Acta Crystallographica Section D
Biological Crystallography

ISSN 0907-4449

Ryan H. Lilien, ${ }^{\text {a,b,c }}$ Chris BaileyKellogg, ${ }^{\text {d }}$ Amy C. Anderson ${ }^{\text {c, }{ }^{*} \text { * }}$ and Bruce R. Donald ${ }^{\text {a }, c, e, f}$,
${ }^{\text {a }}$ Dartmouth Computer Science Department, Hanover, NH 03755, USA, ${ }^{\text {b }}$ Dartmouth Medical School, Hanover, NH 03755, USA, ${ }^{\text {c }}$ Dartmouth Center for Structural Biology and Computational Chemistry, Hanover, NH 03755, USA, ${ }^{\text {d Purdue }}$ Department of Computer Sciences, West Lafayette, IN 47907, USA, ${ }^{\text {e }}$ Dartmouth
Department of Chemistry, Hanover, NH 03755, USA, and 'Dartmouth Department of Biological Sciences, Hanover, NH 03755, USA

Correspondence e-mail:
amy.c.anderson@dartmouth.edu,
bruce.r.donald@dartmouth.edu

## A subgroup algorithm to identify cross-rotation peaks consistent with non-crystallographic symmetry

Molecular replacement (MR) often plays a prominent role in determining initial phase angles for structure determination by X-ray crystallography. In this paper, an efficient quaternion-based algorithm is presented for analyzing peaks from a cross-rotation function in order to identify model orientations consistent with proper non-crystallographic symmetry (NCS) and to generate proper NCS-consistent orientations missing from the list of cross-rotation peaks. The algorithm, $C R A N S$, analyzes the rotation differences between each pair of cross-rotation peaks to identify finite subgroups. Sets of rotation differences satisfying the subgroup axioms correspond to orientations compatible with the correct proper NCS. The CRANS algorithm was first tested using crossrotation peaks computed from structure-factor data for three test systems and was then used to assist in the de novo structure determination of dihydrofolate reductase-thymidylate synthase (DHFR-TS) from Cryptosporidium hominis. In every case, the CRANS algorithm runs in seconds to identify orientations consistent with the observed proper NCS and to generate missing orientations not present in the cross-rotation peak list. The CRANS algorithm has application in every molecular-replacement phasing effort with proper NCS.

## 1. Introduction ${ }^{1}$

When the structure of an homologous protein is known, initial phases for the diffraction data can often be determined using the technique of molecular replacement (Rossmann \& Blow, 1962; Crowther \& Blow, 1967; Rossmann, 1990). To use initial phases from an homologous model, each copy of the molecular-replacement model (henceforth called the model for brevity) must be properly oriented and translated within the asymmetric unit (Tong \& Rossmann, 1990; Rossmann \& Blow, 1962; Crowther \& Blow, 1967; Rossmann, 1990). The task of properly orienting and translating each model is facilitated by exploiting the additional constraint provided by proper non-crystallographic symmetry (NCS).

Traditionally, the initial presence and degree (i.e. threefold, fourfold etc.) of NCS is first identified by a self-rotation function (Rossmann \& Blow, 1962), while the orientations of each copy of the model are subsequently identified using the cross-rotation function (Rossmann \& Blow, 1962). Ideally, after $n$-fold NCS has been identified, the peaks of the crossrotation function will possess two desirable properties. First,

[^0]Received 5 December 2003 Accepted 24 March 2004
© 2004 International Union of Crystallography Printed in Denmark - all rights reserved
the top $n$ cross-rotation peaks should have rotation-function scores significantly higher than the rest and, second, these top peaks should correspond to the correct NCS-consistent orientations. A complication arises when the model is partial (Oh, 1995), when the model shares only moderate structural similarity with the crystallized molecule or when the degree of NCS is high and each model represents only a small fraction of the unit cell's molecular mass. Empirically, in these situations the $n$ top-scoring rotations from a cross-rotation function search may not correspond to the $n$ correct (NCS-consistent) rotations. In fact, some of the correct rotations may not even appear in the cross-rotation peak list, further complicating the search for the correct model orientations. For complex systems it is therefore possible that the wrong cross-rotation peaks are selected for use with the computationally expensive translation function. We have developed an algorithm to (i) compute which cross-rotation peaks generate proper NCSconsistent model orientations and (ii) generate proper NCSconsistent model orientations not specified by rotations in the cross-rotation peak list.

The CRANS (cross-rotation analysis) algorithm identifies and computes sets of rotations (set $R$ in Fig. 1) that, when applied to the model, produce orientations consistent with the NCS. We call these sets of rotations NCS-consistent rotation sets. CRANS therefore reduces the time required to obtain initial phases in two ways. First, by correctly identifying NCSconsistent rotation sets among the rotations identified by the cross-rotation function, we reduce the number of improperly oriented models and thereby the total number of translation searches required to generate initial phases. Second, in the case where one or more NCS-consistent rotations are absent from the cross-rotation peak list, CRANS (i) informs the

crystallographer that a peak is missing without requiring them to perform translation searches on each cross-rotation peak and (ii) computes the missing NCS-consistent rotations.

The core of the CRANS algorithm analyzes the set of rotation differences to identify finite sets of rotations that satisfy a point group with a single rotation axis. Given a set of $w$ cross-rotation peaks (rotations) $C=\left\{r_{1}, r_{2}, \ldots, r_{w}\right\}$, where $r_{i}$ is an element of the group of three-dimensional rotations, CRANS examines the $w^{2}$ rotation differences $d_{i j}$, where $d_{i j}=r_{i}^{-1} r_{j}$ and $r_{i}^{-1}$ is the inverse of $r_{i}$ (such that $r_{i}^{-1} r_{i}$ is the identity). Conceptually, the rotation difference $d_{i j}$ is the rotation that rotates the model oriented by $r_{i}$ into the model oriented by $r_{j}$ (Fig. 1). We can test the NCS-consistency of a set of cross-rotation peaks by examining their rotation differences and verifying that they share a common axis and form a finite subgroup (Appendix $A$ ). In the event of missing rotations, the CRANS algorithm completes each partial set of identified NCS-consistent rotations by generating missing rotations with quaternions (Hamilton, 1969; Salamin, 1979). Although rotation differences can be computed using a number of rotation representations (Evans, 2001), we chose to use quaternions because they have a single compact representation with explicit axis and angle components, are easily composed, are free of singularities, and represent a uniform parameterization of rotation space.

When the model $P$ is a homodimer, the search for NCSconsistent rotation sets is more complex because the orientation $r P$ (where $r P$ is the result of rotating protein $P$ by rotation $r$ ) is equivalent to the orientation $r f P$, where $f$ is the $180^{\circ}$ rotation around the dimer twofold axis (Fig. 2). This rotational degeneracy increases the difficulty of the search because the rotational relationships between the cross-



MR model Identity: $d_{\alpha}=I$ Inverse: $d_{\alpha}=d_{\alpha}^{-1}$ $d_{\beta}=d_{\gamma}^{-1}$
Closure:
$d_{\alpha} d_{\beta}=d_{\beta} d_{\alpha}=d_{\beta}$ $d_{\alpha} d_{\gamma}=d_{\gamma} d_{\alpha}=d_{\gamma}$ $d_{\beta} d_{\gamma}=d_{\gamma} d_{\beta}=d_{\alpha}$

Figure 1
Two-dimensional examples for threefold (a) and fourfold (b) NCS. (a) A model is shown with the results of a simplified cross-rotation search $C$ containing only five rotations. Orientations corresponding to rotations $r_{1}$ (purple), $r_{3}$ (green) and $r_{4}$ (blue) form an NCS-consistent rotation set $R$. For clarity, only a few rotation differences $\left(d_{11}, d_{13}\right.$ and $\left.d_{14}\right)$ are shown in the upper right overlapping orientation figure. The rotation differences $D(R)$ form a complete rotation difference set and satisfy the group properties of associativity (not shown), identity, inverse and closure. (b) A fourfold NCS example is shown using similar notation to $(a)$. In this example, only three of the four NCS-consistent rotations $\left(r_{1}, r_{3}\right.$ and $\left.r_{4}\right)$ are contained in the cross-rotation peak list $C$. The missing rotation $r_{\alpha}=d_{1 \alpha} r_{1}$ is computed using $d_{1 \alpha}$ defined by the axis of the three identified NCS-consistent rotations and the missing angle. The now complete NCS-consistent rotation set $R$ has a complete rotation difference set $D(R)$ which satisfies the subgroup properties.
rotation peaks must be examined modulo $f$. Thus, the dimer axis is explicitly considered by $C R A N S$ when using a homodimer model.

Although the CRANS algorithm only requires a crossrotation peak list as input, if the NCS axis and/or degree are known (i.e. from the self-rotation search) they can be used as an additional constraint in identifying rotation subgroups. In addition to searching for NCS-consistent rotation sets satisfying one particular symmetry, the CRANS algorithm may also be run in 'scan' mode, where the algorithm consecutively attempts to identify subgroups of rotation differences consistent with a user-specified range of NCS degree (i.e. threefold, fourfold, ..., $n$-fold).

Previous work in automated cross-rotation peak analysis includes the programs RFCORR and MOLREP (Vagin \& Teplyakov, 1997; both distributed as part of CCP4; Collaborative Computational Project, Number 4, 1994). RFCORR and MOLREP both compute the set of rotation differences but leave the task of identifying sets of NCS-consistent rotations to the user; CRANS automates this process and identifies NCS-consistent rotation sets even when there are missing cross-rotation peaks.

In summary, the $C R A N S$ algorithm analyzes the output of a cross-rotation search and makes the following contributions:
(i) CRANS computes all sets of $n$-fold NCS-consistent rotation sets, even in the presence of potentially missing peaks. The missing peaks (i.e. those not found by the cross-rotation function) required to form a complete (i.e. no missing rotations) NCS-consistent rotation set are also computed by CRANS. The computed missing peaks can be used to orient the model prior to performing a translation search.
(ii) During a traditional manual examination of crossrotation peaks, one typically only examines the top $p$ crossrotation peaks. In contrast, the CRANS algorithm performs an exhaustive search over all peaks in the cross-rotation function, checking each rotation for inclusion into an NCS-consistent rotation set.
(iii) CRANS computes the NCS axis for each identified NCS-consistent rotation set. If the NCS axis is known from the self-rotation map, CRANS can limit its search to return only those rotation sets with NCS axes close to the known NCS axis.
(iv) CRANS can also be run in 'scan' mode, where the algorithm sequentially searches a range of degrees of NCS and attempts to identify NCS-consistent rotation sets for each degree of symmetry.
(v) Finally, CRANS can perform the four above listed tasks when the model is a homodimer. In our experience, the ability to handle homodimer models was vital in solving the structure of dihydrofolate reductase-thymidylate synthase (DHFR-TS) from Cryptosporidium hominis (O’Neil et al., 2003; §3.4).

The paper is organized as follows. In $\S 2$ we describe the CRANS algorithm, present the algorithm's runtime complexity and describe the data and pre-processing used for each test system. In $\S 3$ and $\S 4$ the performance of $C R A N S$ is demonstrated by analyzing six cross-rotation peak lists generated from six models against four protein crystal systems.

Firstly, we describe the performance of the CRANS algorithm on three test systems of solved crystal structures containing threefold (2-keto-3-deoxy-6-phosphogluconate aldolase; PDB code 1fq0; Wymer et al., 2001), fivefold (cholera toxin B subunit; PDB code 1chp; Merritt et al., 1995) and sevenfold (Gp31 co-chaperonin; PDB code 1g31; Hunt et al., 1997) NCS. Then, in $\S 3.4$ we describe how we used CRANS to solve the de novo structure of dihydrofolate reductase-thymidylate synthase (DHFR-TS) from C. hominis (ChDHFR-TS; PDB code 1qzf; O'Neil et al., 2003), a homodimer, where we used $C R A N S$ to identify the cross-rotation peaks consistent with fivefold NCS. Despite the fact that three of the six tested crossrotation peak lists were missing at least one and in one instance as many as three NCS-consistent rotations, in all cases $C R A N S$ was able to correctly identify NCS-consistent rotations and generate all missing rotations. These results support the general applicability of the $C R A N S$ algorithm for use in analyzing cross-rotation search results for systems with NCS. In $\S 5$ we discuss the use of $C R A N S$ in crystals with improper NCS, multiple point-group symmetries and parallel crystallographic and non-crystallographic axes.

## 2. Methods

### 2.1. CRANS algorithm

The CRANS algorithm takes a list of cross-rotation peaks as input and identifies sets of cross-rotation peaks that generate orientations of the model related by $n$-fold NCS (Appendix $B)$. In describing the $C R A N S$ algorithm, we define the func-


Figure 2
Homodimer-based rotations. The model in orientation $A$ can be rotated by either $r_{1}$ or $r_{2}$ to assume orientations $B$ or $C$. While both orientations $B$ and $C$ can form a fourfold NCS (modulo the dimer flip) with orientations $X, Y$ and $Z$, the rotations $d_{\alpha}$ and $d_{\beta}$ are not equal. The rotation difference $d_{\alpha}$ is not consistent (i.e. does not form a subgroup) with the partial rotation difference set generated from orientations $X, Y$ and $Z$. Thus, it becomes important that when the model is a homodimer, for every rotation $r_{1}$ we generate the dimer-flip-related rotation $r_{2}=r_{1} f$ (where the rotation $f$ flips the model along its twofold dimer axis).
tions axis(•) and angle (•) to return the axis and angle, respectively, of a given rotation. We define the $n$ angles specified by $360 t / n(\mathrm{t} \in\{0,1, \ldots, n-1\})$ as symmetry angles. Let $m$ be the maximum number of missing rotations allowed by the user.

The CRANS algorithm is divided into four stages: process input, filter, partition and patch. In the process-input step, all $w$ original rotations (rotations from the cross-rotation function) $C=\left\{r_{1}, r_{2}, \ldots, r_{w}\right\}$ are read from an input file and converted to quaternions. The CRANS input format is described in Appendix B. After reading the cross-rotation peak-list input file, CRANS computes all $w^{2}$ rotation differences, $d_{i j}$.

In the filter stage, all rotation differences with angle $\left(d_{i j}\right)$ differing from a symmetry angle by more than the user-defined angular threshold $\tau_{\text {angle }}$ are discarded. The intuition behind this filtering step is that two rotations $r_{i}$ and $r_{j}$ that are both members of the correct NCS-consistent rotation set should have a rotation difference angle $\left(d_{i j}\right)$ approximately equal to a symmetry angle.

The core of the algorithm occurs in the partition stage. In general, if rotation differences $d_{i j}$ and $d_{k l}$ are both members of the same NCS-consistent rotation set, then $\operatorname{axis}\left(d_{i j}\right) \approx \operatorname{axis}\left(d_{k l}\right)$. Therefore, in this stage, each remaining (i.e. unfiltered by step 2) rotation difference $d_{i j}$ is clustered with all rotation differences $d_{i k}$ that share a common rotation axis. That is, the differences $d_{i j}$ and $d_{i k}$ are assigned to the same set if $\left|\theta_{i j, i k}\right| \leq \tau_{\text {axis }}$, where $\theta_{i j, i k}\left(0 \leq \theta_{i j, i k} \leq 90^{\circ}\right)$ is the angle between axis $\left(d_{i j}\right)$ and axis $\left(d_{i k}\right)$ and $\tau_{\text {axis }}$ is the user-defined axis similarity threshold. Sets of rotation differences with more than $m$ missing distinct rotations are eliminated. In every set $S_{i}$ of rotation differences of the form $d_{i j}$ (where $i$ is fixed and $j$ varies, i.e. all rotation differences involve rotation $r_{i}$ ) the number of distinct rotations equals the number of unique orientations generated by applying $d_{i j} \in S_{i}$ to $r_{i} P$, where $P$ is the model. Because multiple rotations in a single rotation difference set might generate the same orientation, all possible subsets of rotation differences containing at most $m$ missing distinct rotations are computed. At this point, each set $S_{i}$ of rotation differences has generated zero or more partial rotation difference sets $\Delta^{\prime}$. For each partial rotation difference set $\Delta^{\prime}$, let $D^{-1}\left(\Delta^{\prime}\right)$ be the set of all original cross-rotation peaks used to generate rotations in $\Delta^{\prime}$ and let $\mathbf{v}$ be the average axis of all rotation differences formed by pairs of rotations in $D^{-1}\left(\Delta^{\prime}\right)$. The consistency of the NCS axis for each rotation difference is checked by comparing the axis of rotation difference $d_{a b}$ to the average axis $\mathbf{v}$ for all $r_{a}, r_{b} \in D^{-1}\left(\Delta^{\prime}\right)$ to ensure $\left|\theta_{a b, \mathbf{v}}\right| \leq \tau_{\text {axis }}$, where $\theta_{a b, \mathbf{v}}\left(0 \leq \theta_{a b, \mathbf{v}} \leq 90^{\circ}\right)$ is the angle between $\operatorname{axis}\left(d_{a b}\right)$ and the average axis $\mathbf{v}$. Rotation difference sets not passing this filter are eliminated. After computing all rotation sets, those sets that are subsets of other, more complete, remaining rotation sets are removed.

In the final stage of the algorithm, the patch stage, missing rotations for each remaining rotation set $R$ are computed. For example (Fig. 1b), if $n=4, m=1$ and a rotation set $R$ contains three rotations $r_{1}, r_{3}$ and $r_{4}$ such that $\operatorname{axis}\left(d_{13}\right)=\operatorname{axis}\left(d_{14}\right)=$ $\operatorname{axis}\left(d_{34}\right)$ and angle $\left(d_{13}\right)=90^{\circ}$, angle $\left(d_{14}\right)=270^{\circ}$, angle $\left(d_{34}\right)=$
$180^{\circ}$, then the missing rotation $r_{\alpha}=d_{1 \alpha} r_{1}$ is computed, where $d_{1 \alpha}$ is the quaternion representing a rotation of $180^{\circ}$ (the missing angle in this example) around axis $\left(d_{13}\right)$ (Fig. 1b). By construction, $D(R)$ will be a subgroup (see Appendix $A$ ) and thus $R$ will be an NCS-consistent rotation set.

The NCS-consistent rotation sets identified by CRANS are sorted first on their completeness (i.e. sets with fewer missing rotations are listed first) and second on a quality score. Two quality scores are implemented in CRANS. In the first method, RF-Scoring, the score of each NCS-consistent rotation set is computed as the sum of the rotation-function scores (RF-Scores) of the included cross-rotation peaks. In the second method, axis-angle deviation scoring, the score $s$ of each NCS-consistent rotation set $R$ is computed as the average angular deviation between the axis of each rotation difference and $\mathbf{v}$ plus the average angular deviation between each rotation difference and its closest symmetry angle,

$$
\begin{equation*}
s=\frac{2}{n(n+1)} \sum_{i=1}^{n} \sum_{j=i+1}^{n}\left(\theta_{i j, \mathbf{v}}+\min _{t \in Z_{n}}\left|\operatorname{angle}\left(d_{i j}\right)-360 t / n\right|\right), \tag{1}
\end{equation*}
$$

where $n$ is the number of rotations in $R, \theta_{i j, v}$ is the angle between $\operatorname{axis}\left(d_{i j}\right)$ and the average axis $\mathbf{v}$ and $Z_{n}=\{0,1, \ldots$, $n-1\}$.

When the model is a homodimer, one change is made to the process-input stage of the above algorithm. First, the dimer axis is computed from the model structure and is used to compute $f$, the $180^{\circ}$ rotation around the twofold dimer axis (Fig. 2). The rotation $f$ is then used to generate a dimer-flipped version of each original rotation, which is added to the list of rotations utilized by subsequent stages of the algorithm. Formally, the set of cross-rotation peaks $C$ is replaced by $C \cup\{r f \mid r \in C\}$. The algorithm then proceeds as described above.

Because CRANS does not directly utilize knowledge of crystallographic symmetry, it unable to expand the crossrotation peaks into their crystallographic equivalents. Therefore, NCS-consistent orientations generated by crystallographic symmetry must be identified during the cross-rotation search. The CNS (Brünger et al., 1998) cross-rotation search used in all our examples performs this crystallographic expansion. Therefore, $C R A N S$ can directly analyze the crossrotation search output.

### 2.2. Complexity

Reading, conversion to quaternions and computation of the inverse of each of the $w$ cross-rotation peaks requires $O(w)$ time. Computing all rotation-pair differences requires $O\left(w^{2}\right)$ time. Filtering the rotation-pair differences (for those near the $n$ symmetry angles) requires time $O\left(n w^{2}\right)$. Formation of rotation-difference sets sharing a common axis takes a worstcase time of $O\left(w^{4}\right)$, which occurs when all $w^{2}$ rotation differences have passed the previous symmetry-angle filter. In practice, the number of remaining differences is quite small and only a fraction of the original rotation differences are used to compute rotation-difference sets. Creation and patching of the final sets consistent with the desired NCS requires time
$O(g)$, where $g$ is the number of NCS-consistent rotation sets. Therefore, the CRANS algorithm can search for NCS symmetry among $w$ cross-rotation peaks with an expected runtime of $O\left(n w^{2}+g\right)$ and a worst-case runtime of $O\left(w^{4}+g\right)$. In practice, $g$ is a small constant and we can reduce the runtime to an expected $O\left(n w^{2}\right)$ and a worst-case $O\left(w^{4}\right)$. Our implementation of $C R A N S$ requires only seconds to search for up to eightfold NCS on lists of 120 cross-rotation peaks using an Athlon-based processor.

### 2.3. Source of data and preprocessing

Structure factors for the three test systems 1fq0, 1chp and 1 g 31 were obtained from the Protein Data Bank (PDB; Berman et al., 2000) and converted into CNS (Brünger et al., 1998) format. The first chain from the crystal structure (representing a single monomer) was extracted and used as the model for molecular replacement. A cross-rotation search was performed with $C N S$ using the following default parameters: resolution limits, $15-4 \AA$; data cutoff criteria, 0.0 ; r.m.s. outlier cutoff, 1000; bins for resolution-dependent operations, 10; atoms, known and not hydrogen; scoring function, fastdirect; use automatically determined asymmetric unit; fastdirect grid factor, 5 ; fastdirect maximum number of peaks, 20; cluster threshold, $10^{\circ}$; maximum number of coarse peaks to analyze in fine grid search, 20. The generated cross-rotation function peak list was then analyzed by $C R A N S$. The same axis and angle tolerances $\tau_{\text {axis }}=4.5^{\circ}$ and $\tau_{\text {angle }}=5.0^{\circ}$ were used in all CRANS analyses. Noise in the rotation search will sometimes cause the cross-rotation peaks to deviate from their true NCS-consistent position. Empirically, we found that these values for the axis and angle thresholds allowed multiple NCSconsistent rotation sets to be identified in our test cases; if no NCS-consistent rotation sets are identified but confidence in the presence of NCS is high, the $\tau_{\text {axis }}$ and $\tau_{\text {angle }}$ thresholds may be relaxed.

For comparison, the translation-only RMSD (TRMSD) was computed between the model oriented according to each cross-rotation peak and each chain in the crystal structure. We define the TRMSD of two proteins to be the minimum mainchain $\mathrm{C}^{\alpha}$ RMSD achievable when the molecules are only allowed to translate relative to one another (i.e. rotations are not allowed). Using this measure, we are able to quantify the similarity between models oriented by each cross-rotation peak and each monomer of the crystal structure. We emphasize that the TRMSD is not part of the CRANS algorithm but is rather a tool used to analyze the $C R A N S$ output to verify the correctness of our test cases.

Diffraction data for ChDHFR-TS were collected at Brookhaven National Laboratory (beamline X12C) as previously described (O'Neil et al., 2003). Experiments were performed using three molecular-replacement models. The first model, LmDHFR-TS, consists of the DHFR-TS homodimer of the Leishmania major DHFR-TS protein (Knighton et al., 1994). The second model, PcTSA, consists of the TS homodimer of the Pneumocystis carinii TS protein (PDB code 1f28; Anderson et al., 2001), while the third model, PcTSB, is
simply a rotated version of PcTSA with the flexible loop Asn186-Glu191 removed. Both the PcTSA and PcTSB models only consist of the TS homodimer, which represents approximately $60 \%$ of the entire DHFR-TS homodimer

## 3. Results

The CRANS algorithm was tested on structure-factor data from four different protein crystals exhibiting three-, five- and sevenfold NCS. Three of these systems exhibit planar NCS, whereas ChDHFR-TS exhibits a $5_{1}$ screw NCS (Fig. 3).

### 3.1. 2-Keto-3-deoxy-6-phosphogluconate aldolase (threefold NCS)

Structure factors for 2-keto-3-deoxy-6-phosphogluconate aldolase (1fq0) were obtained from the PDB. The first chain of 1 fq 0 was used as a model in a cross-rotation search using default parameters ( $\S 2.3$ ). The resulting cross-rotation peak list was sorted by rotation-function score and contained 162 rotations. The TRMSD ( $\$ 2.3$ ) was computed between the model oriented by each cross-rotation peak and each chain of the 1 fq 0 crystal structure. While the average and standard deviation (in parentheses) TRMSD between each crossrotation peak oriented model and the closest chain of the refined structure is $16.4(7.2) \AA$, the closest (i.e. smallest TRMSD) peak to chain $A$ has a TRMSD of $0.01 \AA$ (peak 1), to chain $B$ has a TRMSD of $4.00 \AA$ (peak 137) and to chain $C$ has a TRMSD of $3.44 \AA$ (peak 46). In this test case, although peak 1 corresponds to the orientation closest to chain $A$, the top three rotation-function-ranked cross-rotation peaks do not correspond to correct model orientations. The CRANS algorithm was directed to search for complete threefold NCS (no missing peaks) and identified peaks 1 (chain $A$, TRMSD $0.01 \AA$ ), 159 (chain $B$, TRMSD $6.28 \AA$ ) and 59 (chain $C$, TRMSD $4.00 \AA$ ) as the highest scoring NCS-consistent rotation set (Table 1). While the NCS-consistent rotation set consisting of peaks 1,137 and 46 contains those cross-rotation peaks producing orientations with the smallest TRMSD to the refined structure, these orientations do not strictly obey the NCS as tightly as peaks 1,159 and 59 (originally found by $C R A N S$ ). The orientations specified by cross-rotation peaks 1 and 137 have a relative angle of $108.6^{\circ}$ rather than the NCSspecified $120^{\circ}$. Therefore, as a control, we relaxed the values of $\tau_{\text {axis }}$ and $\tau_{\text {angle }}$ and reran $C R A N S$. With the relaxed thresholds, CRANS was able to find the NCS-consistent rotation set consisting of peaks 1,137 and 46 . Despite the fact that the overall quality of the rotations returned by the 1 fq 0 crossrotation function are low compared with the cross-rotation peaks found in the other test cases (1chp, 1 g 31 and ChDHFRTS), CRANS was still able to extract three rotations that result in properly oriented models.

The column 'Best Pk. in top 10' of Table 1 lists the crossrotation peak with the smallest (best) TRMSD among the top ten rotation-function-ranked cross-rotation peaks. This column lists the peak which might be found in a manual molecular-replacement effort. Because CRANS exhaustively
checks all peaks in the cross-rotation list, NCS-consistent rotation peaks that appear at the bottom of the cross-rotation list are found as easily as those that appear near the top of the list.

### 3.2. Cholera toxin B subunit mutant (fivefold NCS)

Structure factors for cholera toxin B subunit mutant (1chp) were obtained from the PDB. The first chain of 1chp was used as a model in a cross-rotation search using default parameters. The resulting cross-rotation peak list was sorted by rotationfunction score and contained 122 rotations. The TRMSD was computed between the model oriented by each cross-rotation peak and each chain of the 1chp crystal structure. In this case, unlike 1 fq 0 , the top five cross-rotation peaks did have the lowest TRMSD to each of the five chains in the crystal structure, $0.10,0.62,1.02,0.68$ and $0.54 \AA$. The CRANS algorithm was directed to search for complete fivefold NCS (no missing peaks) and identified two sets of cross-rotation peaks. The first set contained peaks 1, 2, 3, 4 and 5 , which correspond to the peaks with the lowest TRMSD to the refined structure (Table 1). CRANS also identified a second complete set of cross-rotation peaks containing peaks 59 (chain $D$, TRMSD $1.96 \AA$ ), 75 (chain $E$, TRMSD $2.83 \AA$ ), 58 (chain $F$, TRMSD $2.83 \AA$ ), 22 (chain $G$, TRMSD $1.91 \AA$ ) and 45 (chain $H$, TRMSD $1.97 \AA$ ). Thus, both identified rotation sets are consistent with the crystal structure.

### 3.3. Gp31 co-chaperonin from bacteriophage T4 (sevenfold NCS)

Structure factors for Gp31 co-chaperonin from bacteriophage T4 (1g31) were obtained from the PDB. The first chain of 1 g 31 was used as a model in a cross-rotation search using default parameters. The resulting cross-rotation peak list was sorted by rotation-function score and contained 47 rotations. The TRMSD was computed between the model oriented by each cross-rotation peak and each chain of the 1 g 31 crystal structure. The peaks with the lowest TRMSD to each crystallographic chain are listed in Table 1. High-degree NCS pushes the limits of standard molecular-replacement methods (Oh, 1995) since in these cases the model corresponds to a smaller percentage of the molecular mass in the unit cell (e.g. only $14.3 \%$ for sevenfold NCS). Therefore, it becomes increasingly likely that one or more NCS-consistent rotations will be missing from the cross-rotation peak list. In these cases, the ability of CRANS to generate missing peaks becomes rather useful. Thus, it is not surprising that a simple crossrotation search for 1 g 31 performed with default search parameters is unable to identify rotations specifying orientations similar to chains $D$ and $E$ (i.e. orientations with low TRMSDs; Table 1). In this case, we would not expect the CRANS algorithm to find NCS-consistent rotation sets with less than two missing peaks. Note that the most complete NCS-consistent set found by the CRANS algorithm has three missing peaks (Table 1), indicating that although peak 8 has a TRMSD of $2.31 \AA$ to chain $G$, the rotation differences between peak 8 and the other selected peaks did not satisfy the axis and angle

Table 1
TRMSDs measured for the three test systems 1 fq 0 , 1 chp and 1 g 31 .
Column 2 lists the PDB chain identifier used in computing the TRMSD for the specified row. The 'Best Pk.' column lists the minimum TRMSD observed between the specified chain and all cross-rotation peaks, along with its corresponding cross-rotation peak index (in parentheses). Cross-rotation peaks are ordered by sorting them based on the cross-rotation function score, where a lower index corresponds to a higher (better) cross-rotation score. The TRMSD and peak index of the peak with the smallest TRMSD among the top ten cross-rotation function ranked peaks is listed in the 'Best Pk. in top 10' column. The TRMSD of the CRANS-identified peak of the NCS-consistent rotation set corresponding to the specified chain and its cross-rotation peak index (in parentheses) is listed in column 'CRANS Pk.'. Peaks computed by CRANS (i.e. those missing in the cross-rotation peak list) are shown in italics with a dash for the peak index.

| System | Chain | Best Pk. | Best Pk. in top 10 | CRANS Pk. |
| :--- | :--- | :--- | :--- | :--- |
| 1fq0 | $A$ | $0.01(1)$ | $0.01(1)$ | $0.01(1)$ |
|  | $B$ | $4.00(137)$ | $14.43(6)$ | $6.28(159)$ |
|  | $C$ | $3.44(46)$ | $16.56(3)$ | $4.00(59)$ |
| 1chp | $D$ | $0.10(1)$ | $0.10(1)$ | $0.10(1)$ |
|  | $E$ | $0.62(4)$ | $0.62(4)$ | $0.62(4)$ |
|  | $F$ | $1.02(3)$ | $1.02(3)$ | $1.02(3)$ |
|  | $G$ | $0.68(5)$ | $0.68(5)$ | $0.68(5)$ |
|  | $H$ | $0.54(2)$ | $0.54(2)$ | $0.54(2)$ |
| 1 g 31 | $A$ | $2.33(2)$ | $2.33(2)$ | $2.49(3)$ |
|  | $B$ | $2.37(6)$ | $2.37(6)$ | $2.37(6)$ |
|  | $C$ | $2.49(4)$ | $2.49(4)$ | $2.49(4)$ |
|  | $D$ | $8.13(42)$ | $10.64(9)$ | $2.29(-)$ |
|  | $E$ | $7.04(1)$ | $7.04(1)$ | $2.29(-)$ |
|  | $F$ | $2.29(1)$ | $2.29(1)$ | $2.29(1)$ |
|  | $G$ | $2.31(8)$ | $2.31(8)$ | $2.37(-)$ |

thresholds $\tau_{\text {axis }}$ and $\tau_{\text {angle }}$. The complete NCS-consistent rotation set generated by CRANS is consistent with the 1 g 31 crystal structure. The three missing peaks (generated by CRANS) have TRMSDs of 2.29, 2.29 and $2.37 \AA$ (Table 1), thus demonstrating the ability of the CRANS algorithm to compute correct and complete NCS-consistent rotation sets even in the presence of missing cross-rotation peaks.

### 3.4. C. hominis DHFR-TS ( $5_{1}$-fold NCS)

Molecular replacement was used to determine initial phase angles for the structure of ChDHFR-TS. Diffraction data to $2.8 \AA$ were collected at Brookhaven National Laboratory (beamline X 12 C ) and processed into structure factors as previously described (O’Neil et al., 2003). Analysis of selfrotation peaks indicated the presence of fivefold NCS. A cross-rotation search using default parameters was performed with each of three models (LmDHFR-TS, PcTSA and PcTSB; see $\S 2.3$ ). The resulting cross-rotation peak lists were sorted by rotation-function score and run through the CRANS algorithm (because each model was a homodimer, CRANS was run in homodimer mode). The LmDHFR-TS, PcTSA and PcTSB cross-rotation peak lists contained 45,38 and 38 rotations, respectively. Cross-rotation peak analysis with CRANS was able to find fivefold NCS sets with one missing peak for LmDHFR-TS, no missing peaks for PcTSA and two missing peaks for PcTSB. The NCS axes computed for all $C R A N S$-identified NCS-consistent rotation sets agreed with the axis identified by the self-rotation search. Models oriented according to the rotations of the complete PcTSA NCS-

Table 2
TRMSDs measured for the three homologous models used in solving the ChDHFR-TS structure.

Column 2 lists the identifier of the homodimer used in computing the TRMSD for the specified row. The 'Closest Pk.' and 'CRANS Pk.' columns are as listed in the caption of Table 1. The 'Closest dimer-flipped Pk.' column is similar to the 'Closest Pk.' column; however, TRMSDs are computed between the dimer flip of the specified homodimer and each peak in the cross-rotation peak list. Because CRANS computes model rotations invariant to homodimer flips, the $C R A N S$-computed TRMSDs are taken as the smaller of the TRMSD to the crystallographic dimer or its dimer flip.

|  |  |  | Closest dimer- <br> Model | Dimer |
| :--- | :--- | :---: | :--- | :--- |
| flipped Pk. |  |  |  |  |$\quad$ CRANS Pk. | CmDHFR-TS | $A$ | $24.92(39)$ | $0.90(4)$ | $0.90(4)$ |
| :--- | :--- | :--- | :--- | :--- |
|  | $B$ | $0.87(1)$ | $21.55(14)$ | $0.87(1)$ |
|  | $C$ | $0.85(2)$ | $0.93(5)$ | $0.85(2)$ |
|  | $D$ | $1.07(3)$ | $20.23(32)$ | $1.07(3)$ |
| PcTSA | $E$ | $23.79(4)$ | $21.61(31)$ | $1.33(-)$ |
|  | $A$ | $30.52(24)$ | $1.14(2)$ | $1.14(2)$ |
|  | $B$ | $1.17(1)$ | $15.92(22)$ | $1.17(1)$ |
|  | $C$ | $13.60(17)$ | $1.14(3)$ | $1.14(3)$ |
|  | $D$ | $1.23(5)$ | $20.83(13)$ | $1.23(5)$ |
| PcTSB | $E$ | $1.11(4)$ | $15.76(36)$ | $1.11(4)$ |
|  | $A$ | $21.34(11)$ | $18.47(10)$ | $1.34(-)$ |
|  | $B$ | $1.21(1)$ | $1.15(2)$ | $1.21(1)$ |
|  | $C$ | $1.19(4)$ | $1.15(3)$ | $1.15(3)$ |
|  | $D$ | $1.19(5)$ | $14.39(8)$ | $1.19(5)$ |
|  | $E$ | $26.63(37)$ | $21.24(35)$ | $1.32(-)$ |

consistent rotation set were positioned using a translation search (O'Neil et al., 2003; Crowther \& Blow, 1967). The initial $R$ factor of $52 \%$ was refined to $22.5 \% ~\left(R_{\text {free }}=24.5 \%\right)$. Refined ChDHFR-TS molecules have a non-crystallographic $5_{1}$ axis (O’Neil et al., 2003) (see Fig. 3).

Analysis performed after the structure determination clearly explains the CRANS results. The TRMSD was
computed between the thymidylate-synthase (TS) domain of each model oriented by each cross-rotation peak and each TS homodimer (dimers $A, B, C, D$ and $E$ ) of the refined ChDHFR-TS crystal structure. Although the DHFR domains were used in the LmDHFR-TS cross-rotation search, the DHFR domains were not used in computing the TRMSDs because of the significant difference in the refined DHFR orientations relative to the highly conserved TS homodimer. The TRMSDs are presented in Table 2.
3.4.1. LmDHFR-TS. TRMSD analysis of the cross-rotation search peaks show that while some peaks closely approximate the structures of homodimers $A, B, C$ and $D$, no cross-rotation peaks returned were similar to homodimer $E$ (Table 2). Despite the significant structural and relative orientational differences between the DHFR domains of LmDHFR-TS and ChDHFR-TS, four of five correct rotations were still found in the LmDHFR-TS cross-rotation search using default search parameters. Table 2 shows that peaks 2 and 5 provide redundant information and both correspond to homodimer $C$ (and its dimer flip). Therefore, the naïve selection of the top 5 crossrotation peaks in the hope that they corresponded to the five correct NCS-related rotations would result in only four of five correct orientations and one redundant orientation. While peaks 2 and 5 corresponded to homodimer $C$ and its dimer flip, the only cross-rotation peaks corresponding to homodimers $A, B$ and $D$ matched either the dimer or its dimer flip (but not both). Consequently, the ability of the CRANS algorithm to handle homodimer models was vital in this analysis. Specifically, a search for rotation-difference subgroups that ignored the fact that the model is a homodimer would not have been able to identify all four NCS-related peaks contained in the cross-rotation peak list. The peaks identified by CRANS


Figure 3
(a) A portion of the unit cell containing the $5_{1}$ screw axis of ChDHFR-TS. The dimerization axis of dimer 3 is coincident with a crystallographic twofold. Dimers $x$ and $x^{\prime}$ are related by a crystallographic twofold. The crystallographic $a$ and $c$ unit-cell edges are shown for reference. (b) Top-down view showing the fivefold symmetry.
correspond to those peaks with the smallest TRMSD for each homodimer (Table 2). Furthermore, the rotation generated by $C R A N S$ to complete the NCS-consistent rotation set has a TRMSD to homodimer $E$ of $1.33 \AA$.
3.4.2. PcTSA. Unlike the LmDHFR-TS model, PcTSA consists of only a TS homodimer. Cross-rotation peak analysis with CRANS found one fivefold NCS-consistent rotation set with no missing peaks (Table 2). The peaks identified by CRANS correspond to those with the smallest TRMSD to each of the ChDHFR-TS homodimers. As with LmDHFR-TS, the ability to handle homodimer models was crucial in $C R A N S$ analysis.
3.4.3. PcTSB. Similar to the PcTSA model, PcTSB consists of only a TS homodimer (see $\S 2.3$ ). Analysis of the crossrotation peak sets with CRANS could not identify a fivefold NCS-consistent rotation sets with zero or one missing peaks; however, four NCS-consistent rotation sets were found with peaks corresponding to three of the five TS homodimers. Because peaks 1 and 2 and peaks 3 and 4 provide redundant information and both correspond to orientations of homodimer $B$ and homodimer $C$, respectively, of the refined structure, the four identified sets correspond to the four sets $\{\{1,3,5\},\{2,3,5\},\{1,4,5\},\{2,4,5\}\}=\{1,2\} \times\{3,4\} \times\{5\}$ (numbers are cross-rotation peak numbers sorted by rotationfunction score). The top scoring set was $\{1,3,5\}$, which has TRMSDs of $1.21,1.15,1.19 \AA$ to the crystallographic ChDHFR-TS homodimers. The two computed (missing peaks) for this set have TRMSDs of 1.32 and $1.34 \AA$ (Table 2). Although the top five cross-rotation peaks had a rotationfunction score approximately twice that of the remaining peaks, the direct application of the top five peaks would not have resulted in the five orientations seen in the refined structure. Therefore, CRANS analysis provided the important information that not all five of the orientations seen in the refined structure were seen among the top five peaks of the cross-rotation list.

In summary, although two of the three cross-rotation peak lists did not contain all five NCS-consistent model rotations, the $C R A N S$ algorithm was successfully able to (i) verify that fivefold NCS was present, (ii) find sets of cross-rotation peaks related by an NCS-axis consistent with the self-rotation function for all three models and (iii) compute missing crossrotation peaks corresponding to orientations with TRMSDs of $1.33,1.34$ and $1.32 \AA$ to the final crystal structure. The identified NCS-consistent rotation sets were then used in a translation-function and rigid-body refinement to generate initial phase angles and the final structure of ChDHFR-TS (PDB code 1qzf; O'Neil et al., 2003). The $C R A N S$-identified NCS-consistent rotation sets were correct and agree with the final refined structure.

## 4. Discussion

In all six cross-rotation function searches performed only the default search parameters were used. That is, we did not spend any time optimizing cross-rotation search parameters. It is possible that by tweaking cross-rotation function search
parameters and by optimizing the model (i.e. removing flexible loops, changing residues to Ala etc.) more NCS-consistent rotations could have been returned by the cross-rotation function. The time required to perform these optimizations can be reduced or eliminated by using $C R A N S$ to analyze the output of a cross-rotation search run with default parameters.

As the number of peaks returned by the cross-rotation function increases, the probability that these peaks will conspire to form 'fake' low-degree NCS-consistent rotation sets rises. This is especially true if the axis and angle tolerances are not particularly tight. Therefore, when analyzing lowdegree NCS with large cross-rotation peak lists, the results of a $C R A N S$ search should be treated as a working hypothesis. Confidence in CRANS-identified NCS-consistent rotation sets can be increased by directing CRANS to use the NCS axis identified by the self-rotation map.

CRANS identifies NCS-consistent rotation sets from a cross-rotation search using monomer or homodimer models, since these are the most common model types used in molecular replacement; however, the CRANS algorithm can be extended to handle any degree of oligomerization (e.g. models that are homotrimers, homotetramers or homopentamers). Conceptually, to handle higher order model symmetry all symmetry rotations of the model are computed and then applied to each cross-rotation peak. For example, with $d$-fold model symmetry ( $d$-fold oligomerization), let $f$ be the $360 / d^{\circ}$ rotation around the symmetry axis. Each rotation $r$ identified by the cross-rotation function is replaced by the $d$ rotations $r f^{z}$ $(z \in\{0,1, \ldots, d-1\})$. Heterodimers do not inherently contain a symmetry axis and therefore avoid the rotational degeneracy presented by homodimers. Thus, any model which does not have a symmetry axis (e.g. heterodimers, heterotrimers) should be treated as a monomer by CRANS.

While the tests in this paper were performed with an ordinary cross-rotation function, an alternative is to use a locked cross-rotation function (Tong \& Rossmann, 1990; Tong, 2001) when the NCS axis is clear from the self-rotation function. Although the scoring of the rotation-function peaks is different, in the presence of NCS both the ordinary and locked cross-rotation functions should identify rotations corresponding to each of the $n$ NCS-consistent orientations. When using a locked rotation function, non-crystallographic symmetry mates are generated using the NCS axis initially identified by the self-rotation function. However, if the NCS axis cannot be defined with a high degree of accuracy or the conservation of non-crystallographic symmetry in the crystal is not perfect, then a more accurate orientation for each monomer may be identified by orienting each monomer individually. CRANS can process the results of ordinary or locked cross-rotation functions to return a set of NCSconsistent orientations. The NCS-consistent model orientations identified and generated by $C R A N S$ can be positioned either sequentially using an ordinary translation function (Crowther \& Blow, 1967) or simultaneously using a locked translation function (Tong \& Rossmann, 1990; Tong, 2001) followed by an ordinary translation function (Tong, 2001).

## 5. Limitations and extensions

In this section, we discuss use of the CRANS algorithm in three situations: (i) crystals with improper NCS, (ii) crystals with multiple intersecting proper NCS axes and (iii) crystals with parallel NCS and crystallographic axes.
(i) The orientations of molecules involved in $n$-fold proper NCS are related by rotations of magnitude $360 t / n(t \in\{0,1$, $\ldots, n-1\}$ ) around the NCS axis. By clustering the set of rotation differences between all pairs of cross-rotation peaks, CRANS is able to identify proper NCS-consistent rotation sets. Because there are no a priori constraints on the relative orientations between molecules with improper NCS, CRANS is unable to identify peaks whose orientations are only related by improper NCS. However, when a crystal possesses a combination of proper and improper NCS, the CRANS algorithm should identify the proper NCS, leaving those copies of the model only related by improper NCS to be positioned using alternative means. For example, in this situation, the Matthews coefficient (Matthews, 1968) might indicate the presence of $k$ copies of the molecule per unit cell, while the self-rotation function indicates the presence of only $n$-fold proper NCS $(n<k)$. If $C R A N S$ is able to identify one or more NCS-consistent rotation sets involving $n$ molecules then one might assume that the remaining $k-n$ molecules are related by improper NCS. Alternatively, CRANS could be run in its scan mode to look for proper NCS with degree from 2 to $k$. In this scenario, CRANS might identify the presence of $n$-fold NCS, leaving the remaining $k-n$ molecules to be positioned by other techniques.
(ii) The identification of model orientations for crystals with multiple intersecting non-crystallographic rotation axes (i.e. 532 or 72 symmetry) is a challenge. The $C R A N S$ algorithm can be used to assist in the determination of correct orientations by identifying the cross-rotation peaks involved in each point group. CRANS will output all NCS-consistent rotation sets satisfying the $\tau_{\text {angle }}$ and $\tau_{\text {axis }}$ thresholds. Therefore, if there are multiple point-group symmetries satisfying these thresholds they will all be output by the $C R A N S$ algorithm. For example, in the case of 72 symmetry, CRANS could be directed to identify sevenfold proper NCS, which would identify two sevenfold NCS-consistent rotation sets with parallel axes. In the same situation, if CRANS were instead directed to search for twofold proper NCS, it would identify seven twofold proper NCS-consistent rotation sets with parallel axes. By independently identifying each NCS-consistent rotation set, CRANS could provide supporting evidence for the presence of multiple non-crystallographic rotation axes.
(iii) When an NCS axis is parallel to the crystallographic axis, cross-rotation peaks corresponding to NCS-consistent orientations may be buried under the crystallographic symmetry peaks. When this occurs, the cross-rotation function may have difficulty identifying accurate NCS-consistent orientations of the model. Because CRANS relies on the output of a cross-rotation function, it is limited by the quality of the identified cross-rotation peaks. Therefore, we would expect CRANS to have difficulty identifying NCS-consistent
rotation sets with parallel crystallographic and non-crystallographic axes.

## 6. Conclusions

In this paper, we have presented the CRANS algorithm for analyzing lists of cross-rotation peaks both to extract NCSconsistent rotation sets and to complete partial sets by computing missing NCS-consistent rotations. We showed that the problem of identifying NCS-consistent rotation sets is equivalent to subgroup identification among differences in rotations of the cross-rotation peak list. We then tested the algorithm on four test proteins displaying three-, five- and sevenfold proper NCS using six models for molecular replacement (three from the solved crystal structure itself and three from homologous proteins). We demonstrated the ability of the CRANS algorithm to find NCS-consistent rotation sets both when all appropriate rotations were present and when up to three rotations (in the case of 1 g 31 ) were missing. Furthermore, CRANS successfully identified orientations that were used to generate initial phases in solving the structure of ChDHFR-TS. For all test cases, the CRANS algorithm was able to successfully generate correct NCS-consistent rotation sets. The CRANS algorithm is efficient, requiring only seconds on an Athlon-based processor to search for up to eightfold proper NCS on lists of 120 cross-rotation peaks.

By extracting more information from each cross-rotation peak list, the CRANS algorithm provides two main benefits to the crystallographer. Firstly, in the case where the crossrotation peak list contains $n$ peaks with significantly higher scores than the remainder of the list (where $n$ is the NCS degree), the CRANS algorithm can confirm that these top peaks are indeed consistent with known proper NCS. The importance of this confirmation was demonstrated in the 1 fq 0 , 1 g 31 and ChDHFR-TS systems, where the top $n$ peaks did not correspond to the correct NCS-consistent orientations despite a sharp dropoff in rotation-function score after these top peaks. Secondly, when the model is partial or the NCS degree is high, it becomes likely that one or more NCS-consistent rotations will not be present in the cross-rotation peak list. When this happens, it is not possible to find a complete list of NCS-consistent cross-rotation peaks. The CRANS algorithm can find partial proper NCS-consistent rotation sets and then generate the missing proper NCS-consistent rotations to create a complete set. The ability to correctly generate missing peaks was demonstrated in the 1 g 31 , LmDHFR-TS and PcTSB cases. By using quaternions to generate missing rotations, we avoid the rotational instability (i.e. singularities) that can arise when using other rotation representations.

## 7. Supporting materials

The CRANS program is distributed as a java jar file and is available at http://www.cs.dartmouth.edu/brd/Bio and by contacting the authors. The software is distributed under the Gnu Public License (Gnu, 2002).

## APPENDIX A

## Subgroup search

In this appendix, we show that the search for finite subgroups of $S O(3)$ (the special orthogonal group of three-dimensional rotations) among the rotation differences of cross-rotation peaks provably identifies NCS-consistent rotation sets. The rotation difference set of a set of $n$ rotations $R$ is defined as

$$
\begin{equation*}
D(R)=\left\{r_{i}^{-1} r_{j} \mid r_{i}, r_{j} \in R\right\} \tag{2}
\end{equation*}
$$

(where equal rotations are identified). Recall that $r_{i}^{-1} r_{j}$ is the rotation difference $d_{i j}$ between rotations $r_{i}$ and $r_{j}$. When $D(R)$ is a subgroup of $S O(3)$, we say that $R$ and $D(R)$ are complete. When $R$ is complete, a partial rotation difference set is a subset $D\left(R^{\prime}\right) \subset D(R)$ constructed from a subset of rotations $R^{\prime}$ of $R$. Therefore, a partial rotation-difference set can be made complete by adding missing rotation differences to complete the subgroup.

The goal of our algorithm is the following: given a set of cross-rotation peaks $C \subset S O$ (3), find finite subsets of rotations $R \subset C$ such that $D(R)$ is a subgroup of $S O(3)$.

We now show that an NCS-consistent rotation set $R$ will have a complete rotation-difference set $D(R)$ satisfying the subgroup axioms and, conversely, that a rotation set $R$ with a complete rotation-difference set $D(R)$ that satisfies the subgroup axioms is an NCS-consistent cross-rotation peak set.

For $D(R)$ to be a subgroup, the four group properties of identity, inverse, associativity and closure must be satisfied. All complete rotation difference sets satisfy the properties of identity, inverse and associativity. By definition (2 the rotation difference $d_{i i}=I$ (where $I$ is the identity element) is present in all complete rotation-difference sets. The inverse property for each rotation difference $d_{i j}$ from $r_{i}$ to $r_{j}$ is satisfied by the rotation from $r_{j}$ to $r_{i}$; that is, $d_{i j} d_{j i}=I$. Finally, all rotations compose associatively by construction. Thus, the crux of the proof is in the closure property. To prove the forward direction, we note that all rotation differences are a multiple of $360 / n^{\circ}$ around a common axis (where $n$ is the degree of NCS) and that all $n$ rotations $360 t / n(t \in\{0,1, \ldots, n-1\})$ are represented in the set $D(R)$. Thus, the application of any number of rotation differences results in a rotation that (modulo $360^{\circ}$ ) is already in the complete rotation-difference set. Conversely, to prove the reverse direction, all finite subgroups have an identity and satisfy the closure property (modulo $360^{\circ}$ ); therefore, if they contain $n$ members they represent $n$-fold symmetry and the symmetry angles $360 t / n(t \in\{0,1, \ldots, n-1\})$. This completes the proof.

When the model is a homodimer, each cross-rotation peak and its corresponding dimer-related symmetry mate produce equivalently oriented dimers. Therefore, each cross-rotation peak and its symmetry mate must be used to form finite subgroups and subsequently NCS-consistent rotation sets.

We now present a synthetic two-dimensional threefold NCS example to illustrate the framework presented in this section. Fig. 1(a) shows the model and five resulting cross-rotation function peaks $C=\left\{r_{1}, r_{2}, r_{3}, r_{4}, r_{5}\right\}$. The set of peaks $R=\left\{r_{1}, r_{3}, r_{4}\right\}$ forms an NCS-consistent rotation set. The
complete rotation difference set $D(R)$ contains three rotations around an NCS-axis coming out of the page, $d_{\alpha}$ corresponds to the identity rotation, $d_{\beta}$ corresponds to a $120^{\circ}$ clockwise or $240^{\circ}$ counterclockwise rotation and $d_{\gamma}$ corresponds to a $240^{\circ}$ clockwise or a $120^{\circ}$ counterclockwise rotation. The subgroup $D(R)$ satisfies the four group properties (see Fig. 1a): the group property of associativity is satisfied by all rotations, $d_{\alpha}$ is the identity rotation and also serves as its own inverse, while $d_{\beta}$ is the inverse of $d_{\gamma}$ and $d_{\gamma}$ is the inverse of $d_{\beta}$. Finally, the closure property is satisfied, $d_{\alpha} d_{\beta}=d_{\beta} d_{\alpha}=d_{\beta}, d_{\alpha} d_{\gamma}=d_{\gamma} d_{\alpha}=d_{\gamma}$ and $d_{\beta} d_{\gamma}=d_{\gamma} d_{\beta}=d_{\alpha}$.

## APPENDIX B <br> CRANS input

The CRANS input file is a list of cross-rotation peaks in the $z x z$ Euler angle rotation convention. In the $z x z$ Euler angle format, the first angle specifies the rotation around the $z$ axis, the second angle specifies the rotation around the new $x$ axis and the third angle specifies the rotation around the new $z$ axis. $C R A N S$ requires that the cross-rotation peak list have five columns corresponding to the peak index, $z$ rotation, $x$ rotation, $z$ rotation and rotation-function score. Because this is the format used by CNS (Brünger et al., 1998), the peak-list file generated by a CNS cross-rotation search can be used directly by the CRANS program.

This work is supported by grants to BRD from the National Institutes of Health (GM-65982) and the National Science Foundation (IIS-9906790, EIA-0102710, EIA-0102712, EIA9818299, EIA-9802068 and EIA-0305444) and to ACA from the NIH (GM-067542). We thank Ramgopal Mettu, Robert O'Neil, Anthony Yan, Robert Stroud and all members of the Donald and Anderson Labs for helpful discussions and comments on drafts.

## References

Anderson, A., O'Neil, R., Surti, T. \& Stroud, R. (2001). Chem. Biol. 8, 445-457.
Berman, H., Westbrook, J., Feng, Z., Gilliland, G., Bhat, T., Weissig, H., Shindyalov, I. \& Bourne, P. (2000). Nucleic Acids Res. 28, 235242.

Brünger, A. T., Adams, P. D., Clore, G. M., DeLano, W. L., Gros, P., Grosse-Kunstleve, R. W, Jiang, J.-S., Kuszewski, J., Nilges, M., Pannu, N. S., Read, R. J., Rice, L. M., Simonson, T. \& Warren, G. L. (1998). Acta Cryst. D54, 905-921.

Collaborative Computational Project, Number 4 (1994). Acta Cryst. D50, 760-763.
Crowther, R. A. \& Blow, D. M. (1967). Acta Cryst. 23, 544-548.
Evans, P. R. (2001). Acta Cryst. D57, 1355-1359.
Gnu (2002) The GNU General Public License. http://www.gnu.org/ licenses/licenses.html.
Hamilton, W. (1969). Elements of Quaternions, 3rd ed. New York: Chelsea Publishing Co.
Hunt, J., van der Vies, S., Henry, L. \& Deisenhofer, J. (1997). Cell, 90, 361-371.
Knighton, D., Kan, C., Howland, E., Janson, C., Hostomska, Z., Welsh, K. \& Matthews, D. (1994). Nature Struct. Biol. 1, 186-194. Matthews, B. W. (1968). J. Mol. Biol. 33, 491-497.
Merritt, E., Sarfaty, S., Chang, T., Palmer, L., Jobling, M., Holmes, R. \& Hol, W. (1995). Structure, 3, 561-570.

Oh, B.-H. (1995). Acta Cryst. D51, 140-144.
O'Neil, R., Lilien, R., Donald, B., Stroud, R. \& Anderson, A. (2003). J. Biol. Chem. 278, 52980-52987.

Rossmann, M. G. (1990). Acta Cryst. A46, 73-82.
Rossmann, M. G. \& Blow, D. M. (1962). Acta Cryst. 15, 24-31.
Salamin, E. (1979). Applications of Quaternions to Computation with Rotations. Technical Report, Stanford University, AI Lab. Internal working paper.

Tong, L. (2001). Acta Cryst. D57, 1383-1389.
Tong, L. \& Rossmann, M. G. (1990). Acta Cryst. A46, 783792.

Vagin, A. \& Teplyakov, A. (1997). J. Appl. Cryst. 30, 10221025.

Wymer, N., Buchanan, L., Henderson, D., Mehta, N., Botting, C., Pocivavsek, L., Fierke, C., Toone, E. \& Naismith, J. (2001). Structure, 9, 1-9.


[^0]:    ${ }^{1}$ Abbreviations used: $S O(3)$, group of three-dimensional rotations; MR, molecular replacement; model, molecular-replacement model; DHFR, dihydrofolate reductase; TS, thymidylate synthase; ChDHFR-TS, Cryptosporidium hominis DHFR-TS; LmDHFR-TS, Leishmania major DHFR-TS; PcTS, Pneumocystis carinii TS; NCS, non-crystallographic symmetry (proper, unless otherwise noted); RMSD, root-mean-square distance.

