17.2.11 A RANDOM APPROACH TO CRYSTAL STRUCTURE DETERMINATION. By J.P. Neclarco and G. Germain, Laboratoire de Chimie Physique et de Cristallographie, Université de Louvain, Louvain-la-Neuve, Belgium, and M.W Woolfson and A. Knight, Department of Physics, University of York, Heslington, York Y01 5DD, U.K.

It has been found (Baggio, Woolfson, Neclarco, Germain, Acta Cryst. (1978). A34, 863) that triple-phase relationships treated as linear equations can be used to refine a set of initially random phases, the 70-100 refined phases being used as a starting point for development by the tangent formula. The least-squares method used in refinement is subject to difficulties in weighting the equations and problems arising from the singularity of the matrix can occur. Investigations of alternative methods of refinement show that the method of steepest descents is successful in permitting these difficulties while retaining the advantages of the large starting set method. Novel approaches are made possible by the easier application of weighting schemes and the process is computation-ally economical. Illustrations are shown using output generated by the distributed program YZARC which optionally uses either method of refinement.

17.2.12 RANTAN - RANDOM MULTAN.
By Yao Jia-xing, Department of Physics, University of York, Y01 5DD, England.

RANTAN, a random approach to crystal structure determination, is described. A large number of phases are given random values associated with low weights and then refined, together with reflections fixing origin and enantiomorph, by a weighted tangent formula. An initial random phase is not changed until a phase estimate is obtained with a new weight greater than the initial weight. Then the phase is allowed to vary and to follow its refinement path. In the present method it is possible simultaneously to work with all the reflections and all the relationships ab initio. More than 20 structures, which represent a wide variety of space groups, structural complexity and difficulty, have been successfully solved by RANTAN. One example is a difficult unknown structure containing 100 atoms in the asymmetric unit with space group P21. The first E-map, with very good figures of merit, showed 85 atoms. The experimental results have shown that it is possible to use RANTAN to determine more complex structures which contain more than 100 atoms in the asymmetric unit.

Another application of RANTAN is its use with a partial structure, like Karle recycling. The known phases coming from a partial structure are combined with random phases and then RANTAN is run as usual. Normally only 10% of a structure is enough to develop it completely.

17.2.13 MAGEX: A PROCEDURE FOR PHASE DETERMINATION. By S.E. Hull, D.Viterbo, M.H. Woolfson and Zhang Shao-Hui, Department of Physics, University of York, Heslington, York Y01 5DD, U.K.

This procedure extends and strengthens the PS method (Declarco, Germain and Woolfson, Acta Cryst. (1979)). Long one-dimensional magic-integer sequences are used to express the phases of from 5 to 20 primary reflections. A new concept of multiple definition of secondaries is employed and the error involved in their subsequent application in a conventional g map is thus much reduced. A parameter-shift process, which is based on one or other of two functions, is used to refine 200 sets of from 40 to 100 phases and a number of the best sets of refined phases is selected. A facility is available for limited multi-pathway phase extension before large-scale phase extension by the controlled use of the tangent formula is undertaken. Examples of successful applications of MAGEX are described.

17.2.14 STRENGTHENED TRANSLATION FUNCTIONS: FINDING THE POSITION OF A SMALL, KNOWN FRAGMENT BY SEARCH METHODS IN DIRECT-FOURIER SPACE. By H.M. Doesburg and P.T. Beurskens, Crystallography Laboratory, Twente University, 5525 ED Hooijmag, The Netherlands.

A frequently occurring problem in the determination of crystal structures is the positioning of a correctly oriented molecular fragment, relative to the symmetry elements or relative to another known molecular fragment. Knowledge of the orientation of a fragment may be available from:

- Patterson rotation search techniques.
- Direct methods, when a recognizable fragment is found from an otherwise obscure E map.


Definitions: \( P_{o_f} \) is the electron density of the known fragment and \( P_{o_h} \) are the calculated partial structure factors. \( P_{o} \) is the electron density of the unknown part of the structure and \( P_{o_h} \) are the corresponding structure factors obtained by DIRIDIF, using the known fragment as input. \( P_{o_f} \) is a symmetry related image of \( P_{o} \).

The translation function is defined as

\[
Q(g) = \sum_{h} \frac{1}{\exp(-2\pi i h \cdot g)} P_{o_f}(h) P_{o_h}(h)
\]

From convolution theory it follows:

\[
Q(g) = \sum_{h} \frac{1}{\exp(-2\pi i h \cdot g)} P_{o}(h) P_{o_h}(h)
\]

The maximum of the Q-function gives the position of \( P_{o_f} \) relative to \( P_{o} \). From this the position of the symmetry element can easily be deduced.
For instance, in space group $P2_1$, the $Q$ function takes the form

$$Q(q') = \frac{1}{V} \sum \int \exp\left(2\pi i \mathbf{q} \cdot \mathbf{r}ight) F_{\text{ref}}(\mathbf{g}) (\mathbf{q} - \mathbf{q}') \exp(2\pi i \mathbf{q} \cdot \mathbf{g})$$

while only one Fourier section ($q = x, y, z$) has to be calculated. The shift vector is $-\mathbf{q}'$, where $\mathbf{q}'$ is the position of the highest peak in the $Q$-function.

The shift can be applied in reciprocal space, to produce Fourier coefficients for a new symmetry-enforced electron density map. Examples in various space groups, simultaneous location of the fragment relative to all symmetry elements, and location of fragments relative to each other, will be shown.

Advantages: automatic, high-speed computer program; no Patterson-overlap; for large structures: multisolution technique and application to small or qualitatively bad fragments is possible.

Flow diagram

known

\[ \text{b = reflection data (expand to triclinic)} \]

\[ \text{calc. struct. factors} \]

\[ \text{symm.} \]

\[ \text{DIERDIF = direct method refinement of } \Delta F \]

\[ \text{incl. enantiom. fixation} \]

\[ \text{DIERDIF} \]

\[ \text{shift vector} \]

\[ \text{DIERDIF} \]

\[ \text{to find the rest structure} \]

17.2.15 BREAKING OF PSEUDOSYMMETRY USING DIRECT METHODS. By R. Döhne, Institut für Angewandte Physik, Lehrstuhl für Kristallographie, Universität Erlangen-Nürnberg, Germany (BRD).

Let $p(x)$ describe the electron density distribution of a structure and $F(h)$ its structure factors. If $p(x)$ satisfies a pseudotranslation $\xi$ of index $p$ it is useful to divide the set of all reflections into the set of main and the set of superstructure reflections. The main reflections $H(h, \xi)$ determine the structure of the cell $H = \frac{1}{p} \mathbf{P} \mathbf{P}_0^{-1}$ and the superstructure reflections $U(h, \xi)$ not integer determine the complement structure

$$\tilde{p}(x) = p(x) - \tilde{p}(x) = \frac{1}{V} \sum H(h, \xi) \exp(2\pi i \mathbf{H} \cdot \mathbf{x})$$

(M. Buergers, Vector space 1959). Because $\tilde{p}(x)$ contains equal amounts of positive and negative electron density (J.W. Jeffery 1964, Acta Cryst. 17, 778) this concept is not useful in direct methods. But there exists a structure $\tilde{p}(x)$ which has positive maxima only and the structure factors $\tilde{F}(h)$ of $\tilde{p}(x)$ coincide with $F(h)$, whereas the $\tilde{F}(h)$ remain unknown. If the pseudotranslation has index $p=2$, the maxima in $\tilde{p}(x)$ cannot be rationally dependent of each other, so that $F_{\text{abs}}(u) = \tilde{F}_{\text{abs}}(u)$. Therefore the $E$-normalisation is correct also if the $\tilde{F}(h)$ are known only. Because of $\tilde{E}(h) = E(h)$ the phase determination of $\tilde{E}(h)$ gives phases of $E(h)$ and $\tilde{F}(h)$. If the index $p$ is larger than 2, $\tilde{p}(x)$ normally shows less rational dependency among the coordinates of maxima than $p(x)$. So the assumptions for using direct methods are better fulfilled. In these cases ($p > 3$) it can be shown that the sums of phases of triplets with high values of $x$ are more reliably $\pi$ than 0 under specific conditions. So phase determination is reduced to normalisation of substructure and complement structure. If $n$ atoms having formfactors $f_1, \ldots, f_n$ are expected in the unit cell of $p(x)$, $F$-values can be calculated if the expected intensity $F_{\text{abs}}^2(U)$ and $F_{\text{abs}}^2(U)$ is approximated by

$$<F_{\text{abs}}^2> = a \exp(-B \sin^2 \theta / \lambda^2) \sum_{j=1}^{n} s_j^2$$

using different values of $a$ and $B$ for main and superstructure reflections. This can be done by normalizing $F(U)$ and $F(U)$ separately using a standard $E$-normalising program. If some atoms of the structure satisfy $\xi$ and the others do not this normalisation is equivalent to rescaling the $E$-values. It can be shown by theory and by example (Eukrypt. Tscherry, Schulz & Laves (1972), 2, Krist. 135, 175-182) that in the case where the pseudotranslation $\xi$ is nearly but not exactly fulfilled,

$$<F_{\text{abs}}(U)> \text{ versus } \sin \theta / \lambda \text{ is rather different from the expected intensity of a structure with independent atoms.}$$

The proposed procedure can also be applied to pseudocentrosymmetric structures in handling real and imaginary part of structure factors separately. But in this case it is necessary to determine the coordinates of maxima of the centrosymmetric subcell.

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17.2.16 PROBLEMS OF ENANTIONORM DISCRIMINATION IN DIRECT METHODS. By Suzanne Fortier, William L. Duax and Herbert Hauptman, Medical Foundation of Buffalo, Inc., 73 High St., Buffalo, NY 14203, U.S.A.

The problem of enantiomorph definition in space groups $P1$, $P2_1$, $C2$ and $C3$ is often wrongly attributed to the presumption that it is not possible, in these space groups, to select a starting reflection with a phase orthogonal to the origin defining set.

In simple structures, that is in structures with a well behaved set of triples (with no unexpectedly large deviation from zero) and no significant character of pseudo-centrosymmetry, enantiomorph sensitive phases are automatically identified by a convergence type procedure and are left out as part of the basis set of phases. Moreover, inspection of the phases in the basis set, in particular of their interrelationship in terms of accessibility-inaccessibility (linear-rational dependence) often permits one to single out the best candidate for enantiomorph discrimination.

When the distributions of the normalized structure factors do not suggest any significant degree of pseudo-centrosymmetric character, failure to properly define the enantiomorph is symptomatic of the occurrence of aberrant triplets at important links of the phase development. The possible "ways out", i.e. clean-up techniques (editing the triplets to weed out the aberrant ones), enantiomorph sensitive invariants and seminvariants and the method of strong enantiomorph discrimination, will be discussed.

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