# TRAFO - Coordinate Transformations 

UsER MANUAL

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## 1 Getting Started

This chapter covers the following topics:

| Introduction | An introduction to Trafo |  |
| :--- | :--- | :--- |
| Overviews | An overview of some of the common tasks and actions |  |
| Revision History | Summary of changes versus previous versions |  |
| Known Problems | Summary of known problems |  |
|  | Acknowledgements | Credits for important contributions etc. |

### 1.1 Introduction

Trafo is a program to perform coordinate transformations useful in combination with the analysis of single-crystal NMR spectra, as well as to assist in some related procedures such as diagonalizations of tensors. Although Trafo will often deal with crystallographic data, it is not smart in that respect; it is a simple tool and not an expert system. If you are looking for powerful software to deal with crystallographic questions, consider Platon or Mercury, both available freely.
Trafo is part of a simulation package developed during my stay at the Department of Chemistry, Dalhousie University, Halifax, Canada. The initial version has been written in C++ using Borland C++ 4.52 and has been developed to run on IBM-compatible personal computers under the MSWindows environment in the 16-bit subsystem. For the 32 -bit version, Borland C++5.0 has been used initially. The current version has been produced using Microsoft Visual C++ 2008 Express Edition.

### 1.2 Overview

The actions selected by the menu commands operate

- either directly on the content of the edit window - the default if there is anything available
- or, if the edit window is empty (you can select File-New (p. 8) to clear the window), on an input file, the name of which is requested from the user.
The output is almost always directed to an output file, the name of which is also asked for. Note that currently the edit window has a limited capacity of about 64 KB , which is sufficient for most operations, but can be exceeded easily after application of Action-Generate Block (p. 16); there ís no such low limit on file oriented operations.
The format of the input data should conform to the format expected by Alchemy for crystal data files, although Trafo is a little more flexible in this respect. I have chosen this file format because in the 90ties, when I started the development of Trafo, we used the molecular modeling programs Alchemy II and Alchemy 3 to visualize molecules and the results of single crystal NMR data. Another advantage of this file format is its simplicity. An example is shown in Fig. 1.1. Several of the examples shown later in this document use this example as starting point, hence you could copy and paste the data in the figure into Trafo and save them for perusal later.
Crystal data files to be recognized by Alchemy must have the extension ". CRY". All operations are line-oriented, i.e., each line of the ASCII-type data forms one data record. Basically, there are three different types of data records, each of them with some restrictions as to their format. The three data types are:
- Crystal axis: crystal system data

```
7.502 7.502 7.546 90.00 90.00 90.00
P 0.000 0.000 0.000
N 0.000 0.000 0.500
0
H 0.014 0.111 0.573
H 0.227 0.146 0.122
/* crystal structure of NH4H2PO4 */
/* space group: I-42d (No. 122) */
/* neutron diffraction data */
/* Tenzer, L.; Frazer, B.L.; Pepinsky, R. */
/* Acta Cryst. 1958, 11, 505 */
```

Figure 1.1: Example of an Alchemy crystal data file, here for ammonium dihydrogen phosphate (ADP), $\mathrm{NH}_{4} \mathrm{H}_{2} \mathrm{PO}_{4}$ [1]. You can save the data as ADP_01_fract_coord.cry if you want to repeat some of the example actions shown later.

- Comment lines
- Regular data lines

One disadvantage of the Alchemy crystal data file format is that most other programs do not accept this format. Most of them will be able to deal with Alchemy ".MOL" files, but not the ".CRY" files. However, it is not very difficult to convert the ".CRY" files into a format that is recognized by a wider variety of programs, the so called XYZ format. You can always use Trafo to convert your fractional coordinates into Cartesian coordinates. In the resulting file, simply replace the first line with the Cartesian crystal axes data by the number of atoms in the file, add a second line with a title, and keep the rest with the Cartesian coordinates (an example of a XYZ file is shown in Fig. 2.4). Such XYZ files can be read into Jmol or Mercury, for example.
On this type of data, one can perform transformations, called Actions in Trafo.

Links to some of the software mentioned:

- Jmol: http://www.jmol.org
- Mercury: http://www.ccdc.cam.ac.uk/mercury


## - References

(1) Tenzer, L.; Frazer, B.L.; Pepinsky, R. Acta Crystallogr. 1958, 11, 505, DOI: 10.1107/S0365110X58001389

### 1.2.1 Crystal Axis System Data

The data line with crystal axes data describes the geometry of the unit cell of a crystal and consists of six floating point numbers, $a, b, c, \alpha, \beta, \gamma$, where:
$a, b, c: \quad$ length of the respective crystal axis in Angstrøm
$\alpha, \beta, \gamma$ : angle between pairs of crystal axes ( $\alpha$ is the angle between $b$ and $c, \beta$ between $a$ and $c$, and $\gamma$ between $a$ and $b$ ).

For Cartesian coordinates, this line will read:
1.01 .01 .090 .090 .090 .0

Note: This data line is required for many operations and has always to be the first line in this case. For actions where this line is not required, it is optional but, if present, it also has to be the first line!

### 1.2.2 Comment Lines

Comment lines can be added to the data in order to assist later in identifying the data set. In Alchemy, comment lines can only be added to the end of the data set, but in Trafo they may appear anywhere (except for the first line in cases where crystal axis system data are required). Comment lines are written in the same format as comment lines in the C programming language, as the following example illustrates:

```
/* this is a C-type comment line */
```

In order to qualify as a comment line, the first two characters of the line have to be a slash (/) and an asterisk (*).

### 1.2.3 Regular Data Lines

Each line with regular coordinate data consists of four data entries, all of them are required. The general format will be:

LABEL x.xxxx -y.yyyy z.zzzz
where:

LABEL a label of at least 1 and up to 6 characters describing the atom or data type (no spaces or other whitespace characters, e.g. tabs, within this name are allowed, but underscores etc.); for use with Alchemy, the label should specify a valid Alchemy atom type, but Trafo does not care.
$x, y, z$ the $x, y, z$ coordinates as floating point numbers in free format
A special type of data line will be recognized by Trafo, which is a line describing the origin of the coordinate system:

OR x.xxx y.yyy z.zzz
where Trafo doesn't really care about the coordinates (right now), but this line shall usually be neglected (the position and number of such lines doesn't matter either).
The purpose of this origin is to serve as starting point for the display of principal axes systems, e.g., of chemical shift tensors, whereby the directions of the principal components were indicated as dummy atoms, e.g., d11, d22, d33, that are $1 \AA$ away from the origin.

### 1.3 Revision History

This page summarizes the changes made compared to previous versions of Trafo. Experienced users may use this information as a quick update on new program features.

### 1.3.1 New in Version 2.5 .8 (19.05.2021)

- Bug fix: newer versions of Adobe Acrobat (Reader) should be supported for context sensitive help. (I am installing Acrobat only occasionally, hence I am not always able to check.)
- (Bug) fix: the nuclear tables had a wrong magnetogyric ratio for ${ }^{207} \mathrm{~Pb}$; see Dipolar Second Moment (p. 29)


### 1.3.2 New in Version 2.5.7 (14.05.2015)

- new feature: the help file contains a lot of examples that illustrate the use of Trafo.
- new feature: the help file is now in PDF format because it is suitable for both online viewing as well as printing. To view or print this documentation, any PDF viewer should work. Also, most viewers should be able to deal with the hyperlinks that cross-link topics in this help file. However, opening a PDF file at a specified position is more complicated. Therefore, not every PDF viewer will work with Trafo to display context sensitive help. This matter is aggravated by the fact that every viewer has its own mechanism! Currently, context sensitive help should work with the following PDF viewers:
- Adobe uses with its Acrobat Reader series of products a mechanism called Dynamic Data Exchange (DDE) to open a PDF file at a specific position. For ages, the name of the DDE server has been acroview. However, with the introduction of Acrobat Reader X this tradition has been broken on purpose, breaking many applications that rely on this mechanism. Trafo should be able to work with older versions of the Acrobat Reader as well as with the acroviewR10 DDE server of Adobe Reader X. Because I have been quite dissatisfied with the Adobe products from Reader 5 onwards, I am using a different PDF viewer and will not always check whether Acrobat is still working!
- My preferred PDF viewer is the free PDFXChangeEditor, the successor of PDFXChangeViewer by Tracker Software Products, Ltd. (http://www.tracker-software.com), more powerful and reliable than the Acrobat Reader. The opening of a PDF file at a specific position is achieved via command line parameters.
- Another viewer used by people working with $\mathrm{IATEX}_{\mathrm{E}}$ is the Sumatra PDF Viewer (http: //blog.kowalczyk.info/software/sumatrapdf), also supported by Trafo.
- Other PDF viewers such as FoxIt or NitroPDF do not support DDE or command line parameters, as far as I know.
- bug fix: a few superfluous new lines produced in the output files got eaten now.


### 1.3.3 New in Version 2.5 .3 (21.01.2009)

- converted project to Microsoft Visual C++ 2008 Express Edition
- added diagonalization functions for single crystal NMR
- some minor bug fixes


### 1.3.4 New in Version 2.4.7 (30.04.2004)

- some minor bug fixes


### 1.3.5 New in Version 2.4.3

- this is the first release with a 32 bit version; some internal changes
- comment lines are now copied to output file


### 1.3.6 New in Version 2.3.2

- added calculation of dipolar second moments
- better error reporting


### 1.3.7 New in Version 2.2.10

- Added calculation of normals.


### 1.4 Known Problems

None right now.

### 1.5 Acknowledgements

This section covers the following topics:

| Credits | Credits for important contributions |
| :--- | :--- |
| Trademarks | Trademark acknowledgements |
| Copyright | The obligatory copyright statement |
| Disclaimer | The obligatory disclaimer message |

### 1.5.1 Credits

- Microsoft for providing Visual C++ 2008 Express Edition for free
- Jordan Russell for making Inno Setup available (http://www.jrsoftware.org/)
- Jochen Kalmbach for demonstrating how to statically link against the Microsoft CRT and thus get rid of VCREDIST_X86.EXE (http://blog.kalmbach-software.de)
- "chicks" for demonstrating in his pdfp PDF tools how to establish Dynamic Data Exchange (DDE) with Adobe Acrobat (Reader) (http://www . esnips . com/web/PDFTools, website not available anymore).
Here are some "substitute kudos" pages: http://www.quickpdf.org/forum/printdocument-blank-pages_topic688.html or https://www.techrepublic.com/forums/discussions/want-to-print-pdfs-from-access-form/.
- This manual has been produced using the MiKTEX (http://www.miktex.org) distribution of $\mathrm{LAT}_{\mathrm{E}} \mathrm{X}$ in combination with the TeXstudio editor (http://texstudio. sourceforge.net/).
- Irfan Skiljan's IrfanView (http://www.irfanview.com/) has been used to process bitmapped images, and Inkscape (www.inkscape.org) for dealing with vector graphics.


### 1.5.2 Trademark Acknowledgement

- Microsoft (MS) is a registered trademark and MS-DOS, MS - Word and MS - Windows are trademarks of Microsoft Corporation.
- Alchemy is a copyrighted program and registered trademark of TRIPOS Associates, Inc., a subsidiary of Evans \& Sutherland.
- Other brand and/or product names are used for identification purposes only and are trademarks, registered trademarks or copyrights of their respective owners.


### 1.5.3 Copyright Message

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## 2 Reference

### 2.1 Help on Menu Items

The menu system of Trafo consists of the following pop-up menus:

| - Trafo - 01_ADP_Khan_fract_coord.cry |  |  |  |
| :---: | :---: | :---: | :---: |
| Eile Edit | Action | NMR Help |  |
| 7.4997 | 7.4997 | 7.5494 | 95.0090 .00 |
| P | 0.0000 | 0.0900 | 0.0900 |
| N | 0.0096 | 0.0909 | 0.5090 |
| 0 | $0.08 / 43$ | 0.1466 | 0.1151 |


| Category | Menu Item | Action |
| :---: | :---: | :---: |
| File | File and document management: |  |
|  | New (p. 8) | clears the content of the Trafo edit window |
|  | Open... (p. 8) | read a file into Trafo s edit window |
|  | Save as... (p. 8) | save the content of the edit window to a new file |
|  | Exit (p. 8) | exit Trafo |
| Edit | Editor window actions: |  |
|  | Undo (p. 9) | undo last changes |
|  | Cut (p.9) | cut selection into clipboard |
|  | Copy (p.9) | copy selection into clipboard |
|  | Paste (p. 9) | insert content of clipboard |
|  | Clear (p. 9) | clear content |
|  | Select All (p. 9) | select all content |
| Action | Perform transformations: |  |
|  | Generate: Positions (p. 10) | generate symmetry related sites |
|  | Generate: Block (p. 16) | generate a cube of several unit cells |
|  | Orthogonalize: Axes (p. 17) | transform an oblique axis set into an orthogonal set |
|  | Orthogonalize: Coordinates (p. 19) | transform coordinates from an oblique system into a Cartesian system |
|  | Orthogonalize: Calculate normal (p. 22) | calculate the normal vector |
|  | Rotations: Calculate Euler Angles (p. 23) | calculate Euler Angles from direction cosines |
|  | Rotations: Euler Rotations Forwards (p. 27) | transform from the old system into the new system |
|  | Rotations: Euler Rotations Backwards (p. 28) | transform from the new system into the old system |


|  | Polar angles: Calculate (p. 28) | convert Cartesian coordinates into polar coordinates |
| :---: | :---: | :---: |
| NMR | Perform some NMR relevant calculations: |  |
|  | Dipolar Moment (p. 29) | calculates dipolar second moments for singlecrystal or powder samples |
|  | $\begin{aligned} & \text { Diagonalise (d11 > d22 > } \\ & \text { d33) }(\mathrm{p} .30) \end{aligned}$ | diagonalise a general chemical shift tensor |
|  | Diagonalise (s11<s22< s33) (p. 30) | diagonalise a general magnetic shielding tensor |

### 2.2 File Menu

The File pop-up menu consists of the following items:

| 7. Trafo-01_ADP_Khan_fract_coord.cry |  |  |  |
| :--- | :--- | :--- | :--- |
| File | Edit | Action | NMR | Help

### 2.2.1 File - New

Execution of the command New in the File popup menu clears the edit window. If the current content of the edit window has been modified but not saved, the user has the choice of writing the current content of the edit window to a file. Once the data are gone, they are gone!

### 2.2.2 File - Open File

Execution of the command Open in the File popup menu retrieves the content of a data file into the edit window. If the current content of the edit window has been modified but not saved, the user has the choice of writing the current content of the edit window to a file. Once the data are gone, they are gone!

### 2.2.3 File - Save File As

Execution of the command Save as in the File popup menu saves the content of the edit window, if available, into a data file. To prevent Trafo from adding the extension .cry, enclose your file name in double quotes.
N.B. It overwrites existing data files without a warning!

### 2.2.4 File - Exit

Execution of the command Exit in the File popup menu quits Trafo. If the current content of the edit window has been modified but not saved, the user has the choice of writing the current content of the edit window to a file. Once the data are gone, they are gone!

### 2.3 Edit Menu

The Edit Menu provides various commands which act on the content of the edit window. Depending on previous actions, not all commands might be available at a given time. In their functionality, the commands are typical of a Windows application.

$$
\text { _uly } 8,2021
$$

The Edit pop-up menu consists of the following items:


### 2.3.1 Edit - Undo

This command restores the file in the current window to the way it was before your most recent edit or cursor movement. Undo inserts any characters you deleted, deletes any characters you inserted, replaces any characters you overwrote, and moves your cursor back to a prior position.
This command is available only if an Edit window is currently active and there is something to undo.

### 2.3.2 Edit - Cut

This command removes the selected text from your document and places the text in the Clipboard. You can then choose Edit|Paste to paste the cut text into any other document (or somewhere else in the same document). You can paste the cut text as many times as you want until you choose Edit|Cut or Edit|Copy.
This command is available only if an Edit window is currently active and text has been marked for selection.

### 2.3.3 Edit - Copy

This command leaves the selected text intact and places an exact copy of it in the Clipboard. To paste the copied text into any other document, choose Edit $\mid$ Paste.
To copy text from a Help window, display the text you want to copy, then select Edit|Copy. The entire contents of the window is copied to the Clipboard.
This command is only available if an Edit window is currently active and text has been marked for selection.

### 2.3.4 Edit - Paste

This command inserts the contents of the Clipboard into the current window at the cursor position. This command is available only if an Edit window is currently active and there is something to paste.

### 2.3.5 Edit - Clear

This command removes the selected text but does not copy it to the Clipboard. This means you cannot paste the text as you could if you had chosen Edit|Cut or Edit|Copy.
Although you cannot paste the cleared text, you can undo the Clear command with Edit|Undo.
Clear is useful if you want to delete text, but you do not want to overwrite text held in the Clipboard. This command is available only if an Edit window is currently active and text has been marked for selection.

### 2.3.6 Edit - Select All

This command selects the entire contents of the active Edit window. You can then use Edit|Copy or Edit|Cut to copy it to the Clipboard, or perform any other editing action.
This command is available only if an Edit window is currently active.

### 2.4 Actions Menu

The Actions Menu combines all the commands to perform transformations on input data. Currently, the following actions can be performed:


### 2.4.1 Action - Generate Positions

The Generate Positions command from the Actions|Generate menu allows to create new coordinates (positions) from the existing one by applying symmetry operations.
One possible application is to create all the molecules within the unit cell from the fractional crystal coordinates of one given molecule and applying the generating symmetry operations on them, as detailed in [1]. Also, after generating one complete unit cell, one could create a cube of unit cells by using the Generate Block (p. 16) command from the Actions|Generate menu.


For example, consider the situation illustrated in the figure. The goal is to generate a new molecule which is related to the current one by a twofold screw axis operation. Such a symmetry operation requires the rotation of atomic positions by $180^{\circ}$ about the axis of the screw, followed by a translational element. The detailed procedure together with some examples is explained in the topic symmetry operations (p. 10).

- References
[1] International Tables for X-ray Crystallography, Kynoch Press: Birmingham, England, 1974; Vol. 1.


### 2.4.2 Symmetry Operations

The Generate Positions command from the Actions|Generate menu allows to create new coordinates (positions) from the existing ones by applying symmetry operations. One application is to generate all positions in the unit cell from the content of the asymmetric unit. It calls the dialog box Symmetry Operations. A detailed example at the bottom of this topic illustrates the procedure.
In the dialog box Symmetry Operations, one defines:


- Number of sites to be generated: In order to generate a unit cell, enter the number of molecules (or sites or formula units) as the number of positions. For the first site, apply the identity operation (which is the default symmetry operation). For the following sites, apply the necessary symmetry operations.
- Symmetry operation required to generate the sites: The symmetry operations can be written as combinations of multiplications (Factor) and summations (Shift) applied to the input coordinates, where it is possible that the source of coordinates are interchanged.
The general equation to calculate the new coordinates is:
New $(k)=$ Factor $\times$ Old $(m)+$ Shift
with $k, m=\mathrm{x}, \mathrm{y}, \mathrm{z}$.
It is easier to apply symmetry operations on fractional coordinates rather than Cartesian coordinates.


## - Example: Space Group I42d (No. 122)

For this example, we shall use the fractional coordinates of ammonium dihydrogen phosphate (ADP), $\mathrm{NH}_{4} \mathrm{H}_{2} \mathrm{PO}_{4}$, presented in Fig. 1.1 as a starting point. This example is sufficiently complex to illustrate quite a few points:

- the application of simple symmetry operations;
- how to deal with special positions;
- why it could be necessary to change the source.

The space group symbol I indicates that the crystal structure of ADP is a body centered lattice, therefore there is a so-called centering operation that generates this centered part. Altogether, there are 16 symmetry related general positions, half of them generated by this centering operation. This can be expressed as:

$$
\begin{align*}
(0,0,0 ; 1 / 2,1 / 2,1 / 2)+ & (x, y, z ; \bar{x}, \bar{y}, z ; \bar{y}, x, \bar{z} ; y, \bar{x}, \bar{z} ; \\
& 1 / 2-x, y, 1 / 4-z ; 1 / 2+x, \bar{y}, 1 / 4-z ; 1 / 2+y, x, 1 / 4+z ; 1 / 2-y, \bar{x}, 1 / 4+z) \tag{2.1}
\end{align*}
$$

This generally adopted setting for ADP (at least, the researchers that determined the structure of ADP called it "generally adopted") is obtained by a rotation of $90^{\circ}$ about the $\overline{4}$ axis relative to the setting in International Tables for X-ray Crystallography (1965) for the space group I $\overline{4} 2 d$ [1,2]. If you use the setting found in the International Tables to generate the positions of ADP from the coordinates in Fig. 1.1, you will get a wrong structure!
To generate all 16 general positions of ADP, we will take two steps: in the first step, we will apply the eight operations shown in the second part of Eq. 2.1 and save the result to a file. In this file, we will need to clean up a few positions related to the special positions of P and N before application of the second step, the centering operation.
If you have not done so already, generate your starting file as outlined in the caption of Fig. 1.1. Read this file into Trafo and select Action - Generate Positions. As name for the output file, enter ADP_02_unithalf.cry.
(1) For now, we want to generate eight positions, hence we enter 8 in No. of Positions. For the first position, already queried in this dialog box, the identity operation is used, with a factor of 1 and a shift of zero for all coordinates $X, Y, Z$ (this operation is the default), with the source being $x, y$, and $z$, respectively. This simply copies the input coordinates into the output file. To generate this position, the default looks like:

|  | factor | shift | source |
| :---: | :---: | :---: | :---: |
| $X:$ | 1.0 | 0.0 | $x$ |
| $Y:$ | 1.0 | 0.0 | $y$ |
| $Z:$ | 1.0 | 0.0 | $z$ |

## Click OK.

(2) The second position is related to the first by a two-fold rotation around the $\overline{4}$ axis (the $\overline{4}$ axis generates a two-fold axis), expressed in Eq. 2.1 as $\bar{x}, \bar{y}, z$. To generate this position, enter:

|  | factor | shift | source |
| :---: | :---: | :---: | :---: |
| $X:$ | -1.0 | 0.0 | $x$ |
| $Y:$ | -1.0 | 0.0 | $y$ |
| $Z:$ | 1.0 | 0.0 | $z$ |

(3) The third position, generated by $\bar{y}, x, \bar{z}$, interchanges the sources for $x$ and $y$ :

|  | factor | shift | source |
| :---: | :---: | :---: | :---: |
| $X:$ | -1.0 | 0.0 | $y$ |
| $Y:$ | 1.0 | 0.0 | $x$ |
| $Z:$ | -1.0 | 0.0 | $z$ |

(4) The fourth position, generated by $y, \bar{x}, \bar{z}$, is produced by:

|  | factor | shift | source |
| :---: | :---: | :---: | :---: |
| $X:$ | 1.0 | 0.0 | $y$ |
| $Y:$ | -1.0 | 0.0 | $x$ |
| $Z:$ | -1.0 | 0.0 | $z$ |

5 The fifth position, generated by $1 / 2-x, y, 1 / 4-z$, is produced by:

|  | factor | shift | source |
| :---: | :---: | :---: | :---: |
| $X:$ | -1.0 | 0.5 | $x$ |
| $Y:$ | 1.0 | 0.0 | $y$ |
| $Z:$ | -1.0 | 0.25 | $z$ |

(6) The sixth position, generated by $1 / 2+x, \bar{y}, 1 / 4-z$, is produced by:

|  | factor | shift | source |
| :---: | :---: | :---: | :---: |
| $X:$ | 1.0 | 0.5 | $x$ |
| $Y:$ | -1.0 | 0.0 | $y$ |
| $Z:$ | -1.0 | 0.25 | $z$ |

(2) The seventh position, generated by $1 / 2+y, x, 1 / 4+z$, is produced by:

|  | factor | shift | source |
| :---: | :---: | :---: | :---: |
| $X:$ | 1.0 | 0.5 | $y$ |
| $Y:$ | 1.0 | 0.0 | $x$ |
| $Z:$ | 1.0 | 0.25 | $z$ |

8 The eigth position, generated by $1 / 2-y, \bar{x}, 1 / 4+z$, is produced by:

|  | factor | shift | source |
| :---: | :---: | :---: | :---: |
| $X:$ | -1.0 | 0.5 | $y$ |
| $Y:$ | -1.0 | 0.0 | $x$ |
| $Z:$ | 1.0 | 0.25 | $z$ |

Open the resulting file, ADP_02_unithalf.cry, in Trafo. Because we have applied the symmetry operations on all atoms, including P and N at special positions, we have generated too many N and P positions. There are four formula units of ADP per unit cell, hence our first 8 symmetry operations should generate two formula units: $2 \cdot \mathrm{NH}_{4} \mathrm{H}_{2} \mathrm{PO}_{4}=\mathrm{N}_{2} \mathrm{H}_{8} \mathrm{H}_{4} \mathrm{P}_{2} \mathrm{O}_{8}$ (the protons at oxygen have only an occupation no. of one half). Only one quarter of the P and N positions are required. You can see that P positions 2-4 are the same as the first and 6-8 the same as the fifth, hence can be deleted, leaving two $P$ positions, as required. It is a little more complicated for nitrogen: positions 2,6 are the same as 1 and 5 , while $3,4,7,8$ correspond to 1 and 5 , but shifted by one unit of the $c$ cell axis, therefore they are also superfluous. Clean up the file as outlined and save the result again as ADP_02_unithalf.cry. The result should look like Fig. 2.1.
Now we shall generate the second half of the unit cell by applying the centering operation. With file ADP_02_unithalf.cry in the edit window of Trafo or specifying this file as input file, select Action -

| 7.502 | 7.502 | 7.546 | 90.00 |
| :--- | ---: | ---: | ---: |
| P | 0.000000 | 0.000000 | 90.00 |
| N | 0.000000 | 0.000000 | 0.500000 |
| O | 0.085000 | 0.146000 | 0.115000 |
| H | 0.014000 | 0.111000 | 0.573000 |
| H | 0.227000 | 0.146000 | 0.122000 |
| D | -0.085000 | -0.146000 | 0.115000 |
| H | -0.014000 | -0.111000 | 0.573000 |
| H | -0.227000 | -0.146000 | 0.122000 |
| O | -0.146000 | 0.085000 | -0.115000 |
| H | -0.111000 | 0.014000 | -0.573000 |
| H | -0.146000 | 0.227000 | -0.122000 |
| O | 0.146000 | -0.085000 | -0.115000 |
| H | 0.111000 | -0.014000 | -0.573000 |
| H | 0.146000 | -0.227000 | -0.122000 |
| P | 0.500000 | 0.000000 | 0.250000 |
| N | 0.500000 | 0.000000 | -0.250000 |
| D | 0.415000 | 0.146000 | 0.135000 |
| H | 0.486000 | 0.111000 | -0.323000 |
| H | 0.273000 | 0.146000 | 0.128000 |
| D | 0.585000 | -0.146000 | 0.135000 |
| H | 0.514000 | -0.111000 | -0.323000 |
| H | 0.727000 | -0.146000 | 0.128000 |
| O | 0.646000 | 0.085000 | 0.365000 |
| H | 0.611000 | 0.014000 | 0.823000 |
| H | 0.646000 | 0.227000 | 0.372000 |
| D | 0.354000 | -0.085000 | 0.365000 |
| H | 0.389000 | -0.014000 | 0.823000 |
| H | 0.354000 | -0.227000 | 0.372000 |

Figure 2.1: Alchemy crystal data file for ammonium dihydrogen phosphate (ADP), $\mathrm{NH}_{4} \mathrm{H}_{2} \mathrm{PO}_{4}$, generated from the data in Fig. 1.1, corresponding to one half of the unit cell. Save the data as ADP_02_unithalf.cry if you want to repeat some of the example actions shown later.


Figure 2.2: File "ADP_03_unit_cart.xyz" displayed using Jmol.

Generate Positions and specify ADP_03_unit. cry as output.
(9) As number of positions we specify 2 , and the first operation as identity operation shall simply copy the first eight positions:

|  | factor | shift | source |
| :---: | :---: | :---: | :---: |
| $X:$ | 1.0 | 0.0 | $x$ |
| $Y:$ | 1.0 | 0.0 | $y$ |
| $Z:$ | 1.0 | 0.0 | $z$ |

(11) The last operation is the centering operation:

|  | factor | shift | source |
| :---: | :---: | :---: | :---: |
| $X:$ | 1.0 | 0.5 | $x$ |
| $Y:$ | 1.0 | 0.5 | $y$ |
| $Z:$ | 1.0 | 0.5 | $z$ |

The resulting file ADP_03_unit. cry should look like Fig. 2.3. If you are curious what kind of result we produced, you can load ADP_03_unit.cry into Trafo, select Action - Orthogonalize: Coordinates to convert fractional coordinates into Cartesian coordinates and save the result as ADP_03_unit_cart.cry. Read this file, delete the comment lines, replace the first line (the unit cell data) by the number of atoms in the file, 56 , insert a second line with a title, e.g., ADP, and save the result as "ADP_03_unit_cart.xyz" (note the double quotes; they prevent Trafo from adding the extension .CRY). You can read this XYZ file into $\mathrm{Jmol}(\mathrm{p} .2)$, and the result should look like Fig. 2.2.
In the next section, Generate: Block (p. 16), we shall see how to use this as base for generating packing diagrams.

## - References

(1) Tenzer, L.; Frazer, B.L.; Pepinsky, R. Acta Crystallogr. 1958, 11, 505, DOI: 10.1107/S0365110X58001389.
(2) Khan, A. A.; Baur, W. H. Acta Crystallogr., Sect. B 1973, 29, 2721, DOI: 10.1107/S0567740873007442.

| 7.502 | 7.5027 .546 | $90.00 \quad 90.00$ | 90.00 |
| :---: | :---: | :---: | :---: |
| P | 0.000000 | 0.000000 | 0.000000 |
| N | 0.000000 | 0.000000 | 0.500000 |
| 0 | 0.085000 | 0.146000 | 0.115000 |
| H | 0.014000 | 0.111000 | 0.573000 |
| H | 0.227000 | 0.146000 | 0.122000 |
| 0 | -0.085000 | -0.146000 | 0.115000 |
| H | -0.014000 | -0.111000 | 0.573000 |
| H | -0.227000 | -0.146000 | 0.122000 |
| 0 | -0.146000 | 0.085000 | -0.115000 |
| H | -0.111000 | 0.014000 | -0.573000 |
| H | -0.146000 | 0.227000 | -0.122000 |
| 0 | 0.146000 | -0.085000 | -0.115000 |
| H | 0.111000 | -0.014000 | -0.573000 |
| H | 0.146000 | -0.227000 | -0.122000 |
| P | 0.500000 | 0.000000 | 0.250000 |
| N | 0.500000 | 0.000000 | -0.250000 |
| 0 | 0.415000 | 0.146000 | 0.135000 |
| H | 0.486000 | 0.111000 | -0.323000 |
| H | 0.273000 | 0.146000 | 0.128000 |
| 0 | 0.585000 | -0.146000 | 0.135000 |
| H | 0.514000 | -0.111000 | -0.323000 |
| H | 0.727000 | -0.146000 | 0.128000 |
| 0 | 0.646000 | 0.085000 | 0.365000 |
| H | 0.611000 | 0.014000 | 0.823000 |
| H | 0.646000 | 0.227000 | 0.372000 |
| 0 | 0.354000 | -0.085000 | 0.365000 |
| H | 0.389000 | -0.014000 | 0.823000 |
| H | 0.354000 | -0.227000 | 0.372000 |
| P | 0.500000 | 0.500000 | 0.500000 |
| N | 0.500000 | 0.500000 | 1.000000 |
| 0 | 0.585000 | 0.646000 | 0.615000 |
| H | 0.514000 | 0.611000 | 1.073000 |
| H | 0.727000 | 0.646000 | 0.622000 |
| 0 | 0.415000 | 0.354000 | 0.615000 |
| H | 0.486000 | 0.389000 | 1.073000 |
| H | 0.273000 | 0.354000 | 0.622000 |
| 0 | 0.354000 | 0.585000 | 0.385000 |
| H | 0.389000 | 0.514000 | -0.073000 |
| H | 0.354000 | 0.727000 | 0.378000 |
| 0 | 0.646000 | 0.415000 | 0.385000 |
| H | 0.611000 | 0.486000 | -0.073000 |
| H | 0.646000 | 0.273000 | 0.378000 |
| P | 1.000000 | 0.500000 | 0.750000 |
| N | 1.000000 | 0.500000 | 0.250000 |
| 0 | 0.915000 | 0.646000 | 0.635000 |
| H | 0.986000 | 0.611000 | 0.177000 |
| H | 0.773000 | 0.646000 | 0.628000 |
| 0 | 1.085000 | 0.354000 | 0.635000 |
| H | 1.014000 | 0.389000 | 0.177000 |
| H | 1.227000 | 0.354000 | 0.628000 |
| 0 | 1.146000 | 0.585000 | 0.865000 |
| H | 1.111000 | 0.514000 | 1.323000 |
| H | 1.146000 | 0.727000 | 0.872000 |
| 0 | 0.854000 | 0.415000 | 0.865000 |
| H | 0.889000 | 0.486000 | 1.323000 |
| H | 0.854000 | 0.273000 | 0.872000 |

Figure 2.3: Alchemy crystal data file for ammonium dihydrogen phosphate (ADP), $\mathrm{NH}_{4} \mathrm{H}_{2} \mathrm{PO}_{4}$, generated from the data file shown in Fig. 2.1, corresponding to the unit cell content. Save the data as ADP_03_unit. cry if you want to repeat some of the example actions shown later.

### 2.4.3 Action - Generate Block

The Generate Block command from the Actions|Generate menu allows to create a block (cube) of new coordinates (positions) around the existing one. Specifically, this generates from a unit cell a cube consisting of several unit cells. Such operations might be required to construct packing diagrams or as a convenient tool to create the input coordinates for the calculation of dipolar second moments. The entire content of the unit cell itself should be created using Generate Positions (p. 10). In the dialog box Generate Block, one defines:



The parameter $\mathbf{n}$ describes how many unit cells will get created. For a given value of $n$, this command generates a cube of corner length $2 n+1$, or $(2 n+1)(2 n+1)(2 n+1)-1=(2 n+1)^{3}-1$ new positions around the input data. For a value of $n$ equal to 1 , a cube of 27 positions will get created, as indicated in the figure to the left. In the newly created output file, the original data set will be placed first, followed by the new data (this is important for later use of the results to calculate dipolar second moments).

## Example

In the section Symmetry Operations (p. 10) we demonstrated how to generate the content of the unit cell given the fractional coordinates of the asymmetric unit, using ammonium dihydrogen phosphate (ADP) as an example. If you look at Fig. 2.2, the result is not exactly what is expected naively. Rather than resulting in a sort of block, such generating symmetry operations often generate positions that lie in the neighboring unit cell (e.g., in Fig. 2.3 all atoms that have a negative coordinate or a coordinate greater than one). We would need to shift all these positions by integer multiples of the unit cell axes (i.e. add or subtract 1 ) to bring them into our expected block of $0 \leq x, y, z \leq 1$. Another solution is to simply add more unit cells around our central unit cell:

- Read file ADP_03_unit. cry (Fig. 2.3) into Trafo, remove the comment lines (if present) and save the file under the same name.
- Select Action - Generate Block, specify $n=1$ and save the result as ADP_04_unit_block.cry. Note that this creates 26 additional unit cells, 27 in total, corresponding to 1512 atomic positions.
- The file ADP_04_unit_block.cry is too big ( $>64 \mathrm{~KB}$ ) for Trafo's edit window, but you can still perform file related actions. E.g., to visualize this new result, clear the Trafo edit window (File - New) and select Action - Orthogonalize: Coordinates, leave the order of axes at their defaults, specify ADP_04_unit_block.cry as input file and ADP_04_unit_block_cart.cry as output file. Copy this file to ADP_04_unit_block_cart.xyz and use an editor that has a higher capacity as Trafo (the Windows editor Notepad will do) to replace the Cartesian crystal data line by the number of atoms, 1512, and insert a second line with a title (c.f. Fig. 2.4). Save the file and open it in Jmol (p. 2). Two different views of the result are shown in Fig. 2.5.

| 1512 |  |  |  |
| :--- | ---: | :--- | :--- |
| ADP | 27 | unit cells |  |
| P | 0.000000 | 0.000000 | 0.000000 |
| N | 0.000000 | 0.000000 | 3.773000 |
| O | 0.637670 | 1.095292 | 0.867790 |
| H | 0.105028 | 0.832722 | 4.323858 |
| $\cdots$ |  |  |  |

Figure 2.4: Beginning of file ADP_04_unit_block_cart. xyz, showing the structure of an XYZ file: number of atoms in the first line, a second line with a title, followed by the Cartesian coordinates of the atoms.


Figure 2.5: Two views of file ADP_04_unit_block_cart . xyz using Jmol.

### 2.4.4 Action - Orthogonalize Axes

The Orthogonalize Axes command from the Actions|Orthogonalize menu orthogonalizes the direction cosines of an oblique axis system in the order given by Rollett's procedure. This is an action required when dealing with single crystal NMR. Typically, in this experiment a crystal is mounted in some holder and X-ray diffraction is used to determine the orientation of the crystal axes with respect to the holder. If the crystal axes form an oblique system, i.e., at least one angle differs from $90^{\circ}$, it is useful to transform them into an orthogonal axes set, because one can then determine the Euler angles (p. 24) that relate the crystal axes to the crystal holder frame.
This action requires to assign the crystal axes $a, b, c$ to the coordinate axes $X, Y, Z$ to define the order of axes (p. 22).


Rollett's procedure [1]:

1. The axis selected as the $c$ axis is aligned with the coordinate system Z axis.
2. The axis selected as the $b$ axis is brought into the $Y Z$ plane. To orthogonalize this axis, the normal $n$ to the $b c$ plane is determined as the cross product ( p .23 ) $b \times c$, the new $b^{*}$ axis results from the cross product $c \times n$.
3. The axis selected as $a$ usually points somewhere along the $X$ direction. The orthogonalized $a^{*}$ is the cross product of $b^{*} x c$.

The input data need to contain three regular data lines corresponding to the direction cosines of the oblique axis system. All following data lines are ignored. Crystal axes data lines (p. 2) and origin data lines (p. 3) are neglected.

See also the topic Orthogonalize Coordinates (p. 19) for more information.

## - Example

The example results from a ${ }^{31} \mathrm{P}$ single-crystal NMR experiment on (5-methyldibenzophosphole)pentacarbonylmolybdenum(0) and has been published [2]. A large crystal of this compound has been grown and glued onto a hollow three-sided alumina cube measuring 4 mm on each side. Using X ray diffraction, the orientation of the monoclinic crystal axis system ( $a=31.113(2) \AA, b=7.7917$ (5) $\AA, c=17.9522(12), \beta=122.135(4)^{\circ}$, space group $C 2 / c$ ) with respect to this cube frame was determined (Fig. 2.6). This procedure also yields the direction cosines of the crystal axes with respect to the crystal holder (Fig. 2.7). In the data file shown in Fig. 2.7, I have included the cube frame $x, y, z$ axes (but $1.5 \AA$ away from the origin, OR). We can easily convert this to a XYZ file (compare Fig. 2.4), but Mercury (p. 2) or Jmol (p. 2) will not be able to deal with the result properly. The reason is that we are specifying "atoms" (OR, $\mathrm{a}, \mathrm{x}, \mathrm{z}$ ) that are unknown to those programs. One way around this is to replace them by sensible atoms, e.g., the XYZ file displayed in Fig. 2.8.


Figure 2.6: Orientation of the monoclinic crystal axes $a, b, c$ in the crystal holder frame $x, y, z$.

```
1.00}1.0001.00 90.00 90.00 90.00 
or 0.00000 0.00000 0.00000
a 0.91826 0.31965 -0.23380
b -0.18295 0.87884 0.43607
c -0.19058 -0.46992 0.86036
x 1.5 0.0}00.
y 0.0
z 0.0 0.0}1.
/* (OC)5Mo (Me-DBP) */
/* crystal axis system in */
/* cube axis system */
```

Figure 2.7: File MoMeDBP_01_crys_cube.cry, showing the orientation of the monoclinic crystal axes $a, b, c$ in the crystal holder frame $x, y, z$.

However, now there are many atoms close to each other, therefore the programs Mercury (p. 2) or Jmol (p. 2) will show "clusters" that are not very helpful, one example is shown in Fig. 2.9 that has been produced by reading file MoMeDBP_01_crys_cube.xyz into Mercury (p. 2). The reason is that the XYZ file format contains positions only and no information on bonds between the atoms. We can cure this if we export the XYZ file from Mercury (p. 2) into PDB format. This file format also contains information on connectivities, and we will simply delete the superfluous bonds in this file. The resulting file is shown in Fig. 2.10, where we have also numbered the H and Cl atoms so that it is easier to recognize the $a, b, c$ and $x, y, z$ axes in Fig. 2.11, a representation made using Mercury (p. 2) (Jmol (p. 2) will give the equivalent result). Note the increased complexity of file MoMeDBP_01_crys_cube.pdb (Fig. 2.10) compared to MoMeDBP_01_crys_cube. cry (Fig. 2.7). Now Mercury (p. 2) or Jmol (p. 2) will display the relative orientation of the two frames, crystal axes and crystal holder, in a much clearer way, Fig. 2.11.

```
7
(OC) 5Mo(Me-DBP) crystal axis system in cube axis system
C 0.00000 0.00000 0.00000
H
H -0.18295 0.87884 0.43607
H -0.19058 -0.46992 0.86036
Cl 1.5 0.0
Cl 0.0
Cl 0.0
```

Figure 2.8: File MoMeDBP_01_crys_cube.xyz, showing the orientation of the monoclinic crystal axes $a, b, c$ in the crystal holder frame $x, y, z$.


Figure 2.9: File MoMeDBP_01_crys_cube. xyz displayed by Mercury as cluster.

Using Jmol (p. 2) or Mercury (p. 2) with file MoMeDBP_01_crys_cube.pdb, you can verify that the oblique angle H1-C-H3 corresponds to $\beta=122.135(4)^{\circ}$, whereas the other two angles are $90^{\circ}$.
The purpose of Action - Orthogonalize Axes will be to turn $\beta$ also into a right angle by generating a new crystal axis $a^{*}$ that is orthogonal to $b$ and $c$. Read file MoMeDBP_01_crys_cube.cry into Trafo, select Action - Orthogonalize Axes, keep the Order of Axes settings at their default, specify MoMeDBP_02_crys_cube_orth. cry as output file, and agree to the normalization of the unit vectors. Convert MoMeDBP_02_crys_cube_orth. cry into XYZ file format as outlined above, use Mercury (p. 2) to convert to PDB format, clean up the superfluous bonds, and read the result back into Mercury (p. 2). Measure all "H-C-H" angles and verify that they are at $90^{\circ}$. Measure also the angle between the crystal holder $z$ axis and the crystal $c$ axis $\left(30.52^{\circ}\right)$.
You can now use file MoMeDBP_02_crys_cube_orth. cry in Trafo to calculate the Euler angles (p. 24) between both axes systems (compare the Euler angle $\beta$ with the angle between $z$ and $c$ axes just determined, $30.52^{\circ}$ ).

## - References

(1) Rollett, J. S. Computing Methods in Crystallography, Pergamon Press: Oxford 1975.
(2) Eichele, K.; Wasylishen, R. E.; Maitra, K.; Nelson, J. H.; Britten, J. F. Inorg. Chem. 1997, 36, 3539, DOI: 10.1021/ic970260q.

### 2.4.5 Direction Cosines

### 2.4.6 Action - Orthogonalize Coordinates

For many purposes it is useful to refer to the position of an atom in the unit cell of a crystal to the natural axes $a, b, c$ by means of the fractional coordinates (e.g., symmetry operations). In other cases, it is more convenient to use axes at right angles and measure distances in Angstrom units. The Orthogonalize Coordinates command from the Actions|Orthogonalize menu orthogonalizes


Figure 2.10: File MoMeDBP_01_crys_cube.pdb, showing the orientation of the monoclinic crystal axes $a, b, c$ in the crystal holder frame $x, y, z$.


Figure 2.11: File MoMeDBP_01_crys_cube.pdb, Fig. 2.10, displayed by Mercury and showing the orientation of the crystal axes in the crystal holder frame (cf. Fig. 2.6).
the fractional coordinates of an oblique axis system to Cartesian coordinates in the order given in Rollett's procedure. In fact, it will convert any fractional coordinates into Cartesian coordinates! It requires to assign the crystal axes $a, b, c$ to the coordinate axes $X, Y, Z$ to define the order of axes (p. 22).


Rollett's procedure:

1. The axis selected as the $c$ axis is aligned with the coordinate system Z axis.
2. The axis selected as the $b$ axis is brought into the $Y Z$ plane.
3. The axis selected as $a$ points somewhere along $X$, as required by its angles with $c$ and $b$.

This gives a non-trivial transformation matrix, L:

$$
\begin{aligned}
L & =\left[\begin{array}{ccc}
a \sin \beta \sin \gamma^{*} & 0 & 0 \\
-a \sin \beta \cos \gamma^{*} & b \sin \alpha & 0 \\
a \cos \beta & b \cos \alpha & c
\end{array}\right] \\
\cos \gamma^{*} & =(\cos \alpha \cos \beta-\cos \gamma) /(\sin \alpha \sin \beta)
\end{aligned}
$$

| 1.000 | 1.000 | 1.000 | 90.00090 .000 | 90.000 |
| :--- | ---: | ---: | ---: | ---: |
| or | 0.000000 | 0.000000 | 0.000000 |  |
| a | 0.964246 | 0.074545 | 0.254308 |  |
| b | -0.183882 | 0.879224 | 0.439491 |  |
| c | -0.190832 | -0.470540 | 0.861496 |  |

Figure 2.12: File MoMeDBP_02_crys_cube_orth.cry, showing the orientation of the orthogonalized crystal axes $a^{*}, b, c$ in the crystal holder frame $x, y, z$.

$$
\sin \gamma^{*}=\left(1-\cos ^{2} \gamma^{*}\right)^{1 / 2}
$$

The crystal axes data line (p. 2) is required! Origin data lines (p. 3) are converted as well. Example input: the fractional coordinates of TEPS in a triclinic system

| 8.98 | 8006. | 006 | 500 | 113.0 | 85.200 | 102.500 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| C3 | . 1193 | . 3168 | . 32 |  |  |  |
| C3 | . 2221 | . 1267 | -. 17 |  |  |  |
| C3 | . 2768 | . 4399 | . 41 |  |  |  |
| C3 | . 3915 | . 1312 | -. 17 |  |  |  |
| P3 | . 1197 | . 0484 | . 06 |  |  |  |
| S3 | . 1832 | -. 1916 | . 12 |  |  |  |
| C3 | -. 1193 | -. 3168 | -. 32 |  |  |  |
| C3 | -. 2221 | -. 1267 | . 17 |  |  |  |
| C3 | -. 2768 | -. 4399 | -. 41 |  |  |  |
| C3 | -. 3915 | -. 1312 | . 17 |  |  |  |
| P3 | -. 1197 | -. 0484 | -. 06 |  |  |  |
| S3 | -. 1832 | . 1916 | -. 12 |  |  |  |
| /* Tetraethyl diphosphine disulfide, TEPS */ |  |  |  |  |  |  |
| /* C8 H2O P2 S2 |  |  |  |  | */ |  |
| /* S.N.Dutta, M.M.Woolfson |  |  |  |  | */ |  |
| /* Acta Crystallogr., 14, 178,1961 |  |  |  |  | */ |  |

Output: fractional coordinates of TEPS converted to Cartesian coordinates

| 1.000 | 1.000 | 1.000 | 90.00090 .000 |
| :--- | :---: | :---: | ---: |
| C3 | 1.045919 | 1.667075 | 1.269696 |
| C3 | 1.947180 | 0.354132 | -1.213294 |
| C3 | 2.426743 | 2.115630 | 1.680507 |
| C3 | 3.432333 | 0.077197 | -1.102878 |
| P3 | 1.049426 | 0.072799 | 0.345577 |
| S3 | 1.606139 | -1.465968 | 1.393590 |
| C3 | -1.045919 | -1.667075 | -1.269696 |
| C3 | -1.947180 | -0.354132 | 1.213294 |
| C3 | -2.426743 | -2.115630 | -1.680507 |
| C3 | -3.432333 | -0.077197 | 1.102878 |
| P3 | -1.049426 | -0.072799 | -0.345577 |
| S3 | -1.606139 | 1.465968 | -1.393590 |

## - References

(1) Rollett, J. S. Computing Methods in Crystallography, Pergamon Press: Oxford 1975.

### 2.4.7 Order of Axes

In the dialog box Order of Axes, one defines:


With the dialog box Order of Axes, one assigns which of the crystal $a, b, c$ axes should be aligned to the Cartesian $X, Y, Z$ axes. The default setting assigns the old $a$ axis to the new $X$ axis, $b$ to $Y$, and $c$ to Z.

This information is required for:

- Orthogonalization of axes (p. 17)
- Orthogonalization of coordinates (p. 19)


### 2.4.8 Action - Orthogonalize: Calculate Normal

Given two vectors, this routine calculates the normal vector to the plane defined by the two vectors. The two vectors are defined by the Cartesian coordinates of their common starting point, followed by the Cartesian coordinates of the two end points. Note that the direction of this normal vector depends on the order in which the two vectors are defined, because the normal vector is calculated via the cross product ( $p .23$ ) of the two vectors.

## - Example input

```
1.00 1.00 1.00 90.0 90.0 90.0
OR 
P1 1.00
l1
```


## - Output

```
normal vector: 0.00000000 0.00000000 -4.00000000
normalized: 0.00000000 0.00000000 -1.00000000
```

This command considers only the first three coordinate triples. A Cartesian crystal axis line (p. 2) must precede the data.

### 2.4.9 Cross Product

The Vector or Cross Product of two vectors $A$ and $B$ :

$$
\mathbf{C}=\mathbf{A} \times \mathbf{B}
$$

generates a vector $C$ with direction perpendicular to the plane of $A$ and $B$ such that $A, B$ and $C$ form a right-handed system. The magnitude of $C$ is defined as $C=A B \sin \phi$, where $\phi$ is the angle between $A$ and $B$.
An alternate definition of the cross product consists in specifying the components of $C$ :

$$
\begin{aligned}
& C_{x}=A_{y} B_{z}-A_{z} B_{y} \\
& C_{y}=A_{z} B_{x}-A_{x} B_{z} \\
& C_{z}=A_{x} B_{y}-A_{y} B_{x}
\end{aligned}
$$

or:

$$
C_{i}=A_{j} B_{k}-A_{k} B_{j}
$$

where $i, j, k$ are all different and cyclic permutations of the indices $x, y, z$.

### 2.4.10 Action - Rotations: Calculate Euler Angles

The Calculate Euler Angles command from the Actions|Rotations menu allows to calculate Euler angles (p. 24) from the direction cosines of the new coordinate system with respect to the old coordinate system.
In order to be able to do this, the data are checked for validity with procedures pioneered by Michael D. Lumsden (Halifax, N.S.; Canada):

- check if the unit vectors are indeed unit vectors
- check if the unit vectors are orthogonal
- check if the unit vectors are right-handed; if left-handed, the user has the choice of inverting one axis (p. 24) selectively

If the data pass this test, the Euler angles are calculated using specific components of the unit vectors to get the angles. The result will be displayed by a dialog box, but also be copied to the clip board. From there, the routines to perform Euler rotations will retrieve them automatically and use them as defaults. Note, however, that any editing tasks (copy, cut etc.) may remove these data from the clipboard.
The input data may be preceded by a line defining the crystal axes (p. 2) and may have some comment lines (p. 3) or lines describing the origin (p. 3) interspersed (they are simply ignored). The routine takes the first three regular data lines (p. 3) as input.
An example is presented in Fig. 2.12, the direction cosines of an orthogonalized crystal axis system in a crystal holder frame. Using this file as input, this orientation can also be specified using the triple of Euler angles:


### 2.4.11 Invert Axis

In the dialog box Axis Inversion, one defines:


Inversion of a coordinate axis changes the handedness of the system. Select one axis to invert. Usually this dialog is called if the input data define a left-handed coordinate system. All calculations in Trafo are designed for right-handed coordinate systems!

### 2.4.12 Euler Angles

The triplet of Euler angles $(\alpha, \beta, \gamma)$ is useful to describe rotations or relative orientations of orthogonal coordinate systems. Unfortunately, their definition is not unique and in the literature there are as many different conventions as authors. The convention employed here is one of the more common ones. All rotations are in a counter-clockwise fashion (right-handed, mathematically positive sense).


The Euler angles $(\alpha, \beta, \gamma)$ relate two orthogonal coordinate systems having a common origin. The transition from one coordinate system to the other is achieved by a series of two-dimensional rotations. The rotations are performed about coordinate system axes generated by the previous rotation step (the step-by-step procedure is illustrated in the topic Rotation Matrices (p. 25). The convention used here is that $\alpha$ is a rotation about the Z axis of the initial coordinate system. About the $\mathrm{y}^{\prime}$ axis of this newly generated coordinate system a rotation by $\beta$ is performed, followed by a rotation by $\gamma$ about the new z axis.
Given the Euler angles, the step-by-step procedure illustrates how to move from one coordinate system to the other. However, given the two coordinate systems, how can one determine the Euler angles relating them? This is described in topic Determining Euler Angles (p. 27).

The usual ranges for $\alpha, \beta, \gamma$ are:

$$
\begin{aligned}
& 0 \leq \alpha \leq 360 \\
& 0 \leq \beta \leq 180 \\
& 0 \leq \gamma \leq 360
\end{aligned}
$$

## - References

(1) G. Arfken, Mathematical Methods for Physicists, 3rd. ed., Academic Press: New York 1985.
(2) M. E. Rose, Elementary Theory of Angular Momentum, Wiley: New York 1957.
(3) K. Schmidt-Rohr and H. W. Spiess, Multidimensional Solid-State NMR and Polymers, Academic Press: London 1994.

### 2.4.13 Rotation Matrices

Rotations or transformations from one coordinate system into another are conveniently described by the triplet of Euler angles (p. 24) ( $\alpha, \beta, \gamma$ ). Using the Euler angles, this three-dimensional problem can be dissected into a sequence of two-dimensional rotations, whereby in each rotation one axis remains invariant. Here, all rotations are counter clockwise (right-handed, mathematically positive sense).


## 2D Analogy:

In order to simplify the problem, let us start with a twodimensional rotation:
Suppose the coordinates, ( $x, y$ ), of a point in the twodimensional XY system are known, but we are actually interested in knowing the coordinates of this point in another coordinate system, $\mathrm{X}^{\prime} \mathrm{Y}^{\prime}$, which is related to the XY system by a counter-clockwise rotation by an angle $\varphi$.
As the figure indicates, the coordinates of the given point in the new coordinate system will be:

$$
\begin{aligned}
x^{\prime} & =x \cos \varphi+y \sin \varphi \\
y^{\prime} & =-x \sin \varphi+y \cos \varphi
\end{aligned}
$$

or, in matrix notation:

$$
R(\varphi)=\left(\begin{array}{cc}
\cos \varphi & \sin \varphi \\
-\sin \varphi & \cos \varphi
\end{array}\right)
$$

## Start: Coincidence

Now, transferred to a three-dimensional problem, the goal will be to describe the coordinates in a final rotated system ( $x, y, z$ ) which is related to some initial coordinate system $(X, Y, Z)$ by the Euler angles. The final system is developed in three steps, each step involving a rotation described by one Euler angle. At the start, both coordinate systems, $(X, Y, Z)$ and $(x(1), y(1), z(1))$, shall be coincident.



First Rotation:
The first rotation involves the Euler angle $\alpha$. The $x(1), y(1)$, $z(1)$ axis system is rotated about the $Z$ axis through an angle $\alpha$ counterclockwise relative to $X, Y, Z$ to give the new system $x(2)$, $y(2), z(2)$. It is clear from the figure that this rotation mixes the coordinates along X and Y , completely analogous to the two-dimensional rotation described above, while the coordinate along Z remains unaffected. The rotation matrix to describe this operation is given by:

$$
R_{\mathrm{z}}(\alpha)=\left(\begin{array}{ccc}
\cos \alpha & \sin \alpha & 0 \\
-\sin \alpha & \cos \alpha & 0 \\
0 & 0 & 1
\end{array}\right)
$$

## Second Rotation:

The second rotation involves the Euler angle $\beta$. The $\mathrm{x}(2), \mathrm{y}(2)$, $z(2)$ axis system is rotated about the $y(2)$ axis through an angle $\beta$ counterclockwise to generate the new coordinate system $x(3)$, $y(3), z(3)$. Analogously to the first Euler rotation, this mixes the coordinates along $x(2)$ and $z(2)$, while the coordinate along $y(2)$ remains unaffected. This operation also generates a line of nodes parallel to the direction of $y(2)$. The rotation matrix to describe this operation is given by:

$$
R_{\mathrm{y}}(\beta)=\left(\begin{array}{ccc}
\cos \beta & 0 & -\sin \beta \\
0 & 1 & 0 \\
\sin \beta & 0 & \cos \beta
\end{array}\right)
$$



## Third Rotation:

The last rotation involves the Euler angle $\gamma$. The $x(3), y(3), z(3)$ axis system is rotated about the $z(3)$ axis through an angle $\gamma$ counterclockwise to generate the final coordinate system $x, y, z$. Analogously to the first Euler rotation, this mixes the coordinates along $x(3)$ and $y(3)$, while the coordinate along $z(3)$ remains unaffected. The rotation matrix to describe this operation is given by:

$$
R_{z}(\gamma)=\left(\begin{array}{ccc}
\cos \gamma & \sin \gamma & 0 \\
-\sin \gamma & \cos \gamma & 0 \\
0 & 0 & 1
\end{array}\right)
$$

The combined effect of these three rotations is given by this transformation matrix:
$R_{\mathrm{z}}(\gamma) R_{\mathrm{y}}(\beta) R_{\mathrm{z}}(\alpha)=\left(\begin{array}{ccc}\cos \alpha \cos \beta \cos \gamma-\sin \alpha \sin \gamma & \sin \alpha \cos \beta \cos \gamma+\cos \alpha \sin \gamma & -\sin \beta \cos \gamma \\ -\cos \alpha \cos \beta \sin \gamma-\sin \alpha \cos \gamma & -\sin \alpha \cos \beta \sin \gamma+\cos \alpha \cos \gamma & \sin \beta \sin \gamma \\ \cos \alpha \sin \beta & \sin \alpha \sin \beta & \cos \beta\end{array}\right)$
Note: This type of rotation about sequentially newly generated axes produces the same result as rotations by the same angles about the fixed original axes, if the order of angles is reversed: $R_{\mathrm{Z}}(\alpha) R_{\mathrm{Y}}(\beta) R_{\mathrm{Z}}(\gamma)$ (cf. Mehring's book, appendix [1]).

$$
-26-
$$

## - References

(1) M. Mehring, Principles of High Resolution NMR in Solids; 2nd ed.; Springer-Verlag, Berlin, 1983.

### 2.4.14 Determining Euler Angles

Given the relative orientations of two coordinate systems, how do we go about determining the Euler angles relating them?
First, we need to decide on the coordinate system to take as the reference coordinate system, $\mathbf{X}, \mathbf{Y}, \mathbf{Z}$, and which one as derived coordinate system, $\mathbf{x}, \mathbf{y}, \mathbf{z}$. Because the Euler transformations allow to switch between coordinate systems easily, it does not really matter which one is selected.


The angle $\beta$ is simply the angle between the z axes of both coordinate systems, while $\alpha$ is the angle between the $X$ axis of the reference coordinate system and the projection of $z$ into the $X, Y$ plane. Finally, $\gamma$ is the angle between $y$ and the line of nodes. Computationally, we can work your way backwards from the components of the transformation matrix in Eq. 2.2:

- the zz component gives us $\cos (\beta)$
- use that $\beta$ to get $\alpha$ from components zx and zy
- use that $\alpha$ to get $\gamma$ from components xz and yz


### 2.4.15 Action - Rotations: Euler Rotations Forward

Rotates Cartesian coordinates from the old coordinate system (reference frame) into the new one (derived frame) using Euler rotations.
Example: Given a set of direction cosines specifying unit vectors that define the axes of the reference frame (Fig. 2.13), the rotation forward with Euler angles $\alpha=10, \beta=20, \gamma=30$ creates direction cosines (Fig. 2.14) which specify the coordinates of the reference frame as "seen" from the derived frame (Fig. 2.15). Using the option Calculate Euler Angles on this new set of data yields the Euler angles for going from the derived frame to the reference frame: $\alpha=150, \beta=20, \gamma=170$.

```
1.0 1.0 1.0 90.0 90.0 90.0
x 1.0 0.0 0.0
y 0.0 1.0 0.0
z 0.0 0.0 1.0
```

Figure 2.13: Orientation of the reference frame in its own axis system, 01_unit.cry.

| 1.0 | 1.0 | 1.0 | 90.0 | 90.0 |
| :--- | :--- | :--- | :--- | :--- |
| x |  | 0.0 |  |  |
| y |  | 0.714610 | -0.613092 | 0.336824 |
| z |  | 0.633718 | 0.771281 | 0.059391 |
|  | -0.296198 | 0.171010 | 0.939693 |  |

Figure 2.14: Orientation of the reference frame rotated forward by Euler angles $\alpha=10, \beta=20, \gamma=30$, 02_unit_forward.cry.


Figure 2.15: File 02_unit_forward.cry (Fig. 2.14) after conversion to XYZ and PDB format, as explained in Sect. 2.4.4, displayed using Mercury. It shows the orientation of the transformed reference frame $\mathrm{H} 1, \mathrm{H} 2, \mathrm{H} 3$ as seen from the derived frame $\mathrm{Cl} 1, \mathrm{Cl} 2, \mathrm{Cl} 3$

### 2.4.16 Action - Rotations: Euler Rotations Backwards

Rotates Cartesian coordinates from the new coordinate system (derived frame) into the old one (reference frame) using Euler rotations.
Example: Given a set of direction cosines specifying unit vectors, the rotation backward creates direction cosines which specify the coordinates of the derived frame in the reference frame. Using the option Calculate Euler Angles on this new set of data yields the Euler angles for going from the reference frame to the derived frame (i.e. it yields back the Euler angles used for the transformation, neglecting round off errors).

### 2.4.17 Action - Polar Angles: Calculate

This command transforms Cartesian coordinates into polar coordinates. A Cartesian crystal axis line (p. 2) must precede the data. The output will consist of lines giving the atom label, the azimuth angle $\alpha$, the polar angle $\beta$, and the distance from the origin, in Angstroems.

### 2.5 NMR Menu

### 2.5.1 Dipolar Second Moment



This program option calculates dipolar second moments for single crystal or powder samples, using coordinates supplied in a separate file or in the editor window of Trafo. The coordinates must be Cartesian coordinates (p. 19).
First, it is necessary to specify the spin system in question. For that purpose, you need to specify the isotope corresponding to the observed nucleus. It is assumed that the first atom specified in the input data by a coordinate triple corresponds to this observed nucleus.
Secondly, the isotope of the coupled nucleus is specified. The abundance of this isotope is then automatically set to the natural abundance of this isotope, but can be modified to reflect the actual level of enrichment, if any. It is assumed that all coordinates after the first atom, the observed nucleus, belong to this coupled nucleus. The dipolar second moment of a system where only a fraction $f$ of the crystal sites is occupied by nuclear spins is proportional to this fraction $f$ [6].
If desired, a cutoff distance can be specified. Any nucleus that is more remote than the cutoff distance is not included in the calculation. This is useful if the convergence of the calculation is to be tested but only one input file is used.
The type of moment calculation specifies whether the calculation should be performed for a powder or single crystal sample.

## - Single Crystals

In the case of a single crystal sample, the orientation of the magnetic field needs to be specified. This is done in a way suitable for single-crystal NMR rotation patterns.
The orientation of the rotation progress in the Cartesian frame of reference is specified by Euler angles ( p .24 ), where $\alpha$ and $\beta$ could also be considered as polar angles describing the orientation of the rotation axis, the z axis, while $\gamma$ marks the rotation progress, where the direction of the external magnetic field is parallel to the y axis. An additional constant phase angle $\phi$ for $\gamma$ can be specified; this phase angle is not considered in the output. The actual value of $\gamma$ is calculated according to:

$$
\gamma=\gamma(\text { start })+i * \gamma(\text { incr })+\phi
$$

The sense of rotation specifies whether the field moves as in a single-crystal NMR experiment, i.e. the physical, counter-clockwise rotation of the crystal actually corresponds to a clockwise rotation of the magnetic field $B_{0}$ in the interaction frame.

The dipolar second moment for a heteronuclear spin system is calculated according to:

$$
M_{2}(\text { hetero })=\frac{4}{3} S(S+1) \sum\left[\pi R\left(1-3 \cos ^{2} \theta\right)\right]^{2}
$$

and for a homonuclear spin system:

$$
M_{2}(\text { homo })=3 I(I+1) \sum\left[\pi R\left(1-3 \cos ^{2} \theta\right)\right]^{2}
$$

## - Powder Samples

In the case of a powder sample, no further data are required.
The dipolar second moment for a heteronuclear spin system is calculated according to:

$$
M_{2}(\text { hetero })=\frac{4}{15} S(S+1) \sum[2 \pi R]^{2}
$$

and for a homonuclear spin system:

$$
M_{2}(\text { homo })=\frac{3}{5} I(I+1) \sum[2 \pi R]^{2}
$$

## - Data Source

Most nuclear data have originated from Mason's extremely useful book on Multinuclear NMR [1]. The current version of R has been updated according to data from the IUPAC Recommendations 2001 [6] and IUPAC Recommendations 2008 [7]. The following data differ from those in reference [1], all magnetogyric ratios are according to [6]:

- the nuclear quadrupole moment values are from the "Year 2001 Q Values" collected by Pekka Pyykkö [3];
- Nd-145 apparently has a spin of $7 / 2$ instead of $5 / 2$ [4,5]. Similarly, U-235 has a spin of $7 / 2$ instead of 5/2 [4,5]. U-233 has been added to the tables [4,5].
- the magnetogyric ratio quoted in IUPAC Recommendations 2001 [6] for ${ }^{207} \mathrm{~Pb}$ was in error, in the 2008 recommendations [7] hidden in a footnote


## - References

(1) Joan Mason, Multinuclear NMR, Plenum Press, New York, 1987.
(2) A. Laaksonen and R. Wasylishen, J. Am. Chem. Soc. 1995, 117, 392-400, DOI: 10.1021/ja00106a044.
(3) P. Pyykkö, Mol. Phys. 2001, 99, 1617-1629, DOI: 10.1080/00268970110069010.
(4) Quantities, Units and Symbols in Physical Chemistry (IUPAC)
(5) CRC Handbook of Chemistry and Physics
(6) R. K. Harris, E. D. Becker, S. M. Cabral de Menezes, R. Goodfellow, P. Granger:

- Pure Appl. Chem. 2001, 73, 1795-1818, DOI: 10.1351/pac200173111795.
- Solid State Nucl. Magn. Reson. 2002, 22, 458-483, DOI: 10.1006/snmr.2002.0063.
- Angew. Chem. 2004, 116, 2070-2083, DOI: 10.1002/ange. 200380015.
(7) R. K. Harris, E. D. Becker, S. M. Cabral de Menezes, P. Granger, R. E. Hoffman, K. W. Zilm, Pure Appl. Chem. 2008, 80, 59-84, DOI: 10.1351/pac200880010059.


### 2.5.2 Diagonalise ( $\mathbf{d} 11>d 22>d 33$ )

Diagonalise a general chemical shift tensor. For an application, see section 3.1.2.

### 2.5.3 Diagonalise (s11<s22<s33)

Diagonalise a general magnetic shielding tensor. For an application, see section 3.1.1.

$$
-30-\overline{\text { July } 2,2021}
$$

## 3 Applications

### 3.1 Dealing with NMR Tensors

One of the more frequent questions that I receive with respect to Euler angles and NMR tensors is how to figure out the Euler angles relating two NMR tensors, most often an electric field gradient tensor and a magnetic shielding tensor obtained from ab initio calculations. Because I put some notes on Euler angles and tensor conventions on my home page, people seem to think that I am an expert in this matter and versed in math pretty well. Alas, the opposite is true, my math is pathetic, and the reason I wrote those notes is to remind myself of the essentials and to make it easier to communicate with others. This said, the following procedures will lack mathematical elegance, but they have worked for me, so far!
Before starting, I should advise you that you have to be very careful with such transformations. Frequently, the principal axes systems produced by Gaussian or other programs after diagonalization are either right- or left-handed systems. If you are using a left-handed PAS, Trafo will notify you when you calculate Euler angles. Usually, you can turn a left-handed system into a right-handed system by inverting the direction cosines of one axis. Also, eigenvectors are often reported in column format, but in a Cartesian system they are required row-wise. Thus, the output needs to be transposed.
The following example will assume that you have calculated an electric field gradient tensor and an absolute magnetic shielding tensor for one atom and that you want to determine the Euler angles orienting the shielding tensor in the EFG tensor.

### 3.1.1 The Absolute Magnetic Shielding Tensor

Unless specified explicitly, your ab initio software may not report the eigenvectors, for example the following excerpt from a Gaussian output may only show the shielding tensor elements with respect to the Gaussian Standard orientation (with Gaussian 03, you will get the eigenvectors only if you specify the keyword NMR=PrintEigenvectors):

```
2 Al Isotropic = 
    XX= 541.6395 YX= -1.2823 ZX= 0.5447
    XY= -2.7673 YY= 541.3822 ZY= 0.9831
    XZ= 1.7010 YZ= -1.2516 ZZ= 645.2708
    Eigenvalues: 539.4777 543.5316 645.2832
    Eigenvectors:
        (1) 0.685346 0.728190 -0.006350
        (2) 0.728136 -0.685374 -0.008940
        (3) 0.010862 -0.001504 0.999940
```

Lets pretend you don't have the eigenvectors. You could use Trafo to diagonalize the shielding tensor, but Trafo uses a transposed setup:

```
XX XY XZ
YX YY YZ
```

Therefore, for use with Trafo, your input should look like this (you can copy the three lines in the box into Trafo and save the file as 01_shielding.cry):

```
541.6395 -2.7673 1.7010
    rrrrr
    0.5447 0.9831 645.2708
```

Figure 3.1: Save file as 01_shielding.cry

Select from the Trafo menu: "NMR—Diagonalize (s11 < s22 < s33)" and specify as output file 02_csa_in_mf.cry. This file will now contain the directions of the shielding tensor principal axes in the molecular frame (mf). Open this file in Trafo:

```
1.0 1.0 1.0 90.0 90.0 90.0
OR 0.0 0.0 0.0
T11 
rrrr
```

Figure 3.2: Content of file 02_csa_in_mf.cry
Here, I have deleted the comment lines that provide further information. Compare the result with the eigenvectors reported by Gaussian above: the third vector is inverted because the Gaussian setup is a left-handed setup.
If you want, you can calculate the Euler angles for the orientation of the shielding tensor in the molecular frame; select "Action-Rotations-Calculate Euler angles":

```
alpha= 172.1167014131
beta= 179.3717030882
gamma= 305.3860273406
```

Figure 3.3: Euler angles calculated for the direction cosines in file 02_csa_in_mf.cry

### 3.1.2 The Electric Field Gradient Tensor

The EFG tensor may look like this in a Gaussian file:


Figure 3.4: EFG tensor from Gaussian output
Take this to construct the EFG tensor in the standard orientation (cf. Sect. 3.1.1):

```
-5494.450187 0.008592 -0.013931
    0.008592 -5494.472059 0.009529
    -0.013931 0.009529 -5494.687879
```

Figure 3.5: EFG tensor in standard frame, save as 03_efg.cry

Select "NMR—Diagonalize (s11<s22<s33)" and specify 04_efg_in_mf.cry as output, open the result:

```
1.0 1.0 1.0 90.0 90.0 90.0
OR 0.0 0.0 0.0
T11 0.059759 -0.046104 0.997148
T22 -0.303124 0.950924 0.062133
T33 -0.951076 
* --- Starting Matrix --- */
/* 
/* X: -5494.4502 0.008
/* Y: 
/* --- Symmetrized tensor in reference frame --- */
/* -5494.4502 0.0086 -0.0139 */
/* 0.0086 -5494.4722 
/* -0.0139 0.0095 -5494.6880 */
/* --- Tensor principal components and direction cosines --- */
/* (in reference frame:) */
/* T(ii) x y z */
/* -5494.689453 0.059759 -0.046104 0.997148 */
/* 
/* -5494.446777 -0.951076 -0.305972 0.042852 */
```

Figure 3.6: Content of file 04_efg_in_mf.cry

This tensor is not in the convention that I use in my simulation programs for EFG tensors: traceless and $\left|V_{\mathrm{xx}}\right| \leq\left|V_{\mathrm{yy}}\right| \leq\left|V_{\mathrm{zz}}\right|$ (work out the isotropic part and subtract it from the principal components; order the eigenvectors as specified above, and save the file as 05_efg_in_mf_ordered.cry):

| 1.0 1.0 | 1.0 | 90.090 .090 .0 |  |
| :--- | ---: | ---: | ---: |
| OR 0.0 | 0.0 | 0.0 |  |
| T11 | -0.303124 | 0.950924 | 0.062133 |
| T22 | -0.951076 | -0.305972 | 0.042852 |
| T33 | 0.059759 | -0.046104 | 0.997148 |

Figure 3.7: EFG in molecular frame in the proper convention, save as file 05_efg_in_mf_ordered.cry
Now you need to calculate the Euler angles for the orientation of the EFG tensor in the molecular frame; select "Action-Rotations-Calculate Euler angles":

```
alpha= 322.3498547908
beta= 4.3286051469
gamma= 145.4072983840
```

Figure 3.8: Euler angles calculated for the EFG tensor in file 05_efg_in_mf_ordered.cry

We will need these angles to transform the CSA into the EFG frame. Trafo copies those Euler angles to the clipboard:

```
322.3498547908 4.3286051469 145.4072983840
```

Figure 3.9: Euler angles
and they will stay there, unless you copy something else.

### 3.1.3 Transforming the CSA into the EFG frame

Read the file 02_csa_in_mf.cry, select "Action—Rotations—Euler rotations forwards", select a name for the output file: 06_csa_in_efg.cry. The EFG Euler angles should already be set properly: (unless

```
322.3498547908 4.3286051469 145.4072983840
```

Figure 3.10: Euler angles of the EFG tensor
you put something else into the clipboard)
Read the result file 06_csa_in_efg.cry, (delete the empty line), and select "Action—Rotations-Calculate Euler Angles". This gives the Euler angles for going from the EFG tensor to the shielding tensor:
$209.8756245150 \quad 176.2039520054 \quad 90.9096041873$

Figure 3.11: Euler angles that orient the shielding tensor in the EFG tensor
You are done!

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