Experimental Phasing with SHELX C/D/E

CCP4 / APS School Chicago 2017
22nd June 2017
1 - The Phase Problem
The Crystallographic Phase Problem

1. Crystal diffraction yields intensities $I(hkl)$, and thus structure factor amplitude $|F(hkl)| \propto \sqrt{I(hkl)}$

2. Model building requires a map, i.e. $\rho(x,y,z) = \sum_{h,k,l} |F(hkl)| e^{i\phi(hkl)} e^{-2\pi i(hx+ky+lz)}$

3. **Solving a structure** = determination of the phase angles $\phi(hkl)$ good enough to create an interpretable map

4. Small molecule crystallography: Problem mostly solved with direct method

5. SHELXD solves small molecule data at $d < 1.2\text{Å}$

6. SHELXD solves the substructure even at $d = 5\text{Å}$ (and worse) because substructure atoms are still resolved
The Substructure

- Coordinates of anomalous scatterers
- Anomalous difference

$$\left| F^+(hkl) - F^-(hkl) \right| \approx |F_{\text{sub}}(hkl)|$$

corresponds to small molecule data set
- Shelxd: solve substructure with *direct methods*
- Harker Construction: Expand phases to full data set
SHELX C/D/E [1] in Context with Structure Solution

DIALS, XDS, MOSFLM... shelxc shelxd shelxe REFMAC5 / COOT...

Data Processing Anomalous Differences Substructure Solution Phasing & Density Modification Building & Refinement

Molecular Replacement

22nd June 2017
2 - Using SHELX C/D/E
Graphical User Interface HKL2MAP

http://webapps.embl-hamburg.de/hkl2map/
Most shelx programs issue “short” usage instruction when called without an argument.

tg@slartibartfast:~$ shelxc

++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
+ SHELXC - Create input files for SHELXD and SHELXE - Version 2013/2 +
+ Copyright (c) George M. Sheldrick 2003-13 +
+ Started at 13:59:57 on 10 Jun 2014 +
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++

SHELXC reads a filename stem (denoted here by ‘xx’) on the command line
plus some instructions from ‘standard input’. It writes some statistics to
‘standard output’ and prepares the three files needed to run SHELXD and
SHELXE. SHELXC can be called from a GUI by a command line such as:

shelxc xx <t

which would read the instructions from the file t, or (under most UNIX
systems) by a simple shell script that includes the instructions, e.g.

shelxc xx <<EOF
CELL 49.70 57.90 74.17 90 90 90
SPAG P212121
SAD elastase.sca
FIND 12
<<EOF
sheld x xx_fa
shelxe xx xx_fa -s0.37 -m20 -h -b
shelxe xx xx_fa -s0.37 -m20 -h -b -i

More information including tutorials available at http://shelx.uni-goettingen.de/SHELX/.
Shelxc Data Preparation: Keywords

shelxc can be used for five different phasing scenarios:

- **SAD**
  - SAD
  - (NAT)

- **SIRAS**
  - SIRA
  - NAT

- **MAD**
  - (NAT)
  - PEAK
  - INFL
  - LREM
  - HREM
  - BEFORE
  - AFTER
  - (NAT)

- **SIR**
  - SIR
  - NAT

- **RIP**
  - BEFORE
  - AFTER
  - (NAT)

Each keyword takes the filename of the corresponding integrated dataset.
Running shelxc

1. Create input command file `shelxc.inp` with text editor

```
CELL 49.70  57.90  74.17  90.000  90.000  90.000
SPAG P212121
FIND 12
NTRY 100
SFAC S
SAD elastase.sca
```

2. `shelxc mysad < shelxc.inp`
Shelxc Output Files

The command “shelxc mysad < shelxc.inp” creates three files:

**mysad_fa.ins** Text file with instructions for shelxd

**mysad_fa.hkl** Artificial substructure data set from which shelxd determines substructure coordinates. Each line contains

\[ h, k, l, |F^+(hkl)| - |F^-(hkl)|, \alpha \]

\( \alpha \) is not used by shelxd, but by shelxe to calculate an initial phase estimate for the protein structure as

\[ \phi_T(hkl) = \phi_A(hkl) + \alpha(hkl) \]

MAD/SIRAS: exact \( \alpha \); SIR or SAD: rough estimate of \( \alpha \)

\( \phi_A \) is the phase angle calculated from the substructure coordinates determined by shelxd.

**mysad.hkl** native data used by shelxe for phasing and density modification
Shelxc: Resolution Cut-off for Anomalous Signal

85349 Reflections read from SAD file XDS_pk1pk2.HKL

12186 Unique reflections, highest resolution 7.199 Angstroms
141.7 Friedel pairs used on average for local scaling

<table>
<thead>
<tr>
<th>Resl.</th>
<th>Inf.</th>
<th>16.01</th>
<th>12.71</th>
<th>11.10</th>
<th>10.09</th>
<th>9.36</th>
<th>8.81</th>
<th>8.37</th>
<th>8.01</th>
<th>7.70</th>
<th>7.43</th>
<th>7.20</th>
</tr>
</thead>
<tbody>
<tr>
<td>N(data)</td>
<td>1149</td>
<td>1124</td>
<td>1115</td>
<td>1099</td>
<td>1108</td>
<td>1102</td>
<td>1126</td>
<td>1078</td>
<td>1091</td>
<td>1122</td>
<td>1072</td>
<td></td>
</tr>
<tr>
<td>Chi-sq</td>
<td>1.30</td>
<td>1.22</td>
<td>1.24</td>
<td>1.27</td>
<td>1.37</td>
<td>1.55</td>
<td>1.58</td>
<td>1.62</td>
<td>1.49</td>
<td>1.35</td>
<td>1.35</td>
<td>1.43</td>
</tr>
<tr>
<td>&lt;I/sig&gt;</td>
<td>41.7</td>
<td>28.7</td>
<td>24.4</td>
<td>22.1</td>
<td>16.5</td>
<td>13.3</td>
<td>9.2</td>
<td>7.4</td>
<td>5.3</td>
<td>3.8</td>
<td>2.2</td>
<td></td>
</tr>
<tr>
<td>%Complete</td>
<td>98.9</td>
<td>99.9</td>
<td>100.0</td>
<td>100.0</td>
<td>100.0</td>
<td>100.0</td>
<td>100.0</td>
<td>100.0</td>
<td>100.0</td>
<td>98.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;d&quot;/sig&gt;</td>
<td>4.82</td>
<td>2.79</td>
<td>1.90</td>
<td>1.53</td>
<td>1.26</td>
<td>1.04</td>
<td>1.00</td>
<td>0.89</td>
<td>0.85</td>
<td>0.80</td>
<td>0.71</td>
<td></td>
</tr>
<tr>
<td>CC(1/2)</td>
<td>94.8</td>
<td>81.8</td>
<td>65.2</td>
<td>53.8</td>
<td>39.6</td>
<td>25.0</td>
<td>23.6</td>
<td>9.3</td>
<td>5.3</td>
<td>-6.6</td>
<td>-14.4</td>
<td></td>
</tr>
</tbody>
</table>

- anomalous signal where $CC(1/2) > 30\%$

- anomalous signal where $<d"/sig> > 1.3$

- $CC > 30\%$ usually more reliable than $<d"/sig> > 1.3$
3 - Shelxd
Shelxd — Finding the Substructure

shelxd mysad_fa

reads the “substructure data” mysad_fa.hkl and its instructions from mysad_fa.ins. The most important entries in mysad_fa.ins:

- **SFAC SE**  atom type to look for
- **FIND 12**  expected number of substructure atoms, should be within 20 % of the actual number (try several for e.g. a soak where the number is not known)
- **SHEL 999 3.3** resolution limits of the **anomalous signal**. High resolution limit can be critical, but the default of $d_{\text{min}} + 0.5$ Å works well in many cases.
- **NTRY 10000** number of trials.
**Shelxd Output**

While **shelxd** runs, the best solution is written to **mysad_fa.res** which contains the substructure coordinates in fractional coordinates and which is later read by shelxe.

---

```
REM Best SHELXD solution:   CC  60.74    CC(weak) 49.22    CFOM 109.96
TITL  mysad_fa.ins MAD in C2
CELL  0.98000  109.02  61.75  71.74  90.00  97.08  90.00
LATT -7
SYMM -X, Y, -Z
SFAC SE
UNIT  192
SE01  1  0.758774  0.508636  0.246391  1.0000  0.2
SE02  1  0.792908  0.398262  0.138903  0.8845  0.2
[...]
SE10  1  0.925819  0.231575  0.191291  0.5569  0.2
SE11  1  0.495239  0.183609  0.416278  0.5352  0.2
SE12  1  0.643097  0.029221  0.210653  0.4897  0.2 <---
SE13  1  0.811539  0.048553  0.227752  0.1453  0.2 <---
SE14  1  0.600281  0.156860  0.149628  0.0764  0.2
HKLF  3
END
```

The sixth column contains the occupancy of the corresponding atom. A sharp drop (here between SE12 and SE13) is a promising sign of a correct solution. The correlation coefficient (CC and CCweak) in the first line measures the reliability of the solution.

For SAD, a CC of more than 30 % is a safe sign of a correct solution, for MAD the limit is about 40 %.
Shelxd Output

SHELXD is very robust. Attention should be paid to

1. The **resolution** at which the data are truncated, *e.g.* where the internal CC (CC1/2) between the signed anomalous differences of two randomly chosen reflection subsets falls below 30%.

2. The **number of sites** requested should be within about 20% of the true value.

3. In the case of a soak, the rejection of sites on **special positions** should be switched off.

4. For S-SAD, DSUL (**search for disulfides**) can be very useful.

5. In difficult cases it may be necessary to run more trials (say 50000).
Fine tuning SHELXD substructure solution

SHELXD is very fast and robust, but achieves this with the help of drastic assumptions.

In borderline cases it may be worth using the LLG (log likelihood gain) to distinguish substructure solutions, e.g. using the programs SHARP, CRANK2 or PHASER. For details see:


These programs could also be used to refine and extend the heavy atom substructure before density modification and poly-Ala tracing with SHELXE. In general LLG-based methods require more detailed information (e.g. which elements are present) than SHELXC/D/E, and they tend to be slower.
4 - Shelxe
Shelxe: Phasing, Density Modification, Model Building

No .ins-tructions file. All parameters provided as command line options after data file names.

A typical and one of the most simple command line could be

```
shelxe mysad mysad_fa -s0.65 -h -a
```

```
mysad read native data mysad.hkl
```

```
mysad.fa read angle estimate for $\alpha$ from mysad_fa.hkl, substructure coordinates from mysad_fa.res (the shelxd output)
```

```
-s0.65 Assume a solvent content of 65%. It should be reasonably well estimated.
```

```
-h substructure atoms present in native data mysad.hkl
```

```
-a run 5 (default) cycles of poly-ALA autotracing.
```
**Shelxe \(-i\): Inverted Substructure**

It is impossible to distinguish the substructure from its enantiomorph with the anomalous data and there is a 50% chance that the coordinates in \texttt{mysad_fa.res} are inverted w.r.t. the correct substructure.

Therefore shelxe must always be run **twice**

- with the **direct substructure**
- with the **inverted substructure**, \emph{i.e.} with the same options as the direct hand \emph{plus} the switch \(-i\). This inverts the hand and takes care of everything necessary
  - inversion of screw axes, P4\(_1\) to P4\(_3\)
  - off-axis inversion for \(I4_1\) (1-x, 1/2-y, 1-z); \(I4_1\)\_2 (1-x, 1/2-y, 1/4-z); \(F4_1\)\_32 (1/4-x, 1/4-y, 1/4-z)
- output files are automatically amended by \(-i\) to distinguish the two runs.

**N.B.** if the inverted hand turns out to be the correct hand, your **space group may change** - \emph{e.g.} in the **presence of screw axes**. Keep this in mind when you convert your native data to \emph{e.g.} mtz-format!
Caveat: Substructure Resolution

- “Normal” macromolecular structure: Determine atom positions even at e.g. 5 Å resolution because of restraints.

- Substructure unrestrained

⇒ coordinates only know within resolution of anomalous signal, often much worse than 3 Å

Way out:

1. e.g. Sharp improves substructure coordinates before density modification

2. “substructure recycling” with shelxe
Shelxe: Substructure Recycling

- Better substructure = better starting phases = better map
- **Caveat**: If the inverted structure turns out to be the correct hand (*i.e.* mysad_i.hat from the -i-run of shelxe), the second run of shelxe must be run **without** the -i switch:
Shelxe: Did it work?

Criteria to tell if phasing worked:

1. Correct hand shows better **Contrast**, especially at early cycles of density modification.

2. Correct hand has higher **map correlation coefficient** throughout resolution range:

   \[
   \begin{array}{cccccccccccc}
   d & \text{inf} & 4.66 & 3.70 & 3.23 & 2.93 & 2.72 & 2.56 & 2.43 & 2.33 & 2.24 & 2.15 \\
   <\text{mapCC}> & 0.626 & 0.795 & 0.775 & 0.754 & 0.819 & 0.804 & 0.756 & 0.694 & 0.620 & 0.582 & \text{direct} \\
   <\text{mapCC}> & 0.810 & 0.877 & 0.845 & 0.844 & 0.874 & 0.856 & 0.840 & 0.830 & 0.839 & 0.809 & \text{inverse} \\
   \end{array}
   \]

3. A reasonable poly-ALA trace (average 10 residues per chain) and a **CC > 25%**

When using the auto-tracing option (–a) in shelxe, the first two figures (contrast/ mapCC) become meaningless, but in this case the poly-ALA trace is much more conclusive.
Shelxe: Structure Solved?

Indicators from shelxe:

<table>
<thead>
<tr>
<th>TITLE</th>
<th>elastase.pdb</th>
<th>Cycle</th>
<th>CC</th>
<th>226 residues in 4 chains</th>
</tr>
</thead>
<tbody>
<tr>
<td>TITLE</td>
<td>elastase_i.pdb</td>
<td>Cycle</td>
<td>CC</td>
<td>61 residues in 7 chains</td>
</tr>
</tbody>
</table>

1. CC>25%

2. average chain length > 10 (here: 56.5 vs. 8.7)

3. jump in CC over many cycles (e.g. with $-a50$)
Shelxe: Structure Solved?

Coot reads `mysad.pdb` (poly-ALA trace) and `mysad.phs` (map).

Elastase SAD tutorial
SHELXE: Current and Future Developments

- SHELXE started with -x and a reference PDB file name.ent is present: the mean phase error is output at various stages. The necessary origin shift is determined on the fly.

- If -h is also set, the program finds the atom in the reference file nearest to each heavy atom site. This is particularly useful for checking the substructure.

- The density modification has been improved for SAD phasing. For 20 test structures the mean phase improvement after the first round of density modification was 4.6°.

- These improvements are already in the current distributed version. In addition, SHELXC is being adapted to handle multiple SAD datasets and a major rewrite of SHELXE is in progress.
References and Further Reading


2. Visit the shelx web page for documentation, tutorials, etc.: http://shelx.uni-goettingen.de

Availability

SHELX is available free for academic use via the SHELX homepage http://shelx.uni-goettingen.de/. Extensive documentation and many links to useful programs may also be found there. SHELX C/D/E are also distributed along with CCP4.