

## 19. OTHER EXPERIMENTAL TECHNIQUES

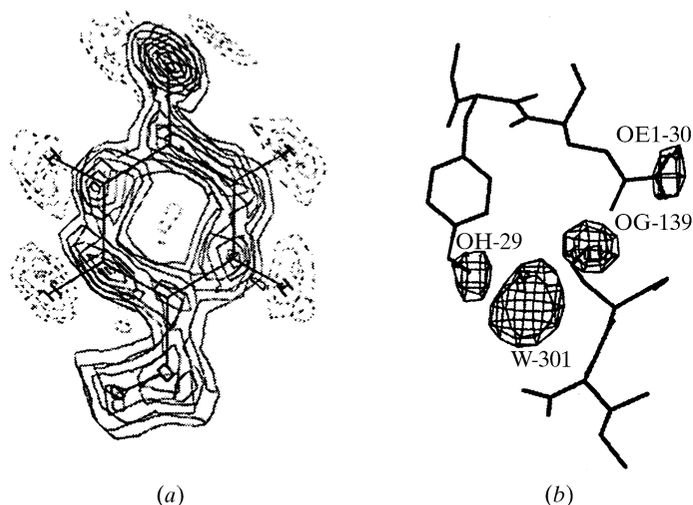


Fig. 19.1.3.1. Information content in neutron density maps. (a) A well ordered tyrosine ring in the 1.4 Å refined structure of crambin (Teeter & Kossiakoff, 1984). (b)  $D_2O - H_2O$  difference density map of a hydrogen-bonding network in trypsin: Gln30 O $\epsilon$ 1–Ser139 O $\gamma$ , Ser139 O $\gamma$ –W301, W301–Tyr29 DO. Water density and H/D exchange density shown.

illusion that the peak has been translated. The hydroxyl deuterium orientation is readily determined by its position in positive density.

Use of  $D_2O - H_2O$  neutron difference maps provides a high level of stereochemical information (see below) (Kossiakoff *et al.*, 1992; Shpungin & Kossiakoff, 1986). Fig. 19.1.3.1(b) displays a network of three hydrogen bonds involving three side-chain types and an occluded water. With knowledge of the heavy atoms alone, it is not possible to define the donor/acceptor character of any of the side chains, because they can act in either capacity, as can the water. The assignments can be made unambiguously from the  $D_2O - H_2O$  density, as can the orientation of the water molecule. These maps have allowed detailed analysis of hydroxyl orientations in protein molecules (Kossiakoff *et al.*, 1990; McDowell & Kossiakoff, 1995).

Neutron diffraction is an ideal method for investigating methyl-group conformation, because it allows direct observation of hydrogen-atom positions (Fig. 19.1.3.2) (Kossiakoff & Shteyn, 1984). Although methyl groups in proteins are not held in fixed positions, but spin rapidly around their rotor axes, the time-averaged character of the diffraction experiment establishes the low-energy conformer and the degree of disorder. Accurate methyl-group analysis requires relatively higher resolution (1.5 Å or better) than characterizing other structural features.

#### 19.1.4. Phasing models and evaluation of correctness

Neutron diffraction does not lend itself to the multiple isomorphous phasing approach. This is because the range in atomic scattering power is much narrower than for the X-ray case. There are a few relatively rare isotopes where a significant anomalous effect exists; however, they are not adequate for getting primary phasing information (Schoenborn, 1975). In practice, the initial phasing model has to be derived from the X-ray-determined structure. This is done by applying the appropriate neutron scattering lengths to the refined X-ray coordinates (Norvell & Schoenborn, 1976). Thus, at least in the early stages of analysis, the neutron model relies heavily on the accuracy of the X-ray structure. The importance of an accurate phasing model is borne out by the fact that in several investigations the phasing models were not accurate enough to allow the structure to be refined successfully.

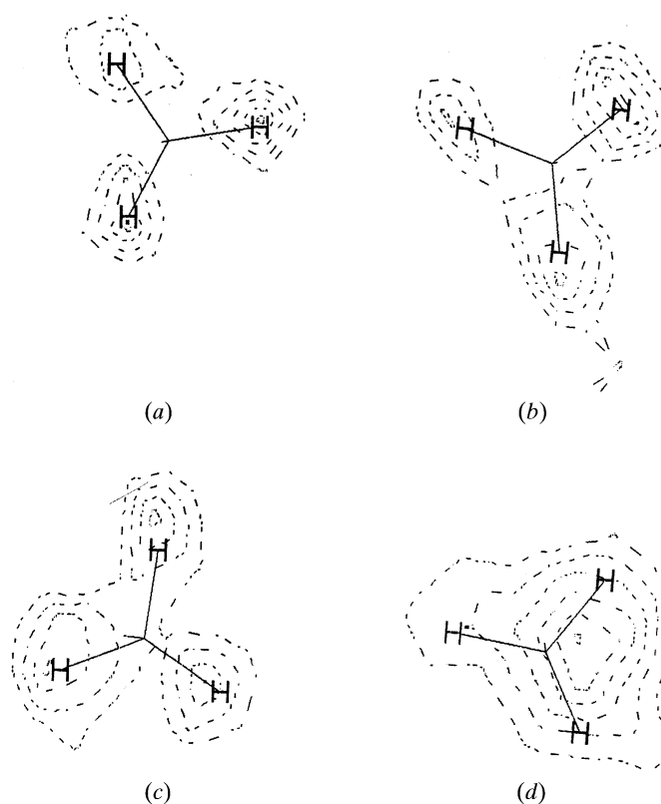


Fig. 19.1.3.2. Sections of a neutron difference Fourier map showing methyl hydrogen densities for several representative methyl groups. No phasing information about the methyl hydrogens was included in the model; therefore, hydrogens should appear in the difference map at their true positions but at reduced density ( $\sim$  half weight). The groups shown are: (a) Ala24, (b) Thr21, (c) Thr28, and (d) Ala45.

#### 19.1.5. Evaluation of correctness

It is an important first step in the structural analysis to determine the quality of the phases derived from the X-ray structure (Kossiakoff, 1983). Several methods have been used. Using the initial phasing model, the most powerful tests examine an unbiased neutron Fourier map for the appearance of features that are independent of the model. The presence or absence of these features, especially those resulting from the scattering of hydrogen and deuterium atoms, is the most reliable measure of the phasing model. One such test is to evaluate the appearance of the water structure, *i.e.*, the water molecules hydrogen-bonded to the surface of the protein. The water molecules observed in the X-ray analysis are excluded from the neutron-phasing model. The test is applied in cases where the crystals have been soaked in  $D_2O$ . The peaks in the neutron density map that correspond to the strongly coordinated water-molecule positions owe their existence solely to the neutron data and phasing model. Even at an early stage, because of the large neutron-scattering potential of  $D_2O$ , many of these tightly bound waters found in the X-ray structures should also be observable in the neutron density map.

Another aspect to test phasing reliability is the ability to identify the orientation of side-chain amide groups of asparagine and glutamine. The difference in neutron scattering between O and the two deuteriums and the N $\delta$ 2 (5.8 f versus 22.6 f) is large enough to be detectable in the Fourier map when these groups are well ordered (Fig. 19.1.5.1). The use of unexchangeable hydrogens for evaluation is considerably more complicated, despite the fact that they constitute about one-half the total number of atoms in the molecule. The difficulty arises from the negative scattering character of the hydrogens, which displaces their apparent positions

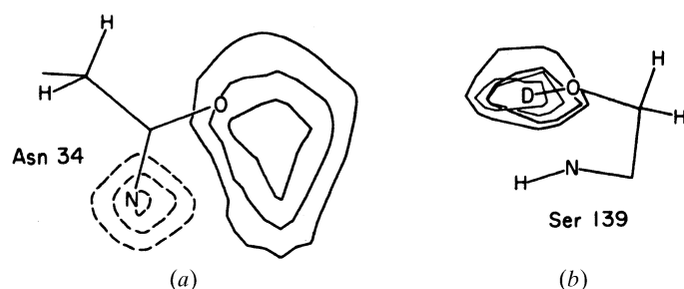


Fig. 19.1.5.1. Difference map of Asn34 in the trypsin structure. (a) In a protein X-ray analysis, the difference in scattering intensity between O and NH<sub>2</sub> is much too small to be detected. In contrast, the neutron-scattering magnitudes of oxygen and nitrogen (5.8 f versus 9.4 f) are quite dissimilar, and there is additional scattering at the nitrogen site from the two bound deuterium atoms. The resulting differential is over 350%, quite large enough to be detected for well ordered side chains. The nitrogen and oxygen positions shown are from the X-ray model. The difference density indicates that the orientation of the nitrogen and oxygen atoms is incorrect and should be rotated by 180° around the C<sub>β</sub>-C<sub>γ</sub> bond. (b) Difference map for Ser139. On well ordered hydroxyl side chains, the orientation of deuterium atoms can sometimes be assigned.

in the Fourier map from the true positions and, coupled with their short bond lengths, complicates the interpretation of the results. Additionally, it has been shown that small errors in positional and thermal parameters of the parent atoms can further complicate the identification of hydrogen-atom positions (Kossiakoff & Spencer, 1981).

### 19.1.6. Refinement

The methodologies employed to refine neutron data are essentially the same as those used in most X-ray studies. These include real-space (Hanson & Schoenborn, 1981; Norvell & Schoenborn, 1976; Schoenborn & Diamond, 1976), reciprocal-space (Bentley & Mason, 1980; Phillips, 1984; Wlodawer & Hendrickson, 1982; Wlodawer & Sjolín, 1981) and restrained difference-map refinement (Kossiakoff & Spencer, 1980; 1981). A joint refinement technique in which the neutron and X-ray data are refined simultaneously has been developed (Wlodawer & Hendrickson, 1982). In addition to the normal difficulties encountered in the refinement of any protein structure, there are several that are peculiar to the neutron-diffraction technique. These special problems arise from the close proximity of hydrogen atoms to their parent atoms, coupled with the effects of the negative scattering length of the hydrogen atoms. Potential problems exist when the difference density generated from positional errors of one atom overlaps an adjacent atom site. The situation is further complicated by the fact that, because of its negative scattering length, an error in a hydrogen-atom position is minimized by moving the atom down the gradient, that is, in the opposite direction to that required for correcting parent-atom positions. To evaluate the extent of this problem in refinement, a test was devised using a 2.2 Å data set (Kossiakoff & Spencer, 1981). The coordinates of the protein trypsin were perturbed by a varying, but known, amount from their ideal positions. It was determined that, in general, convergence towards the true coordinate could be obtained when the coordinate errors were less than 0.3 Å; however, if the parent atom (an atom with one or more hydrogens attached to it) was displaced by more than 0.6 Å from its correct position, the effect of neighbouring hydrogens rendered the calculated shifts inaccurate. The results of this study support the observations of other investigators that it is absolutely crucial that the starting phasing

model be of high quality, because the range of convergence of a neutron analysis is relatively small.

### 19.1.7. D<sub>2</sub>O – H<sub>2</sub>O solvent difference maps

D<sub>2</sub>O – H<sub>2</sub>O solvent difference maps provide an unbiased method for identifying water molecules and exchangeable hydrogens (Kossiakoff *et al.*, 1992). For several years, the large difference in the scattering characteristics of neutrons by H<sub>2</sub>O compared to D<sub>2</sub>O has been effectively exploited by using density matching and exchange labelling in small-angle neutron-scattering experiments. This difference can likewise be exploited in neutron protein crystallography to determine the detailed structural characteristics of protein hydration through the calculation of solvent difference maps (Shpungin & Kossiakoff, 1986; Kossiakoff *et al.*, 1992). In practice, such maps are obtained by comparing the changes in diffracted intensities between two sets of data – one obtained from a crystal having H<sub>2</sub>O as the major solvent constituent, and a second where D<sub>2</sub>O is the solvent medium. To a good approximation, the protein-atom contributions to the scattering intensities in both data sets are equal and cancel, but since H<sub>2</sub>O and D<sub>2</sub>O have very different scattering properties, their differences are accentuated to reveal an accurate and nearly unbiased representation of the solvent structure.

The features of a solvent difference map of this type are not as affected by errors in the phasing model as conventional difference Fourier maps. In addition, there are refinement procedures that can be applied to them that lead to significant enhancement in signal/noise discrimination. The basic feature of the method is a set of density-modification steps based on the fact that a considerable amount of information about the density distribution of the crystallographic unit cell is known. For instance, it is known that

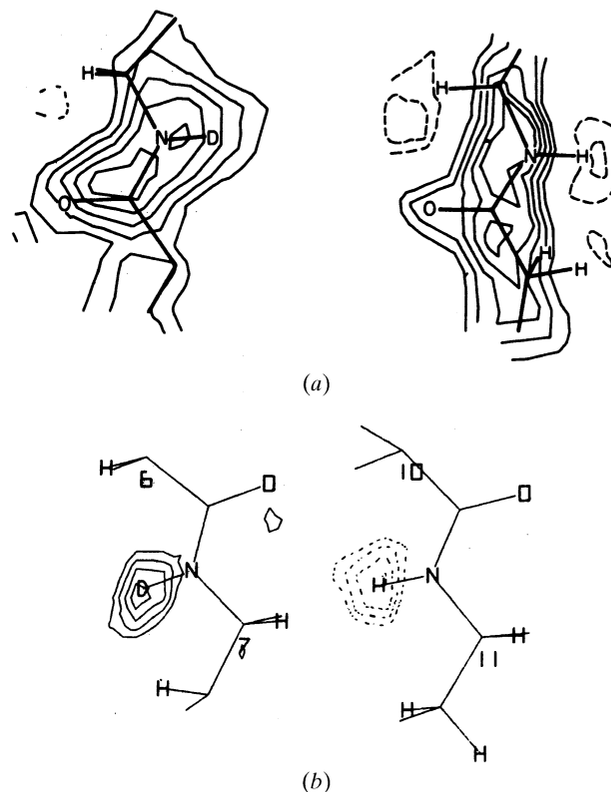


Fig. 19.1.8.1. Sections of neutron density maps taken in the plane of the peptide group. (a)  $2F_o - F_c$  maps showing an example of an exchanged and unexchanged amide peptide group. (b) D<sub>2</sub>O – H<sub>2</sub>O difference density map showing the same.