures (see, for example, Navaza & Vernoslova, 1995) using bulk-solvent-corrected structure factors instead of those calculated directly from the atomic model.

(c) Such an approach does not provide an adequate bulk-solvent correction for the positions where the models overlap, and spurious peaks in the translation function are eventually possible at such points; these spurious peaks, however, will be eliminated by the packing criterion and will not appear in the final list.

In order to construct a good molecular envelope, an atomic model is supposed to be more or less complete. Such a situation is usual when NMR models are used as templates [for a review of the MR searches with NMR models, see Chen et al. (2000); cases reported in this work as most difficult were chosen for our tests below (Table 1)].

All test calculations were performed using experimental data, and the orientation of the search models was supposed to be known (it may be noted that typical errors in model orientation practically did not influence the searches when low-resolution reflections were included; not shown in this paper). All translation searches were made using CNS (Brünger et al., 1998) using the fast translation function (Navaza & Vernoslova, 1995). The translation search parameters were taken without any optimization; complete NMR models were taken as they are in the PDB; the $B$ factors for all atoms of the search models were assigned to be equal to 20 Å$^2$ which is far from the optimal scheme (Chen et al., 2000). In each test, a single NMR model was used for the translation.

### Table 1
Test structures: summary information.

<table>
<thead>
<tr>
<th>Protein name (reference)</th>
<th>Crystal structure</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>PDB ID</td>
</tr>
<tr>
<td>Human interleukin-4 (Müller et al., 1995)</td>
<td>1hik</td>
</tr>
<tr>
<td>PS3 tetramerization domain (Müller et al., 1998)</td>
<td>1aie</td>
</tr>
<tr>
<td>Corn Hageman factor inhibitor (Behnke et al., 1998)</td>
<td>1bea</td>
</tr>
</tbody>
</table>

### 3. Results and discussions

Fig. 1 shows the results of the translation searches performed with and without the bulk-solvent correction, using three experimental data sets. Each pair of diagrams shows the results of the translation search at a given resolution shell:
the right-hand diagrams show the peaks obtained in the translation search without bulk-solvent correction; the left-hand diagrams show the peaks obtained under the same conditions when the bulk-solvent correction was taken into account, as suggested above. The height of each peak is shown as a percentage of the height of the first peak of the corresponding search, and the correct solution is indicated in grey.

For human interleukin-4 (Müller et al., 1995), the translation search performed at the standard resolution of 4–15 Å without solvent correction gave the solution as the second peak (Fig. 1a, left). When all available reflections with resolution lower than 15 Å were also included in the calculations, this brought the correct peak to the first position without significant contrast (Fig. 1a, right). The following bulk-solvent correction increased the contrast of the signal drastically; the best results were obtained when low-resolution data were included.

In the case of p53 tetramerization domain (Mittl et al., 1998), no experimental data below 15 Å are available. Without solvent correction, the translation search at the standard 4–15 Å resolution gave the correct solution hidden in noise (Fig. 1b, left) and the search at 3–15 Å resolution gave it slightly higher in the list (Fig. 1b, right). With the solvent correction, the peak for the solution became the first with the best contrast at 4 Å when higher-resolution data were suppressed [see Urzhumtsev & Podjarny (1995b) for similar observations when searches are performed with molecular envelopes].

Corn Hageman factor inhibitor (Behnke et al., 1998) was reported as the worst case among all NMR-based searches (Chen et al., 2000). The orientation of the search model can be found very surely and precisely by a multiple rotation function (Urzhumtsev & Urzhumtseva, 2002). Without the bulk-solvent correction, the solution did not appear among the ten highest peaks, neither at 4–15 Å nor at 5–15 Å resolution (Fig. 1c, left), and appeared only as the seventh peak when all available magnitudes with resolution lower than 5 Å were used (Fig. 1c, right). At the same time, with the bulk-solvent correction, this peak became the first for a resolution lower than 5 Å while the contrast was not as high as for the two previous cases. Better contrast can be obtained by variation of the MR parameters, which is the subject of further analysis.

More complete results are out of the scope of this paper.

4. Conclusions

Low-resolution reflections are very useful for the translation search in molecular replacement. Because the solvent contribution to them is very significant, a corresponding correction is necessary in order to use these reflections properly. The flat model for the bulk-solvent correction, the best model currently available, can be easily included in the translation search. The use of low-resolution reflections, especially with the bulk-solvent correction, solves a number of translation problems which are difficult to solve by standard procedures.

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References


