### Manual

# singlepowder

# Integrating single-crystal area-detector data as powder diffractogram

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# Installation (Linux)

# Requirements

- C++11
- std library
- cmake
- Google-Test

Google-Test is installed as follows:

```
sudo aptitude install libgtest-dev
cd /usr/src/googletest/
sudo cmake . \\
sudo cmake --build . --target install
```

#### **Build**

In the directory singlepowder/build/:

```
cmake ..
```

#### **Testing**

In the directory singlepowder/build/test/:

./Test

#### Installation

In the directory singlepowder/build/src/:

sudo cp singlepowder /usr/bin/

# Running

singlepowder is run with one parameter, that is the name of the parameter file, i.e. parameters.txt:

singlepowder parameters.txt

## Input files

Figure 1 shows an example for input files. The directory

~/singlepowder\_test/TD015S001apex004/

contains the data files TD015S001apex004\_01\_0001.out, TD015S001apex004\_02\_0001.out, etc.

These files are listed in ~/singlepowder\_test/list.txt. This list file contains one line per image file. So far, only the columns filename, detectordistance and weight are used. The program only needs the name of the parameter file parameters.txt and gets all further information from this.

The output is written to the file given in the line output\_filename of the parameter file. In the example, the output file is named output.txt.

Everything behind the character # in the parameter file or the list file is a comment and ignored by the program. Empty lines or lines consisting of only a comment are allowed. The order of the parameters in the parameter file does not matter. Appart from comments, every line in the parameter file must consist of exactly two words. Thus, spaces in file names are not allowed.

The parameters in the parameter file are the following (all parameters must be given, there are no default values):

```
Width of one pixel in mm
          pixel_width
         pixel_height
                           Height of one pixel in mm
                           x \begin{cases} \text{index of the pixel hit by the direct} \\ y \end{cases}  beam at 2\theta = 0 (non-integer index possible)
      centre_pixel_x
      centre_pixel_y
             angle_min
                              The output powder diffractogram
             angle_max
                             covers 2\theta from angle_min to angle_max
                   step
                             with stepsize step.
                           path and name of the list file
image_list_filename
                           path for the data files
      data_directory
     output_filename
                           path and name of the output file
       output_format
                           format of the output file (standard or detailed)
```

# Geometry of the diffractometer

All lengths are in mm and all angles in deg.

Figure 2 shows a four circle diffractometer. The angles are shown with the directions used in the program (only  $2\theta$  is actually used so far). In order to avoid confusion,  $2\theta$  names the position of the detector while "powder angle"  $\varepsilon$  is used for the angle between the diffracted beam and the direct beam for a certain pixel. The pixel indices x and y and the pixel width w and height h are shown with the direction used by the classes Geometry and DetectorImage.

The variables in the figure and the following calculation refer to the following variables in the program code:

$d: {\it detector\_distance}$	(mm)
$2\theta$ : two theta	$(\deg)$
$\varepsilon$ : powder_angle	$(\deg)$
$x: pixel\_x$	(index)
$y: \operatorname{pixel\_y}$	(index)
$w: \mathbf{pixel\_width}$	(mm)
$h: \mathbf{pixel\_height}$	(mm)
$x_c: \text{centre\_pixel\_x}$	(index)
$y_c: \text{centre\_pixel\_y}$	(index)
$\Delta x$ : delta_x	(mm)
$\Delta y$ : delta_y	(mm)

# ~/singlepowder\_test/parameters.txt

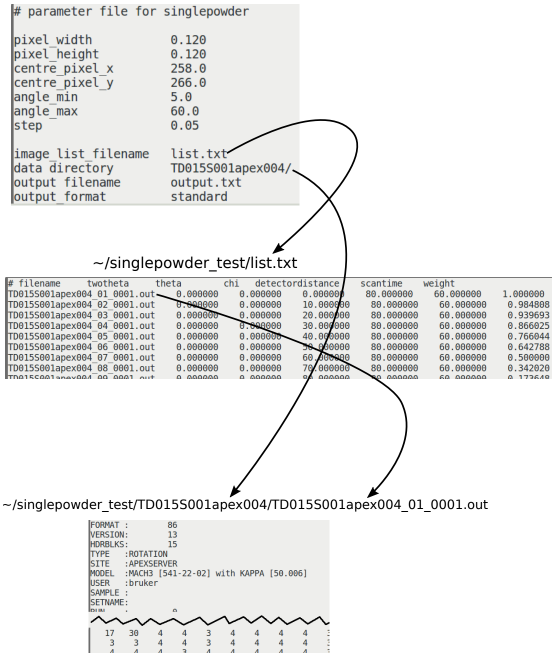


Figure 1: Example for input files. In the directory ~/singlepowder\_test/, the command singlepowder parameters.txt runs the program.

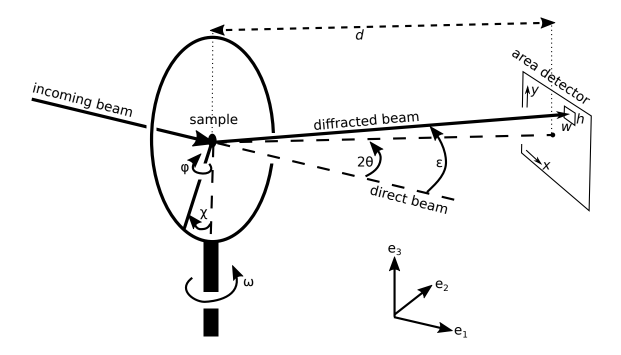


Figure 2: Direction of the angles  $2\theta$ ,  $\omega$ ,  $\chi$ ,  $\varphi$ , the pixel indices x, y and the Cartesian basis vectors  $e_1, e_2, e_3$ .

 $x_c$  and  $y_c$  are the indices of the pixel that is hit by the direct beam when all angles are set to zero. The deviation (in mm) from this pixel is for the pixel with indices (x, y):

$$\Delta x = w(x - x_c)$$
$$\Delta y = h(y - y_c)$$

Using the basis vectors<sup>1</sup> defined in Figure 2 (the origin is placed at the pivot point of the goniometer, i.e. the sample), we get for  $2\theta = 0$  the following coordinates of the pixel:

$$\boldsymbol{p} = \begin{pmatrix} d \\ -\Delta x \\ \Delta y \end{pmatrix}$$

The rotation matrix around  $e_3$  depends on  $2\theta$ :

$$R = \begin{pmatrix} \cos(2\theta) & -\sin(2\theta) & 0\\ \sin(2\theta) & \cos(2\theta) & 0\\ 0 & 0 & 1 \end{pmatrix}$$

<sup>&</sup>lt;sup>1</sup>The basis vectors are dimensionless and the coordinates have unit mm.

The coordinates of the pixel with indices (x, y) depend on d and  $2\theta$  and are thus:

$$\mathbf{p'} = R\mathbf{p}$$

$$= \begin{pmatrix} d\cos(2\theta) + \Delta x \sin(2\theta) \\ d\sin(2\theta) - \Delta x \cos(2\theta) \\ \Delta y \end{pmatrix}$$

The direct beam intersects the detector circle in the following point:

$$r = \begin{pmatrix} d \\ 0 \\ 0 \end{pmatrix}$$

For the angle  $\varepsilon$  between the direct and diffracted beam, the following condition holds:

$$|\boldsymbol{r}||\boldsymbol{p}|\cos(\varepsilon) = \boldsymbol{r}\cdot\boldsymbol{p'}$$
,

where  $\cdot$  denotes the scalar product.

From this, we can calculate the powder\_angle  $\varepsilon$ :

$$\varepsilon = \arccos \frac{\boldsymbol{r} \cdot \boldsymbol{p'}}{|\boldsymbol{r}||\boldsymbol{p'}|}$$

$$= \arccos \frac{d(d\cos(2\theta) + \Delta x\sin(2\theta))}{\sqrt{(d\cos(2\theta) + \Delta x\sin(2\theta))^2 + (d\sin(2\theta) - \Delta x\cos(2\theta))^2} d}$$

$$= \arccos \frac{d\cos(2\theta) + \Delta x\sin(2\theta)}{\sqrt{(d\cos(2\theta) + \Delta x\sin(2\theta))^2 + (d\sin(2\theta) - \Delta x\cos(2\theta))^2}}$$

This is the formula in Geometry::calculate\_powderangle().

## Integration and error propagation

The algorithm loops over all pixels of all detector images. For each detector image, the detector distance and  $2\theta$  are given in the list file. From the geometric parameters, the powder\_angle is calculated. The counts of this pixel are summed in the diffractogram at the according powder\_angle. The bin of the histogram is used, where the angle deviates maximally step/2. If the powder\_angle of the pixel lies outside the interval [angle\_min - step/2, angle\_max + step/2], the counts are discarded. The counts are weighted by the weight given in the list file.

Let i, j run over all pixels of all detector images for a fixed powder\_angle. The counts are  $c_i$  and the weights  $w_i$ . The intensity at this angle is:

$$I = \frac{\sum_{i=1}^{n} w_i c_i}{\sum_{j=1}^{n} w_j}$$

The error on the counts are  $\sigma_{c_i} = \sqrt{c_i}$ , so the error on the intensity can be computed in the following way:

$$\sigma_{I} = \sqrt{\sum_{i=1}^{n} \left(\frac{\partial I}{\partial c_{i}}\right)^{2} \sigma_{c_{i}}^{2}}$$

$$= \sqrt{\sum_{i=1}^{n} \left(w_{i} / \sum_{j=1}^{n} w_{j}\right)^{2} c_{i}}$$

$$= \frac{1}{\sum_{i=1}^{n} w_{j}} \sqrt{\sum_{i=1}^{n} w_{i}^{2} c_{i}}$$

During the integration, the following sums are collected for each powder\_angle:

$${\tt sum\_of\_weights} = \sum_{i=1}^n w_i$$
 
$${\tt sum\_of\_weighted\_counts} = \sum_{i=1}^n w_i c_i$$
 
$${\tt sum\_of\_squareweighted\_counts} = \sum_{i=1}^n w_i^2 c_i$$

These values are written to the output file when the parameter output\_format in the parameter file is set to detailed.

The sums are calculated in Diffractogram::add\_counts().

The intensity and its error is then calculated as follows:

intensity = 
$$I = \frac{\text{sum\_of\_weighted\_counts/sum\_of\_weights}}{\sigma_I = \sqrt{\text{sum\_of\_squareweighted\_counts/sum\_of\_weights}}}$$

This is the formula in Diffractogram::calculate\_intensities\_and\_errors().

# Structure of the program

The structure of the program is shown in Figure 3.

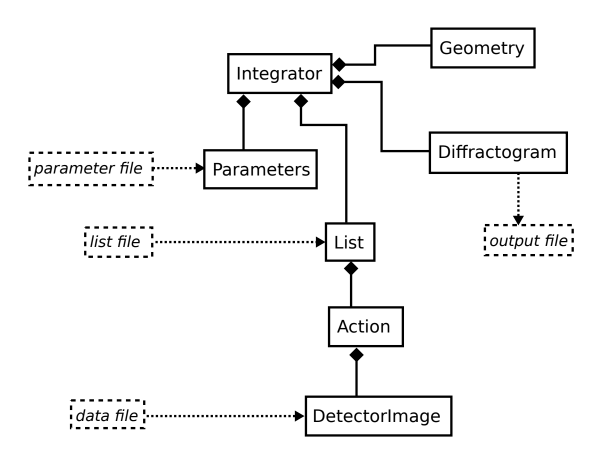


Figure 3: Diagram showing the hierarchy of the classes and the files they read and write.