Embarrass de richesses – It is not good to be too anomalous: Accurate structure of selenourea, a chiral crystal of planar molecules

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Abstract

Selenourea, SeC(NH₂)₂, recently found an application as a derivatization reagent providing a significant anomalous diffraction signal used for phasing macromolecular crystal structures. The crystal structure of selenourea itself was solved about 50 years ago, from data recorded on films and evaluated by eye and refined to \( R = 0.15 \) with errors of bond lengths and angles about 0.1 Å and 6°. In the current work this structure is re-evaluated on the basis of synchrotron data and refined to \( R_1 = 0.021 \) with bond and angle errors about 0.007 Å and 0.5°. The nine planar molecules of selenourea pack either in the \( P_3_1 \) or in the \( P_3_2 \) unit cell. All unique molecules are connected by a complex network of Se•••H-N hydrogen bonds and Se•••Se contacts. The packing of selenourea molecules is highly pseudosymmetric, approximating either of the \( P_3_1(2)12, \ R_3, \) and \( R_32 \) space groups. Because the overwhelming majority of diffracted X-ray intensity originates from the anomalously scattering selenium atoms, the measurable anomalous Bijvoet differences are diminished and it was not possible to solve this crystal structure based on the anomalous signal alone.

Introduction

Selenourea, SeC(NH₂)₂ (SeUrea) is a small molecule, the selenium derivative of urea (Fig 1), which was the first ever organic compound that was synthesized chemically by Wöhler in 1828 [1]. After several early attempts [2,3], the crystal structure of SeUrea was published almost 50 years ago by Rutherford and Calvo ([4], thereafter R&C), with diffraction data recorded on photographic film using Weissenberg and precession cameras. This structure, containing nine molecules of SeUrea in the asymmetric unit in the space group \( P_3_1 \), was refined to \( R_1 = 0.120 \) at the room temperature and \( R = 0.116 \) at -100°C. In the latter structure, the estimated standard deviations (esd’s) of bond lengths were about 0.1 Å and esd’s of the bond angles were about 6°.

Our interest in SeUrea was triggered by its use as a convenient reagent for derivatization of macromolecular crystals leading to determination of structures based on the anomalous signal...
of the incorporated Se atoms [5]. We have obtained accurate diffraction data from SeUrea crystals with the synchrotron radiation and refined the atomic model to estimated bonds and angles accuracy of about 0.007 Å and 0.5˚. This accurate model of the SeUrea molecule provides appropriate geometrical targets for refinement of this ligand in complexes with macromolecules, including the hydrogen bonds involving SeUrea.

In addition, the crystal structure of SeUrea presents an interesting and exceptional case from the crystallographic methodology point of view, in the context of its high degree of pseudosymmetry and unusual content of the anomalous signal in diffraction data. These features are discussed and illustrated in detail.

Methods

The crystalline powder of SeUrea (Sigma-Aldrich) was soaked in 50% (v/v) 2-methyl-2,4-pentanediol (MPD) for 5 min. The remaining undissolved SeUrea crystals were picked up by a nylon loop and vitrified in liquid nitrogen for diffraction data collection. The MPD was used as a cryoprotectant to prevent the ice formation after cooling crystals to 100 K in the stream of cold nitrogen gas. Two data sets from two separate crystals were collected at the wavelength of 0.8 Å on the Rayonix MX300HS detector at the SER-CAT 22ID beam line of the Advanced Photon Source, Argonne National Laboratory, USA. The data were processed by HKL2000 [6] without merging the Friedel mates and converted to SHELX format by XPREP [7]. The crystal structure was solved by SHELXT [8] and refined by SHELXL [9]. The two crystals turned out to have an opposite chirality, in space groups P31 and P32. The better refined structure in P31 is discussed in the following sections, but the diffraction data and refinement statistics are presented in Table 1 for both crystals. The atomic parameters of the P31 structure are listed in Table 2. The bond lengths and angles in all nine independent molecules of SeUrea are listed in Table 3.

Structure

In general, the structure of SeUrea is isomorphous to that described by R&C. The unit cell of the P31 symmetry contains 27 molecules of SeUrea arranged in five helical columns around three non-equivalent crystallographic 31 axes and two similar columns forming the non-crystallographic, approximate 31 axes, as illustrated in Fig 2. The details of this highly pseudosymmetric arrangement are discussed in the next section.

The van der Waals radius of the Se atom is 1.90 Å [10], but many Se...Se contacts between neighboring columns are slightly shorter than twice of this value, down to 3.55 Å (Fig 3A and 3B, Table 4). The SeUrea molecules within each 31 column are connected through hydrogen bonds between one of the amide NH2 groups of one SeUrea molecule (H-bond donor) and the Se atom of the SeUrea molecule from the next layer (Fig 3C and 3D and Table 5). In addition,
the amide groups of SeUrea molecules are engaged in hydrogen bonds with Se atoms form neighboring columns. In effect, all hydrogen atoms of the amide groups serve as donors in hydrogen bonds except for one (on the N1_3 atom, Table 5), and some are engaged in bifurcated H-bonds with two Se acceptors. The donor-acceptor lengths of all H-bonds vary between 3.4 and 3.8 Å, within the range observed also in other small molecule crystal structures containing the selenocarbonyl group [11].

The crystal structure of SeUrea is noncentrosymmetric, even though the individual SeUrea molecules are planar and have the mm2 (C\textsubscript{2v}) symmetry. The chirality of the space group P3\textsubscript{1} is therefore a result of specific packing of SeUrea molecules in the crystal. Indeed, the crystal structure of SeUrea described here is in the space group P3\textsubscript{1}, which is supported by the Flack parameter [12] refined to -0.003(6), whereas after inversion of atomic coordinates and change of the space group to P3\textsubscript{2}, the Flack parameter became 1.013(6). However, the structure of the P3\textsubscript{2} crystal, obtained in exactly same way, refined to R1 = 0.032 with the Flack parameter of 0.009(11) in the space group P3\textsubscript{2} (and of 1.003(11) after inversion to P3\textsubscript{1}), confirming that the chirality of the SeUrea crystals is selected at random.

The final model of SeUrea discussed here was refined against the diffraction data in the space group P3\textsubscript{1} with merged Friedel mates.

### Pseudosymmetry

As pointed out by R&C, the arrangement of 27 SeUrea molecules in the whole unit cell is characterized by a certain degree of pseudosymmetry and this was utilized by them in the original structure solution. Each of the three crystallographic 3\textsubscript{1} axes at (0,0,z), (1/3,2/3,z) and (2/3,1/3, z) is surrounded by three SeUrea molecules forming right-handed helices. Two triples of SeUrea molecules form very similar helices around approximately (1/3,1/3,z) and (2/3,2/3,z) lines, according to the two additional, approximate (non-crystallographic) 3\textsubscript{1} axes, Fig 2A.
effect 9 of the total 27 SeUrea molecules in the unit cell are arranged around the crystallographic 3 axes and the remaining 18 molecules are analogously arranged around the non-crystallographic 3 axes. All SeUrea molecules are positioned in three layers with respect to the c-axis with the fractional z coordinates of the Se atoms in the following three ranges: 0.13–0.28, 0.46–0.61, 0.80–0.95, with average values separated by 1/3 of the c cell parameter (Fig 2B). The location of Se atoms in the unit cell is presented in Fig 4A, which shows both crystallographic and approximate, non-crystallographic three-fold axes of the 3, 3, and 3 types. The arrangement of all three kinds of the threefold axes corresponds to the R3 space group with three times smaller unit cell with the edges transformed as follows: 

\[ a^R = \frac{2a^2 - b^2}{3}, \quad b^R = \frac{2a^2 + b^2}{3}, \quad c^R = c^p. \]

Table 2. Atomic coordinates, Ueq values (in Å²). The uncertainties (related to the last decimal digit) are in parentheses.

| Atom | x     | y     | z     | Ueq  |
|------|-------|-------|-------|------|
| Se_1 | -0.03486(4) | -0.06824(4) | 0.27914(5) | 0.00817(15) |
| C_1  | 0.0452(4) | 0.0743(4) | 0.2769(4) | 0.0120(12) |
| N1_1 | 0.0567(4) | 0.1257(4) | 0.1908(4) | 0.0199(12) |
| N2_1 | 0.0933(4) | 0.1257(4) | 0.3615(4) | 0.0241(12) |
| Se_2 | 0.64231(5) | 0.26430(4) | 0.15816(5) | 0.01333(15) |
| C_2  | 0.6720(4) | 0.3998(4) | 0.1619(4) | 0.0108(12) |
| N1_2 | 0.6613(4) | 0.4447(4) | 0.0788(4) | 0.0186(11) |
| N2_2 | 0.7017(4) | 0.4528(4) | 0.2490(4) | 0.0226(12) |
| Se_3 | 0.28985(5) | 0.60531(4) | 0.13361(5) | 0.01084(15) |
| C_3  | 0.3820(4) | 0.7454(4) | 0.1339(4) | 0.0110(12) |
| N1_3 | 0.4000(4) | 0.7995(4) | 0.0483(4) | 0.0173(11) |
| N2_3 | 0.4240(4) | 0.7924(4) | 0.2232(4) | 0.0182(11) |
| Se_4 | 0.29828(4) | 0.35726(4) | 0.23551(4) | 0.00683(14) |
| C_4  | 0.3585(4) | 0.2755(4) | 0.2253(4) | 0.0076(11) |
| N1_4 | 0.4147(4) | 0.2827(4) | 0.1435(3) | 0.0137(10) |
| N2_4 | 0.3458(4) | 0.2093(4) | 0.2991(4) | 0.0188(11) |
| Se_5 | 0.29979(4) | 0.26410(5) | 0.55963(4) | 0.00569(13) |
| C_5  | 0.4185(4) | 0.3930(4) | 0.5505(4) | 0.0077(10) |
| N1_5 | 0.4373(4) | 0.4118(3) | 0.4620(4) | 0.0162(11) |
| N2_5 | 0.4738(3) | 0.4360(4) | 0.6349(3) | 0.0127(10) |
| Se_6 | 0.38732(4) | 0.35966(4) | 0.89154(4) | 0.00566(14) |
| C_6  | 0.2443(4) | 0.2873(4) | 0.8962(4) | 0.0079(11) |
| N1_6 | 0.1906(4) | 0.2320(4) | 0.8158(3) | 0.0125(10) |
| N2_6 | 0.1956(4) | 0.2916(4) | 0.9795(4) | 0.0161(11) |
| Se_7 | 0.71170(4) | 0.68529(4) | 0.18036(4) | 0.00535(14) |
| C_7  | 0.5685(4) | 0.6225(4) | 0.1820(4) | 0.0064(11) |
| N1_7 | 0.5141(4) | 0.5738(4) | 0.0995(3) | 0.0144(11) |
| N2_7 | 0.5199(4) | 0.6252(4) | 0.2669(3) | 0.0121(10) |
| Se_8 | 0.62817(4) | 0.69335(4) | 0.51909(4) | 0.00597(14) |
| C_8  | 0.7309(4) | 0.6582(4) | 0.5228(4) | 0.0113(12) |
| N1_8 | 0.7765(4) | 0.6542(4) | 0.4373(3) | 0.0140(10) |
| N2_8 | 0.7586(4) | 0.6367(4) | 0.6116(4) | 0.0242(12) |
| Se_9 | 0.63120(4) | 0.60300(4) | 0.85061(4) | 0.00529(13) |
| C_9  | 0.6509(4) | 0.7353(4) | 0.8606(4) | 0.0101(11) |
| N1_9 | 0.6339(4) | 0.7791(4) | 0.7803(3) | 0.0151(11) |
| N2_9 | 0.6808(4) | 0.7864(4) | 0.9476(4) | 0.0206(11) |
However, the repetition of Se atom positions in the a,b-plane is more symmetric than along the c direction. Superposition of all five individual helical triples of Se atoms in the a,b-plane (disregarding differences in z coordinates) gives the root-mean-square (rms) deviation from the average position of 0.20 Å. The rms deviation of the Se atoms z-coordinates from their respective average values of the three layers is 0.67 Å. The rms deviation of Se atoms from the ideal R3 symmetry is 1.07 Å.

In addition to the arrangement of Se atoms according to the set of the exact and approximate 3-fold axes, they also conform to a number of the approximate 2-fold symmetry axes perpendicular to the c axis, which together with the crystallographic 3_1 axes form the approximate P3_112 space group, as illustrated in Fig 4B. The rms deviation of the Se atoms from this idealized arrangement is 1.33 Å. The combination of all approximate 3-fold and 2-fold axes of various types forms the R32 space group presented in Fig 4C (within the three times smaller unit cell) where the Se atoms agree with this symmetry within the rms deviation of 1.61 Å.

The pseudosymmetric arrangement of SeUrea molecules in the unit cell consequently modulates the reflection intensities in the reciprocal space, evident in the statistics of the normalized structure factors. The \( \langle E^2 \rangle \) value for all 2797 reflections is 0.993, for 931 reflections with \( -h+k = 3n \) is 2.086, and for 1866 reflections with \( -h+k \neq 3n \) is 0.448, clearly identifying the

| Molecule | Se-C   | C-N1   | C-N2   | Se-C-N1 | Se-C-N2 | N1-C-N2 |
|----------|--------|--------|--------|---------|---------|---------|
| 1        | 1.873(6) | 1.317(7) | 1.327(8) | 120.7(4) | 120.8(4) | 118.5(5) |
| 2        | 1.867(6) | 1.322(7) | 1.323(8) | 121.3(4) | 120.7(4) | 118.0(5) |
| 3        | 1.866(6) | 1.313(7) | 1.337(7) | 122.0(4) | 119.5(4) | 118.6(5) |
| 4        | 1.872(6) | 1.325(7) | 1.326(7) | 120.4(4) | 121.0(4) | 118.6(5) |
| 5        | 1.882(5) | 1.312(7) | 1.330(7) | 121.0(4) | 119.8(4) | 119.2(5) |
| 6        | 1.876(6) | 1.326(7) | 1.323(7) | 120.5(4) | 120.4(4) | 119.2(5) |
| 7        | 1.881(6) | 1.321(7) | 1.331(7) | 120.7(4) | 120.4(4) | 118.7(5) |
| 8        | 1.863(6) | 1.320(7) | 1.316(7) | 121.2(4) | 120.3(4) | 118.6(5) |
| 9        | 1.876(6) | 1.323(7) | 1.309(7) | 120.9(4) | 120.8(4) | 118.2(5) |
| Average  | 1.873(6) | 1.322(7) | 1.307(7) | 120.7(5) | 118.6(4) |

Fig 2. SeUrea crystal packing viewed along c axis (A) and a axis (B) in the unit cell. The Se atoms are colored black and the remaining atoms are colored separately for each group of symmetry equivalent molecules. The atoms are represented as thermal ellipsoids of 50% probability. The Figure was created by Mercury [20].

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Table 3. Bonds lengths (Å) and angles (°).

Average structure of selenourea

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prominence of the smaller cell. The statistics of reflections of the pseudo-rhombohedral cell, with indices transformed accordingly \([H = (-h+k)/3, \ K = (h+2k)/3, \ L = l]\) are also indicative for the small cell, with \(<E^2>\) value for all 931 reflections of 0.982, for 309 reflections with—\(H+K+L = 3n\) of 2.099, and for 622 reflections with—\(H+K+L\neq3n\) of 0.428. It may be pointed out

\[\text{Fig 3. The Se•••Se (black dash) contacts and hydrogen bonds (grey dash) between SeUrea molecules. (A) The Se•••Se contacts between neighboring SeUrea molecules viewed along the c axis. (B) The same contacts viewed along the a axis. (C) H-bonds between SeUrea molecules in the neighboring 3\_1 columns viewed along the c axis. (D) H-bonds between SeUrea molecules within the same 3\_1 columns viewed along the a axis. The three leftmost columns show the crystallographic and the two rightmost columns show the non-crystallographic 3\_1 axes. The Figure was created by Mercury \[20\].}
\]

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| Table 4. Intermolecular contacts between Se atoms (Å) comparable with the sum of van der Waals radii (2 x 1.90 Å). The uncertainties of these distances are 0.001 Å. |
| Se_1   | Se_2   | Se_3   | Se_4   | Se_5   | Se_6   | Se_7   | Se_8   | Se_9   |
|--------|--------|--------|--------|--------|--------|--------|--------|--------|
| Se_2   | -      | -      | -      | 3.563  | -      | -      | -      | 3.668  | -      |
| Se_3   | -      | -      | -      | 4.039  | -      | -      | -      | 3.705  | -      |
| Se_4   | -      | 3.668  | -      | -      | -      | -      | -      | 3.724  | -      |
| Se_5   | 3.563  | -      | -      | -      | -      | -      | -      | 3.761  | -      |
| Se_6   | -      | 3.668  | -      | -      | -      | -      | -      | 3.724  | -      |
| Se_7   | 3.992  | -      | -      | 3.761  | -      | -      | -      | 3.724  | -      |
| Se_8   | -      | 3.548  | 3.705  | -      | -      | -      | -      | -      | -      |
| Se_9   | -      | 3.659  | -      | -      | -      | -      | -      | 3.724  | -      |

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that the intensity statistics depend mainly on the positions of the Se atoms, since they are responsible for $Z^2(\text{Se})/ [Z^2(\text{Se}) + Z^2(\text{C}) + 2 Z^2(\text{N}) + 4 Z^2(\text{H})] = 0.90$, i.e. about 90% of the total reflection intensities. All diffraction data merge in the $P_3_12$ symmetry with the $R_{\text{merge}} = 0.203$ and the subset of rhombohedral reflections $(H,K,L)$ merge in the $R_32$ symmetry with $R_{\text{merge}} = 0.154$, confirming a high degree of pseudosymmetry present in the reciprocal space. No merohedral twinning, theoretically possible in the $P_3_12$ symmetry, was evident in the diffraction data statistics or during refinement of the structure by SHELXL [9].

**Anomalous signal in the diffraction data**

At the wavelength of 0.80 Å used for collecting the diffraction data, the anomalous scattering coefficients of Se are: $\Gamma' = -0.64$ e, $\Gamma'' = 2.72$ e. It was therefore expected that a significant anomalous signal would be present in the diffraction data (Fig 5). However, the crystal of SeUrea constitutes an interesting and somewhat complicated case in the context of the anomalous scattering effects.

If crystal structure consists of a number of “normally” scattering atoms (without any anomalous component) and of several atoms displaying observable anomalous effect, the total structure factor $F_T$ of a selected reflection can be presented as a vector sum of contributions from individual atoms:

$$F_T = \sum_N^N (f_j) + \sum_A (f_j' + i f_j'') = F_N + F_A$$

as illustrated in Fig 6A, where black vector $F_N$ is the total component of $N_N$ normal atoms, the red vector $F_A$ represents the normal scattering component of the $N_A$ anomalous scatterers (based on the real part of atomic form factors $f_j'$), the pink vector $F''$ corresponds to the purely anomalous contribution $f''$ of the $N_A$ anomalous scatterers. The direction of the $F''$ vector is always rotated with respect to the direction of the $F_A$ vector by $-\pi/2$, if all anomalous atoms are of the same kind. The blue $F_{T''}$ vector represents the total, observable structure factor of the reflection $F(hkl)$. 

| Table 5. Hydrogen bond lengths (Å) between Se atoms and amide nitrogen atoms. Hydrogen bonds within the same $3_1$ columns are in bold. The uncertainties of these distances are ±0.005 Å. |
|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|
| Se_1 | Se_2 | Se_3 | Se_4 | Se_5 | Se_6 | Se_7 | Se_8 | Se_9 |
| N1_1 | 3.447 | - | - | 3.628 | - | - | 3.655 | - |
| N2_1 | 3.474 | - | - | 3.697 | - | - | - | - |
| N1_2 | - | 3.472 | - | - | - | - | 3.574 | - |
| N2_2 | - | 3.454 | - | - | - | - | 3.566 | - |
| N1_3 | - | - | 3.488 | - | - | - | - | - |
| N2_3 | - | - | 3.470 | - | - | 3.426 | - | - |
| N1_4 | - | 3.597 | - | - | - | - | 3.549 | - |
| N2_4 | - | - | - | 3.614 | - | - | 3.670 | - |
| N1_5 | - | - | - | 3.519 | - | - | 3.482 | - |
| N2_5 | - | - | - | - | - | 3.537 | - | 3.711 |
| N1_6 | 3.618 | - | - | 3.574 | - | - | 3.748 | - |
| N2_6 | 3.718 | - | - | 3.619 | - | - | - | - |
| N1_7 | - | 3.682 | 3.713 | - | - | - | - | - |
| N2_7 | - | 3.758 | 3.774 | - | - | - | - | - |
| N1_8 | - | - | - | 3.670 | 3.557 | - | - | - |
| N2_8 | - | - | - | - | - | 3.498 | - | - |
| N1_9 | - | - | - | 3.684 | - | - | 3.602 | - |

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Fig 4. (A) Projection of the SeUrea unit cell along the c-axis with all positions of Se atoms marked in three colors. The red color corresponds to Se atoms within the lowest z-layer, the green color to Se atoms within the layer higher by c/3 and the blue color to Se atoms again higher by c/3. The crystallographic 3, axes are presented in black, the approximate, non-crystallographic 3-fold axes of various kind and the corresponding R3 cell are shown in grey. (B) The selenium atom positions presented in the approximate P3,12 unit cell, with 1.33 Å of the rms difference from ideally symmetric constellation of this space group. The crystallographic 3, axes are in black and the approximate two-fold axes in grey. (C) The idealized approximate symmetry elements of the R32 space group relate the positions of selenium atoms within the rms deviation of 1.61 Å.

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If a large number of atoms are spread in the unit cell randomly, as was e.g. assumed in the classic derivation of the Wilson probability distribution of reflection intensities [13] and in many other crystallographic theories, the atomic contributions form a random walk of small vectors in the Argand diagram of Fig 6A. The rms distance after $N$ random unit steps of length $d$ is $d \sqrt{N}$ [14]. Hence, the large vectors in Fig 6A can be approximated as follows:

$$|F_N| = f_N \sqrt{N_A}, |F_{A}| = (f_A' + f_A) \sqrt{N_A}, |F''| = f_A'' \sqrt{N_A}.$$ 

In the usual practice of phasing macromolecular crystal structures by the Multi- or Single-wavelength Anomalous Diffraction (MAD or SAD, [15]) methods, the great majority of the structure is built by the non-anomalous atoms and the normal contribution of the anomalous atoms is small, a situation which is presented in Fig 6B. The blue vector $F^+$ is the total, measurable amplitude of reflection $F(hkl)$, and the vector $F^-$ is (the complex conjugate of) its Friedel mate $F(-h-k-l)$ (the original $F^-$ reflection has a negative phase and would lie below the abscissa axis). It is evident that the largest measured Bijvoet difference $\Delta F^\pm$ occurs when the red vector $F_A$ is perpendicular to the total (averaged for both Friedel mates) green vector $F_T$, i.e. when $|\alpha_A - \alpha_N| = \pi/2$, or $\sin\gamma = F_A / F_N$. The average measured Bijvoet difference $\Delta F^\pm$ corresponds to the average projection of the pink vector $2F''$ on the direction of $F_T$, so that $<\Delta F^\pm> = 4F''/\pi$, since $<|\cos\alpha|> = 2/\pi$.

The effective Bijvoet ratio, is usually estimated from the following formula given by Hendrickson [15,16]:

$$\text{rms}(\Delta F^\pm)/\text{rms}(F_T) \approx \sqrt{N_A/2N_T} \times (2f'' / Z_{\text{eff}})$$

where $Z_{\text{eff}}$ of 6.7 is the number of electrons in the “average” protein atom.

Taking into account the characteristics of the random walk and the fact that $<|\cos\alpha|> = 2/\pi$, not 1/2 assumed by Hendrickson, the formula should be modified to:

$$\text{rms}(\Delta F^\pm)/\text{rms}(F_T) \approx \sqrt{N_A/2N_T} \times (4f'' / \pi Z_{\text{eff}}) \approx 0.2f'' \sqrt{N_A/2N_T}$$

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**Fig 5.** The correlation coefficient between the anomalous differences in the two randomly selected and merged half-data sets ($CC_{anom}$) versus resolution.

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Fig 6. Vector representation of the anomalous scattering effects. (A) The contributions of individual normal and anomalous atoms to the total structure factor. (B) The case of majority of normal atoms and a few anomalous atoms, $F_N > F_A$. (C) The case with majority of the anomalously scattering atoms $F_A > F_N$ as in the crystal of SeUrea. See the text for explanation of all vector relations.

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However, the situation is different if the anomalous atoms constitute the majority of the total structure, as is the case in the crystal of SeUrea, illustrated in Fig 6C. Here, the red vector $\mathbf{F}_A$ is much larger than the black vector $\mathbf{F}_N$, and it can never be perpendicular to the total vector $\mathbf{F}_T$, so that the Bijvoet difference $\Delta F$ will never reach the maximum value of $2F'$. The maximum projection of $2F'$ onto $\mathbf{F}_T$ occurs when $\mathbf{F}_N$ is perpendicular to $\mathbf{F}_T$, and the angle $\gamma$ is in maximum, with $\sin \gamma_{\text{max}} = \mathbf{F}_N/\mathbf{F}_A$, and then $(\Delta F')_{\text{max}} = 2F' \sin \gamma_{\text{max}}$. The average $\langle \Delta F' \rangle = 2F' \int \sin \gamma \, d\gamma$ from 0 to $\gamma_{\text{max}} = 2F'/\sqrt{1-(\mathbf{F}_N^2/\mathbf{F}_A^2)}$. It is evident that if the structure consists of only anomalous atoms of the same kind, the anomalous Bijvoet differences vanish completely [17,18], although the measured amplitudes depend on the X-ray wavelength through the changing $f$ values.

For SeUrea with $\mathbf{F}_N^2 = 6^2 + 2 \times 7^2 = 134$ and $\mathbf{F}_A^2 = 1156$, the $\Delta F'/2F'$ factor is equal to 0.06, meaning that the anomalous signal is diminished more than 10 times in comparison with the cases when $\mathbf{F}_A < \mathbf{F}_N$, and the $\Delta F'/2F'$ factor is $2/\pi = 0.64$. Nevertheless, the diffraction data measured for SeUrea contain a significant amount of anomalous signal with the averaged $\langle \Delta F' \rangle / \langle F \rangle = 4.5\%$. That value is much higher than what is available in many macromolecular diffraction data sets, where often the anomalous signal of sulfur smaller than 1\% is enough to solve structures of native proteins by the SAD approach. It was, however, not possible to locate the positions of selenium atoms in the SeUrea structure from their anomalous signal by the SHELXD program [19], in spite of many attempts. This structure can be, of course, easily solved by direct methods from the native data.

The failure of the SAD approach is due to the fact that the measured Bijvoet differences do not correlate effectively with the amplitudes $\mathbf{F}_A$ representing diffraction of the anomalous atoms. The phase estimation in the SAD and MAD methods is based on the fact that (for the largest anomalous differences) the total reflection phases are larger by about 90° than the anomalous substructure phases. In case of SeUrea this is not true, since all substructure phases are close to the total phases, as is evident from Fig 6B. The anomalous difference map calculated in a standard way, i.e. with phases smaller by 90° from the total phases will not identify the correct positions of anomalous atoms.

Conclusions

The high resolution diffraction data obtained for selenourea from the synchrotron source provided the structure about ten times more precise than available previously. The accurate values of bond lengths and angles as well as hydrogen bonds in SeUrea can serve as geometrical restraints used in the refinement of macromolecular structures solved with SeUrea as a phasing vehicle. The structure of SeUrea revealed a high degree of pseudosymmetry and some unusual anomalous diffraction properties. The high proportion of anomalously scattering selenium atoms in the structure precluded the use of the standard SAD phasing procedures, suggesting that too many anomalous scatterers in the structure create "embarras de richesces", instead of making the structure solution easier. However, when SeUrea is soaked into crystals of macromolecules [5], the selenium atoms constitute only a small fraction of the total structure and their full anomalous scattering potential can be utilized for estimation of phases by the standard SAD or MAD methods.

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