Iterative build OMIT maps: Map improvement by iterative model-building and refinement without model bias

Thomas C. Terwilliger\textsuperscript{a}, Ralf W. Grosse-Kunstleve\textsuperscript{b}, Pavel V. Afonine\textsuperscript{b}, Nigel W. Moriarty\textsuperscript{b}, Paul D. Adams\textsuperscript{b}, Randy J. Read\textsuperscript{c}, Peter Zwart\textsuperscript{b}, Li-Wei Hung\textsuperscript{a}

\textsuperscript{a}Los Alamos National Laboratory, Los Alamos, NM 87545, USA.
\textsuperscript{b}Lawrence Berkeley National Laboratory, One Cyclotron Road, Bldg 64R0121, Berkeley, CA 94720, USA.
\textsuperscript{c}Department of Haematology, University of Cambridge, Cambridge CB2 0XY, UK

Abstract A procedure for carrying out iterative model-building, density modification and refinement is presented in which the density in an OMIT region is essentially unbiased by an atomic model. Density from a set of overlapping OMIT regions can be combined to create a composite “Iterative-Build” OMIT map that is everywhere unbiased by an atomic model but also everywhere benefiting from the model-based information present elsewhere in the unit cell. The procedure may have applications in the validation of specific features in atomic models as well as in overall model validation. The procedure is demonstrated with a molecular replacement structure and with an experimentally-phased structure, and a variation on the method is demonstrated by removing model bias from a structure from the Protein Data Bank.

Keywords: Model building; model validation; macromolecular models; Protein Data Bank; refinement; omit maps; bias; structure refinement; PHENIX

1. Introduction

Model bias is a continuing problem in macromolecular crystallography. It results from using an atomic model to calculate crystallographic phases, in which case the resulting electron density map will tend to have the features present in the model, even if they are not actually present in the structure (Ramachandran & Srinivasan, 1961; Read, 1986; Bhat, 1988; Hodel et al., 1992; Adams et al., 1999; Kleywegt, 2000). Once an atomic model is refined, model bias can be indirect as well as direct because the positions and other parameters describing correctly-placed atoms are adjusted during refinement to compensate for the incorrectly-placed atoms. Consequently even if the incorrectly-placed atoms are removed from the model before calculation of phases, a memory of their positions can remain and the resulting map can retain incorrect features. Refinement of a model omitting incorrectly-placed atoms should reduce this indirect bias, but there remains a question of how extensive the
refinement must be to reverse all compensating adjustments. Model bias can make interpretation of electron density maps difficult, particularly in cases where molecular replacement (Rossmann, 1972) is used to solve a structure.

There are many ways that model bias could be defined. In this work model bias refers to the situation where a map has peaks of density resembling atomic density due only to atoms in the working model used in phasing or density modification and not to the presence of atoms in the real structure.

There have been many methods developed to reduce the effects of model bias. These fall into two general classes. The first class consists of methods to remove the model bias after it has been introduced. The second class consists of methods in which a model is never introduced in a particular location of a map, so that in that location there is never any model bias at all.

The reason to use an approach in which model bias is introduced and then removed is that the process of building and refining a model greatly improves the overall accuracy of phase information, and often this is required to obtain a structure at all. Once an initial structure is determined, it becomes important to know which details are correct, and removal of model bias can be important for this. A well-established method for reducing model bias is to calculate a map using $\sigma_A$-weighted $(2mFo-DFc) \ e^{i\phi}$ coefficients (Read, 1986, Lunin et al., 2002), typically using coefficients $m$ and $D$ calculated using a test set of reflections (Urzhumtsev et al., 1996, Pannu & Read, 1996). The $\sigma_A$ procedure yields high-quality maps, but they can retain some model bias because the procedure is based on the assumption of random errors in the model, while actual errors after model refinement are typically correlated (Read, 1997). Simulated-annealing OMIT maps (Hodel et al., 1992; Brünger et al., 1998), “kicked” OMIT maps (Guncar et al., 1998), and model rebuilding with randomization (Zeng et al., 1997; Reddy et al., 2003) can all reduce model bias by removing, at least to some extent, the memory of some or all of the atoms in the model. Prime-and-switch density modification (Terwilliger, 2004) reduces the effects of model bias in a different way. This approach uses a model to calculate phases, yielding a map that is biased but nearly correct, and then using a characteristic of this map that is relatively uncorrelated with the model, such as the flatness of the solvent region, to calculate less-biased phases. Although these procedures can reduce the effects of model bias, they all have the disadvantage that it is not possible to know for certain that model bias is removed. Additionally, techniques that involve a solvent mask have some potential for model bias as the mask can be influenced by the model.

The second class of methods are those that never introduce model bias, and consequently can yield a higher degree of confidence in electron density that is in the same location as an
atom in a model. The simplest method in this class is an OMIT map in which the atoms in a particular region of the map are never included in phase calculation or in refinement of the other atoms in the structure (Bhat, 1988). In a molecular replacement structure determination, this type of OMIT map can be easily calculated early in the process, after the molecule has been placed in the correct location in the unit cell, but before any atomic refinement has occurred. In this case the memory of the presence of the omitted atoms is likely to be minimal, and the resulting map is unbiased in the OMIT region. The disadvantage of OMIT maps is that they are typically very noisy and consequently difficult to interpret. Additionally, once the structure (with all atoms) has been refined, OMIT maps are of less utility because of the indirect model bias described above.

Another approach that avoids introducing model bias is to carry out the usual process of iterative model-building and refinement, but to avoid building a particular part of the model. The part of the model deliberately not built might be a ligand, a side-chain, or any other part of the model. This approach is commonly used for poorly-defined portions of electron density maps and has been used in some cases specifically to obtain unbiased information (James et al., 1980). Such poorly-defined regions are typically not interpreted until the improvement in the rest of the model is sufficient to yield clear electron density for a model in those regions. This approach has recently been extended into a systematic procedure (“Ping-pong refinement”) that allows each of the side-chains of a structure to be built into density that is unbiased, while gradually building up a complete atomic model (Hunt & Deisenhofer, 2003).

Here we combine the ideas of OMIT maps and Ping-pong refinement in an “iterative-build OMIT” procedure for obtaining a partial or complete composite electron density map that is essentially free from model bias, yet that benefits from the power of iterative model-building and refinement.

2. Methods

2.1. Calculation of an iterative build OMIT map for a single omit region

Obtaining an iterative build OMIT map for a single region of the asymmetric unit of a crystal is in principle quite straightforward. First the OMIT region is defined as a contiguous region representing part of the asymmetric unit. Then a border (typically 2 Å thick) is added to this region. Finally an iterative model-building, refinement and density modification procedure is carried out in a standard fashion (Terwilliger, 2003), except that any atom that is located within the OMIT or border regions is given a zero occupancy in all calculations. We call this overall process the “Iterative-build omit procedure.” In this procedure all atoms in the omit region are included in geometric restraint calculations and are included in all rebuilding
steps. In this way the geometry of the model is retained. Due to their zero occupancy values, however, the atoms in the OMIT region do not contribute to the structure factor calculation. This prevents direct model bias. This procedure also prevents indirect model bias in the density calculated within the OMIT region, as parameters of atoms outside the OMIT region are never adjusted to compensate for electron density of atoms in the OMIT or border regions.

To ensure that model bias is not indirectly introduced through the application of NCS, in the RESOLVE density modification steps no NCS-based target electron density is transferred into the OMIT or border regions. This is accomplished by defining the boundaries of the OMIT region and specifying that no NCS information is to be transferred into this region. As density modification with NCS is done point-by-point, using the density from N-1 copies as a target for density modification for the remaining copy (Terwilliger, 2002), it is straightforward to leave out NCS information for all points in the OMIT region. Once the final electron density map is obtained from the iterative-build omit procedure, the OMIT region (but not the boundary region or other parts of the map) will have essentially no bias due to structure factor contributions from the atomic model. As parameters describing the NCS relationships are refined there is in principle some possibility of model information to be transferred between NCS regions As in the case of rigid-body refinement of an MR model, however, it is not likely that significant information about the density in a particular location in the map is going to be transmitted through the very small number of parameters refined in this step. In the standard PHENIX model-building procedure in the presence of NCS, all building is done independently for all copies, and in a specific step the structure of each NCS copy is transformed to match each other one, and the best parts of the structure from each NCS copy are kept. This step is not carried out when an OMIT map is constructed, so that no information about the structure within the OMIT region is transmitted among the NCS copies. NCS restraints are applied during model-building with the OMIT procedure. The effect of this is that the (zero-occupancy) atoms in the OMIT region may be placed in incorrect positions because of the NCS restraints from copies outside the OMIT region.

Other potential sources of model bias are the bulk-solvent correction and geometric restraints. In the procedure carried out here, a bulk solvent model and geometric restraints are applied throughout, as not applying them would lead to a poorer atomic model in the regions outside the OMIT boundaries and would therefore result in an OMIT map with greater artefacts and less utility.

We note that there are some circumstances where the density within an OMIT region can be affected by the positions of atoms in the model inside the OMIT region. For example, an atom inside an OMIT region may be bonded to an atom outside the OMIT region, so that the positions are correlated. Similarly, the position of an atom inside the OMIT region could
affect the placement of a solvent molecule outside the OMIT region, or the allowed conformations of a side-chain outside the OMIT region. Further, during automated model-building, large units (helices, strands) may be placed based on density both inside and outside the OMIT region. In all these cases, however, this coupling between atoms inside and outside the OMIT region is unlikely to lead to density at the positions of the atoms inside the OMIT region. Consequently there is unlikely to be any model bias (density at the coordinates of atoms in the model, due to the presence of those atoms in the model) in the resulting maps.

In our procedure OMIT regions are constructed so that they tile to fill the asymmetric unit. Normally approximately 10-20 OMIT regions are used to cover the asymmetric unit, but more (as many as 132 in our tests) may be chosen so as to have a minimal impact on the density modification procedure. Due to this way in which OMIT regions are chosen, there may be some OMIT regions that contain no atoms from the macromolecule and others with many atoms. Those OMIT regions that contain many atoms typically have poor electron density compared to those with few atoms, as density from the atoms in the OMIT region is excluded from contributing to the density modification procedure. It may be possible to improve the procedure by defining variable-sized OMIT regions that contain more equal numbers of atoms.

The reason for adding a border region around the OMIT region is that a peak of density in electron density maps calculated from a model containing a particular atom has a substantial contribution within a radius that may depend on the resolution of the data and atomic displacement factors. We use the value of 2 Å for the thickness of the border region based on previous experience with composite omit maps in CNS (Brünger et al., 1998).

The approach of never including atoms in an omit region is applicable in a straightforward way to cases where a model is being built without reference to an existing model. In cases where molecular replacement is used, ideally the atoms in the omit region should be omitted from the very start of the procedure, so that the placement of the molecule and any rigid-body refinement carried out are not affected by the atomic positions of these atoms. In practice this is probably unnecessary however, as the number of parameters being refined in placement of the molecule are so few that little information about positions of specific atoms in the OMIT region can be retained.

We note that the OMIT procedure described here has a kind of negative model bias. In the OMIT region a model is built, but occupancies are set to zero. No solvent atoms are placed at the locations of these atoms, and no bulk solvent model is placed there. Consequently there is low (zero) density near the locations of atoms in the model within the OMIT region. It seems possible that the procedure could be improved by setting the density in these locations to an intermediate value rather than to zero.
Though this procedure is straightforward in concept, it is somewhat less simple in implementation as all programs that operate on models that are built during iterative model-building, density modification and refinement need to keep track of which atoms are in the OMIT and border regions. These steps have been implemented in the PHENIX (Adams et al., 2002) AutoBuild Wizard by using the RESOLVE omit box generation procedure to specify the OMIT region and its boundary and to identify which atoms in a model are within these regions. Additionally, all density modification procedures with RESOLVE are called with the specification of the omit region and boundary so that that no model-based information is transmitted into these regions through the application of NCS. Standard procedures for iterative model-building, density modification and refinement are used as implemented in the PHENIX AutoBuild Wizard.

2.2. Calculation of a composite iterative-build OMIT map

A composite iterative-build OMIT map can be calculated by dividing the asymmetric unit of the crystal into a set of OMIT regions, calculating an iterative-build OMIT map for each region and its boundary as described above, and then simply combining the OMIT regions of all the iterative-build OMIT maps. This method of combining OMIT maps is similar to that used to create composite omit maps in CNS (Brünger et al., 1998). The resulting iterative-build composite OMIT map has the property that the density at each point in the map has never been affected by the presence of a model atom near that point (or near any NCS-related point). It should be noted that due to the way such a map is constructed, it can potentially have discontinuities at the boundaries between OMIT regions, although we have not noticed any in the maps we have examined.

The calculation of an iterative-build OMIT map can be carried out automatically using the “omit_type=composite_omit” keyword in version 1.3b or higher of the PHENIX AutoBuild Wizard (Adams et al., 2002), available at http://www.phenix-online.org.

3. Results and Discussion

3.1. Iterative build composite OMIT map for antitrypsin with molecular replacement

We tested the use of an iterative-build composite OMIT map by applying it to repeat the structure solution of antitrypsin (Kim et al., 2001, entry 1HP7 in the PDB; Bernstein et al., 1977; Berman et al, 2000) by molecular replacement, using the structure of antichymotrypsin (PDB entry 1AS4; Lukacs et al., 1998) as a search model. We used the PHENIX AutoMR Wizard which calls Phaser (McCoy et al., 2005) to obtain an initial molecular replacement
solution. The PHENIX AutoBuild Wizard was then used to edit the sequence of the search model to match that of antitrypsin, truncating side-chains at the C$_\alpha$ or C$_\beta$ atoms if the remainder of the atomic positions were unknown. The AutoBuild Wizard was then used to define 128 OMIT regions covering the asymmetric unit of the crystal.

To examine the properties of OMIT maps, the starting molecular replacement model was then used to calculate electron density maps in two ways. First, the occupancies of all atoms in one OMIT region were set to zero, then the entire structure was refined with the PHENIX refinement package phenix.refine (Afonine et al., 2005b) without rebuilding manually or with RESOLVE. The standard refinement procedure used included three macrocycles of refinement with automated estimation of parameters in a bulk solvent model and placement of solvent molecules (Afonine et al., 2005a), individual atomic coordinate shifts, and isotropic atomic displacement parameters. This refinement yielded an R/Rfree of 0.41/0.48, and a $\sigma_A$-weighted (2mFo-DFc) $e^{i\phi}$ OMIT map was calculated (Read, 1986). This $\sigma_A$-weighted OMIT map (Fig. 1A) has some features corresponding to a helix in the omitted region, but it is very difficult to interpret.

The second type of map calculated was an iterative-build OMIT map (Fig. 1B) in which the occupancies of all atoms in the same OMIT region as in Fig. 1A were maintained at a value of zero for 10 cycles of iterative model-building, density modification and refinement. This map has very clear features of helical density in the omitted region, despite the fact that density was never calculated using a model for any atoms in this region. Figs. 1C and 1D show that OMIT regions such as the one illustrated in Figs. 1A and 1B can be joined together to form a composite iterative-build OMIT map that has clear electron density for much of the structure. Fig. 1E shows the starting model superimposed on the final composite iterative-build OMIT map. It can be seen in Fig. 1E that the helix in the starting model is offset from the final position of the helix density by about 2 Å. Fig. 1F shows that for this OMIT region, a map similar to the OMIT map can also be obtained by deleting all the atoms that are in the OMIT region completely, then carrying out a standard iterative-build procedure (with no OMIT regions).

The reason why the density in the iterative-build OMIT map in Fig. 1B is so much improved over the standard OMIT map in Fig. 1A is that the model outside the omitted region was much more accurate after the iterative model-building process, and this model is used as a source of information for density modification. For the 128 OMIT procedures carried out, the mean final R and Rfree (including only the part of the model outside the omitted region) were 0.29 and 0.34, respectively (compared with the starting R/Rfree of 0.41/0.48). The range of R-factors was from 0.25 to 0.38 and the range of free R factors from 0.30 to 0.44 (the low R-factors are paired with low free R factors). The range of both sets of R-factors is quite...
large, and the higher R-factors typically correspond to OMIT regions containing larger numbers of atoms.

3.2. Iterative build composite OMIT map for SAD-phased gene 5 protein

We also tested the use of an iterative-build composite OMIT map for a case where experimental phases were available. It seemed possible that a density-modified map could be created that had no potential of model bias, but that benefited from the use of iterative model-building, density modification and refinement. Experimental phases were obtained using SOLVE SAD phasing (Terwilliger & Berendzen, 1999) with single wavelength anomalous dispersion (SAD) from gene 5 protein in this test. The structure of gene 5 protein was determined previously by MAD (PDB entry 1VQB; Skinner et al., 1994); in this test only the data corresponding to the peak wavelength were used, simply to yield a poorer starting set of phases.

The AutoSol Wizard was used to re-solve the structure of gene 5 protein, yielding a starting density-modified SAD-phased electron density map with a correlation coefficient of 0.71 to the model electron density map calculated from the refined gene 5 structure (Fig. 2A). The AutoSol Wizard uses the initial density-modified map for model-building and does not do iterative model-building and refinement. It built 64 of 87 residues of the protein, and (correctly) docked 9 residues into sequence. The R and Rfree for this model were 0.46 and 0.47, respectively. The AutoBuild Wizard was then used to calculate an iterative-build composite OMIT map (Fig. 2B). During refinement the experimental phases were used as restraints (Pannu et al., 1998). The mean R and Rfree for the final models built to construct the 132 OMIT regions were 0.26 and 0.34, respectively. The composite iterative-build map has an improved correlation coefficient of 0.82 to the model electron density map calculated from the refined gene 5 structure. This improvement relative to the starting density-modified SAD-phased map comes from including model information from outside each omit region in the phase calculation for that omit region. In effect, both the map in Fig. 2A and the map in Fig. 2B can be thought of as density-modified maps. They differ in that the density modification used in Fig. 2B includes model information, while that in Fig. 2A does not, and both are essentially free of model bias.

We note that iterative-build OMIT maps may normally be unnecessary for models autobuilt into experimental electron density maps. The procedures typically used in autobuilding have a high cutoff for density, and incorrectly-placed atoms are normally removed in subsequent cycles, so that it is uncommon for atoms to be repeatedly placed in very incorrect positions, as would be required to introduce model bias. A map for this structure can also be calculated by carrying out the iterative building procedure with no
OMIT regions; this map is slightly better than the OMIT map, with a correlation coefficient with the model map of 0.85.

### 3.3. Using iterative-build OMIT maps to remove existing model bias

Although the principal intent of the iterative-build OMIT procedure described here is to avoid model bias entirely, it seemed possible that the process of rebuilding a model outside an OMIT region might be useful in removing existing model bias as well. In particular, we would expect that extensive rebuilding should effectively remove adjustments to atoms in the rest of the model that compensate for incorrectly-placed atoms. To test this idea, we identified an entry in the PDB with some features that were likely to be incorrect, but which remained in a $\sigma_A$-weighted $(2mFo-DFc)\phi$ map (Read, 1986) calculated after refinement of the structure. PDB entry 1ZEN (Cooper et al., 1996) was such a structure (G. Kleywegt, personal communication). It was obtained at a resolution of 2.5 Å, and the closely-related structure 1B57 (Hall et al., 1999), determined later at a resolution of 2 Å, differs in the sequence register of residues 6-16 by one residue.

Fig. 3, panels A and B show a $\sigma_A$-weighted $(2mFo-DFc)\phi$ map calculated after re-refinement of 1ZEN (without omitting any atoms) with phenix.refine using deposited structure factors and re-generating a test set of reflections for refinement (the original test set was not available). The final R/Rfree after refinement was 0.25/0.29. Fig. 3A shows the map with residues 5-10 from 1ZEN, centered on residues F6 and K8 which are likely to be misaligned by one residue in this structure. Fig. 3B shows the map with residues 3-10 of chain A from PDB entry 1B57, after superimposing this chain from 1B57 on the structure from 1ZEN by least squares. The $\sigma_A$-weighted $(2mFo-DFc)\phi$ map has features in common with both the structure used to generate the phases for this map (1ZEN) and with the structure derived from the higher-resolution model (1B57), including several features that appear to be examples of model bias in the map. In particular the map shows density for the side chain of K8 from the 1ZEN structure even though the more likely 1B57 structure has only a carbonyl oxygen pointing towards this location. Overall, in the neighborhood of residues 5-10 of 1ZEN, the $\sigma_A$-weighted $(2mFo-DFc)\phi$ map is somewhat more similar to a model map calculated from the higher-resolution structure, 1B57, than to a model map calculated from 1ZEN. This is shown numerically by local map correlation coefficients, summarized in Table 1 for this map and the maps discussed below.

It seemed possible that the model bias found in the $\sigma_A$-weighted $(2mFo-DFc)\phi$ map shown in Figs. 3A and 3B could be due in part to the fact that we did not have access to the original test set used in refinement. To examine this possibility, we carried out a second
refinement in which the atoms in the structure were displaced by an rms distance of 1.0 Å with the “shake” procedure (Brünger et al., 1998; Guncar et al., 1998), yielding a starting R/Rfree of 0.48/0.47, followed by 6 cycles of refinement with phenix.refine, leading to a final R/Rfree of 0.25/0.30. The resulting $\sigma_A$-weighted $(2mFo-DFc)$ $\epsilon^{\phi}$ map is shown in Figs. 3C and 3D. It appears to have slightly less model bias than the map calculated after standard refinement, but overall the maps are very similar.

Next we carried out a more extensive re-refinement procedure to try to reduce model bias. The partially-randomized model from the “shake” procedure above was taken as a starting point, all solvent molecules (waters) were removed, and 10 cycles of refinement and water picking, including two cycles with simulated annealing, were carried out. Once again the starting R/Rfree was 0.48/0.47 and the final R/Rfree was 0.25/0.30. The resulting map has less model bias than the starting $\sigma_A$-weighted $(2mFo-DFc)$ $\epsilon^{\phi}$ map (Figs. 3E, 3F), and lowered correlation with the 1ZEN model (Table 1), but it would still be difficult to decide which of the two models is correct because of the model bias showing density for the entire side chain of K8 from the 1ZEN structure.

Three types of OMIT maps were then calculated using the coordinates and structure factors from 1ZEN. In each case, the occupancies of all the atoms inside a small OMIT box were set to zero before initial refinement and throughout the procedures. The OMIT box was defined as a region with edges parallel to the cell edges and 4 Å from the nearest atom in residues 5-9 of 1ZEN. The OMIT maps calculated in these procedures are therefore not based on any density coming from any atoms in residues 5-9 of 1ZEN.

The first OMIT map calculated (Figs. 3G, 3H) was a simple omit map, in which the structure of 1ZEN was refined with zero occupancies for the atoms in the OMIT region and a $\sigma_A$-weighted $(2mFo-DFc)$ $\epsilon^{\phi}$ map was calculated. This map still shows model bias from 1ZEN at the side chain of K8 (Fig. 3G, Table 1).

The second OMIT map was a simulated-annealing OMIT map (Brünger et al., 1998) calculated with phenix.refine (Afonine et al., 2005b; Figs. 3I, 3J). In calculating this map the refinement started at a pseudo-temperature of 5000K and cooled to a final temperature of 300K. This map shows substantially less model bias but has relatively weak density for the entire segment, resulting in lower correlations with the density from both models (Table 1).

The third OMIT map was an iterative-build OMIT map (Figs. 3K, 3L). To create this map the model, with zero-occupancy atoms in the OMIT region, was rebuilt three times using the PHENIX rebuild-in-place algorithm. In this rebuilding procedure the polypeptide chain is rebuilt by iteratively removing a segment and retracing the chain for that segment. Then the parts of the resulting models that best fit a density-modified map are combined and side-
chains are re-fit into the density. This iterative-build OMIT map shows little model bias from 1ZEN (Fig. 3K), and matches the model from 1B57 well (Fig. 3L, Table 1).

We also tested whether a \( \sigma_A \)-weighted \((mFo-DFc) e^{i\phi_c}\) map calculated from the 1ZEN model might be as informative as the OMIT maps we have calculated. We downloaded \( \sigma_A \)-weighted \((2mFo-DFc) e^{i\phi_c}\) and \( \sigma_A \)-weighted \((mFo-DFc) e^{i\phi_c}\) maps from the EDS server (Kleywegt et al., 2004). Fig. 3M shows the \((2mFo-DFc) e^{i\phi_c}\) map, which is as expected essentially identical to the map calculated by phenix.refine and shown in Fig. 3A. Fig. 3N shows the difference \((mFo-DFc) e^{i\phi_c}\) map, however there is no negative (or positive) difference density at the coordinates of the mis-placed lysine side chain.

Considering all of these maps, along with the differences between the lower-resolution 1ZEN and higher-resolution 1B57 models, the simplest interpretation of these results is that the higher-resolution 1B57 model is the more accurate of the two in the region that we have examined, and that the iterative-build OMIT map is particularly useful in reducing model bias without much cost to overall map quality.

4. Conclusions

The iterative-build OMIT procedure can be thought of as a type of density modification that involves the use of a model outside of the OMIT region. In statistical density-modification procedures, model density can be used as the expected value of the electron density. Phases are then adjusted to better match the map to this expected density (Terwilliger, 2003). In the iterative-build OMIT procedure, the model is built wherever it exists, but only model density outside the OMIT region is used in density modification. In this way the model density can improve the quality of the crystallographic phases, yet not directly bias the density in the OMIT region.

The iterative-build OMIT procedure can be of substantial use in molecular replacement in situations where initial refinement of the molecular replacement model yields relatively poor R-factors, but in which iterative model-building, density modification and refinement yields a greatly improved model with lowered R-factors. In such a case a \( \sigma_A \)-weighted \((2mFo-DFc) e^{i\phi_c}\) OMIT map calculated after initial refinement may be relatively uninformative because the model is not yet good enough to produce phases that lead to a clear map (e.g., Fig. 1A). On the other hand, after iterative model-building, density modification and refinement, the model built outside the OMIT region can be accurate enough to yield phases that clearly show the density inside the OMIT region (e.g., Fig. 1B). In the case shown in Fig. 1 for example, the starting R-factor after refinement of the model was 0.41, but the iterative building procedure yielded much lower R-factors (mean of 0.29), and produced a very clear map for the omitted region.
The generation of iterative-build composite OMIT maps can be computation-intensive, particularly in cases where there is only just enough phase information for the iterative model-building procedure to improve upon the starting model. In such cases, the size of the OMIT regions must be very small or no phase improvement results. Consequently in some cases many OMIT regions must be constructed, and the entire iterative model-building procedure must be carried out many times (132 OMIT regions were combined in the case shown in Fig. 2B). The procedure is readily made parallel, so as highly parallel machines and large clusters become increasingly available, the procedure may become practical for even very large structures. At present, the procedure can be time-consuming for a very large structure when run on a single processor, as several cpu-days can be required for each OMIT region. A use of the iterative-build OMIT procedure that is quicker is to construct an unbiased map for a small region within an electron density map, such as for the density in the vicinity of a ligand or side-chain of interest. In such a case a single OMIT map can often be calculated, setting to zero the occupancies of all atoms in a box containing the region of interest during the iterative-build process. This local application of the iterative-build OMIT procedure may also be of substantial use in checking for errors that may be partially masked by model bias in completed structures, as shown in Fig. 3.

An additional potential use of composite iterative-build OMIT maps is as a source of relatively unbiased phasing information. The density in each OMIT region of these composite maps is not biased by the model within that OMIT region. Consequently it seems possible that the phase information obtained by using these maps as a target for density modification (Terwilliger, 2001) might be of high quality. It might also have lowered model bias compared to that obtained directly from a model. It would not necessarily be completely unbiased, however, because the inverse Fourier transformation would combine phase information from different OMIT regions and the separation of model information within each OMIT region from the map in that region would therefore no longer be complete. Preliminary experiments indicate that these maps can be improved over standard density-modified maps.

We conclude by noting that increasing use of automated procedures for iterative model-building, density modification and refinement (Perrakis et al., 1999; Terwilliger, 2003; DePristo et al., 2005; Ondrácík, 2005) has the potential for reducing the effects of model bias and the incidence of significant errors to very low levels, particularly for experimentally-phased structures, because a complete check of the fit of model to electron density map can be easily carried out during every cycle of automated building. If poorly-fitted parts of a model are removed promptly, instead of remaining for many cycles of refinement in which the remainder of the model adjusts to compensate for errors in building, then subsequent electron density maps will be unlikely to have substantial model bias.
Figure 1  Iterative build OMIT and composite iterative-build OMIT maps for molecular replacement solution of 1HP7 (Kim et al., 2001). Maps are contoured at 1\(\sigma\). A. OMIT map calculated with \(\sigma_A\)-weighted \((2mFo-DFc) e^{i\phi_c}\) coefficients (Read, 1986) after refinement of molecular replacement model, omitting all atoms in one OMIT region. The atoms in the structure that were not omitted are shown. B. Iterative build OMIT map for the same region shown in A, after 10 cycles of iterative model-building, density modification and refinement. Shown is the model that was built outside of the OMIT region. C. Composite iterative-build OMIT map constructed by combining all OMIT regions obtained as in B. The model is the same as shown in B. D. Composite iterative-build OMIT map as in C with refined structure of 1AS4 superimposed. E. Composite iterative-build OMIT map shown in C and D with MR starting model superimposed. F. Standard iterative-build density-modified map and model built starting from MR starting model after removing all the atoms that are omitted in panel A.

Figure 2  RESOLVE density modified and composite iterative-build OMIT maps for SAD experimental phasing solution of 1VQB (Skinner et al., 1994). Maps are contoured at 1.5\(\sigma\). A. RESOLVE density-modified SAD-phased map (Terwilliger, 2000). B. Iterative build OMIT map for the same region shown in A. The model shown is the refined structure of 1VQB.

Figure 3  \(\sigma_A\)-weighted \((2mFo-DFc) e^{i\phi_c}\) and OMIT maps for 1ZEN (Cooper et al., 1996) compared with structures 1ZEN and of chain A from 1B57 (Hall et al., 1999) superimposed on the structure from 1ZEN. Maps in A-M are contoured at 1\(\sigma\). A, B. \(\sigma_A\)-weighted \((2mFo-DFc) e^{i\phi_c}\) map (Read, 1986) calculated after refinement of the 1ZEN structure with phenix.refine (Afonine et al., 2005b), compared with structure of 1ZEN (A) and with chain A from 1B57 (B). C, D, as in A and B, except that the atoms in the 1ZEN structure were moved randomly by an rms distance of 1 Å (“shake” procedure), then refined for 6 cycles with phenix.refine. E, F, as in C and D, except that solvent water molecules were removed after the shake procedure and 10 cycles of refinement including simulated annealing were carried out. G, H, as in A and B, except the map shown is a simple OMIT map calculated by omitting all atoms in an OMIT box with edges parallel to the cell edges and 4 Å from any atom in residues 5-9 of 1ZEN (setting their occupancies to zero), refining the resulting structure, and calculating a \(\sigma_A\)-weighted \((2mFo-DFc) e^{i\phi_c}\) map. I, J, SA-OMIT map calculated as in G, H except the map is a simulated-annealing OMIT map in which the refinement step in C and D is replaced by simulated-annealing refinement (Brünger et al., 1998). K, L, Iterative build OMIT map calculated as in G, H except that the 1ZEN structure was iteratively rebuilt using the rebuild-in-place option of the PHENIX AutoBuild Wizard, always setting the occupancies of all atoms in the OMIT box to zero during the procedure. M, N. Maps downloaded from the EDS density server (Kleywegt et al., 2004). M, \(\sigma_A\)-weighted \((2mFo-DFc) e^{i\phi_c}\) map and model for 1ZEN. N, \(\sigma_A\)-weighted \((mFo-DFc) e^{i\phi_c}\) map for 1ZEN, contoured at \(\pm 2\sigma\), with coordinates of 1ZEN (green) and 1B57 (blue) superimposed.

Table 1  Map correlation coefficients near residues 3-10 of 1ZEN
| Map (all based on 1ZEN structure except as noted) | map correlations* with |
|-----------------------------------------------|-------------------------|
|                                               | 1ZEN model | 1B57 model |
| Initial $\sigma_A$ map, no OMIT               | 0.68        | 0.75       |
| $\sigma_A$ map after “shake” procedure, no OMIT | 0.67        | 0.74       |
| $\sigma_A$ map after “shake”, removal of waters, refinement and water picking, no OMIT | 0.64        | 0.75       |
| Simple refined OMIT                            | 0.63        | 0.71       |
| Simulated-annealing OMIT                       | 0.60        | 0.71       |
| Iterative-build OMIT                           | 0.65        | 0.75       |
| 1B57 Fcalc map                                 | 0.66        | 0.98       |

*Map correlations were calculated with RESOLVE (Terwilliger, 2000), including grid points within 2 Å of each atom in the corresponding model. Residues 3-10 from 1ZEN were chosen because they were largely within the OMIT region, and residues 3-10 from 1B57 were selected to match the 1ZEN fragment.
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Fig. 3F
