TOPAS 64, Version 6

What’s New

by Alan A. Coelho

September 8, 2016

64 bit address space for GUI and command line executables

Works with 64 bit Windows 7, 8 and 10

Multithreaded

Stacking faults at speed

PDF refinement at speed

Charge-Flipping for neutron data

Surface Plots

Macro enhancements
Contents

1 New kernel functionality............................................................................................................. 4
  1.1 Introduction.......................................................................................................................... 4
  1.2 Miscellaneous ...................................................................................................................... 4
    1.2.1 TC-INPS.BAT and the aac$ macro ............................................................................. 4
    1.2.2 TOPAS is now 64 bit .................................................................................................. 4
    1.2.3 Indexing - Figure of merit ......................................................................................... 5
    1.2.4 INP file enhancements Miscellaneous ..................................................................... 5
    1.2.5 out_dependences and out_dependences_for .............................................................. 11
    1.2.6 The peaks buffer, speed and memory considerations ................................................ 12
    1.2.7 Threading ................................................................................................................... 12
    1.2.8 Using local to assist in using “for ... {}” loops ......................................................... 14
    1.2.9 Charge Flipping and neutron_data ........................................................................... 15
    1.2.10 Error determination using SVD ............................................................................... 16
    1.2.11 Error Propagation using prm_with_error ............................................................... 17
    1.2.12 New Keywords ......................................................................................................... 17
    1.2.13 New Functions .......................................................................................................... 19
  1.3 Stacking faults ..................................................................................................................... 19
    1.3.1 Generating the same stacking sequences each run .................................................... 20
    1.3.2 The SF_Smooth macro ............................................................................................... 20
    1.3.3 Fitting to DIFFaX test diamond data ....................................................................... 21
    1.3.4 Stacking faults from layers of different layer heights ............................................... 21
    1.3.5 Rietveld-Generated example ..................................................................................... 22
    1.3.6 Refining on layer heights ......................................................................................... 22
  1.4 PDF refinement .................................................................................................................... 23
    1.4.1 Inter and Intra molecule FWHMs .............................................................................. 26
    1.4.2 Instrument Sinc function Sinc-1.INP ....................................................................... 27
    1.4.3 Weighting of PDF and 2-Theta type data ..................................................................... 28
    1.4.4 Test_examples\pdf\BEQ-2.INP .................................................................................. 28
    1.4.5 Test_examples\pdf\BEQ-3.INP .................................................................................. 28
    1.4.6 Speeding up refinement with rebin_with_dx_of ....................................................... 28
    1.4.7 Refining on beq parameters ...................................................................................... 29
1.4.8  Structure Solution, Simulated Annealing.......................................................... 29
1.4.9  Rigid bodies with PDF data ........................................................................... 30
1.4.10 Occupancy merging with PDF data .............................................................. 30
1.4.11 Equivalence of pdf_gauss_fwhm and beq when there’s one atom type .......... 30

2  New GUI functionality.............................................................................................. 31
2.1  TOF x-axis can be displayed in d-spacing, Q and tof ...................................... 31
2.2  Displaying many files at once............................................................................. 31
2.2.1 Surface plots – 2D with offsets ........................................................................ 31
2.2.2 Inserting peaks and identifying scans .............................................................. 32
2.2.3 2D-offset Surface plots..................................................................................... 32
2.2.4 2D-offset Planview plots ................................................................................ 34
2.2.5 OpenGL Surface plots..................................................................................... 35
2.2.6 OpenGL – Weighted difference for colours ..................................................... 36
2.3  Normalizing scans to the maximum scan value within a Scan Window .......... 37
1 NEW KERNEL FUNCTIONALITY

1.1 Introduction

Directory test-mag have been removed to reduce file distribution size.

1.2 Miscellaneous

1.2.1 TC-INPS.BAT and the aac$ macro

The bath file TC-INPS.BAT runs through over 150 test examples in a time of a few minutes. These examples play an important role in program testing. Arguments passed via the command line to the test examples can contain the aac$ macro; if defined aac$ is expanded at the bottom of the INP file. For example, to terminate refinement after 100 iterations the following could be used:

```
tc test_examples\pdf\alvo4\rigid "macro aac$ { iters 100 verbose 0 }")
```

1.2.2 TOPAS is now 64 bit

Version 6 utilizes 64 bit addressing. The command line TC.EXE and the GUI TA.EXE both run on the Windows 64 bit operating system. It means that all available memory can be used. The 64 bit compile has resulted in a 10 to 20% increase in execution speed.

1.2.2.1 Limiting Memory Usage – MaxMem.TXT

Accidental INP file errors coupled with 64 bit address space can lead to excessive memory usage. A wrong decimal place in a lattice parameter for example could lead to the generation of billions of hkl's. In cases where all of RAM is used the Windows 7 and Windows 10 operating systems hang with the Task Manager being unresponsive. This reason for the ‘hang’ is due to the system swapping data/programs to and from virtual memory (typically a swap file on the hard disc). This ‘hang’ scenario is typically avoided using option (1) below which is the default. The file MaxMem.TXT, found in the main TOPAS directory, comprises two floating point numbers A and B and their use is as follows (all values in Gbytes):

1) If A=0 then the maximum allowed memory usage becomes:

\[ \text{Max\_Mem\_Allowed} = \text{Max\_Physical\_Memory} \times B \]

In other words 75% of the total physical memory

2) If the number in MaxMem.TXT is negative then the maximum allowed memory usage becomes:

\[ \text{Max\_Mem\_Allowed} = \text{Max\_Physical\_Memory} + A \]

3) If the number in MaxMem.TXT is positive then the maximum allowed memory usage becomes:

\[ \text{Max\_Mem\_Allowed} = A \]

The default value in MaxMem.TXT is zero which corresponds to case (1). For all cases, if memory usage exceeds Max\_Mem\_Allowed then TC.EXE aborts with the message “Memory allocation limit reached, increase limit in file MaxMem.TXT”. TA.EXE aborts without a message; instead it creates
the empty file called MaxMem-CHK.TXT. Checking the time/date stamp of MaxMem-CHK.TXT reveals whether TA.EXE aborted due to excessive memory usage.

1.2.3 Indexing - Figure of merit

The figure of merit $M$ used in indexing is as follows:

$$M = \frac{1}{\left( (1 + N_{\text{uni}}) \right) \sum \left( d_{o,\text{min}}^2 \right) \left( N_c / N_o \right)} \sum \left| d_{o,i}^2 - d_{c,i}^2 \right| Q_i$$

$$M = \frac{1}{\sum_j w_j}$$

where $Q_i = N_o \ w_i / \sum_j w_j$

Where $d_o$ and $d_c$ are the observed and calculated d-spacings, $N_o$ and $N_c$ the number of observed and calculated lines used, $N_{\text{uni}}$ the number of unindexed lines found and the summations are over the used observed indexing lines. $Q_i$ is a weighting that assists in the determination of extinction subgroups where $w_i$ could for example be the inverse of the error in the peak positions from a Pawley refinement (see indexing\MgIr\index.inp). The keyword index_I corresponds to $w_i$. The formulation of $Q_i$ is such that with or without $Q_i$ the figure of merit $M$ is of the same order of magnitude. The reciprocal-space lattice relationship solved during the indexing process (Coelho, 2000) includes $Q_i$ as follows:

$$2 \sin(2 \lambda / (4 \pi (X_{hk} h^2 + X_{hk} k^2 + X_{kl} l^2 + X_{hl} h k + X_{hl} l h + X_{lk} k l) (\pi / 360) (4 / \lambda^2) \sin(2 \theta)) \right) W_{hkl} = W_{hkl} / d_o^2$$

$$W_{hkl} = Q_{hkl} d_o^{\Delta \theta_{hkl}}$$

1.2.3.1 Extinction subgroup determination

At the end of an indexing further indexing runs are internally performed across extinction subgroups in an attempt to determine the most likely subgroup. These internal runs are seeded with already determined lattice parameters and in most cases the correct extinction subgroup is obtained without the need for $Q_i$.

1.2.4 INP file enhancements Miscellaneous

1.2.4.1 The num_runs keyword and Preprocessor improvements

[num_runs #]
[out_file = E;]
[system_before_save_OUT { $system_commands }]
[system_after_save_OUT { $system_commands }]

$num_runs$ defines the number of times the program executes (Runs) the INP file. Typically an INP file is run once; $num_runs$ changes this behaviour where the refinement is restarted and then performed again until it is performed $num_runs$ times. Information from one run to the next can be exchanged via the $out$ keyword and the include keyword. The INP file is read each Run but is not updated when $num_runs > 1$ and out_file is empty. Equations during a Run could well simplify into a constant, or indeed, the Constant keyword can be used such that during a Run a parameter is not
refined. From TB.EXE and Launch mode the Rwp graphical plot is appended such that it looks like `continue_after_convergence`. The following INP segment:

```plaintext
num_runs 10
yobs_eqn aac##Run_Number##.xy = Gauss(Run_Number, 1 + Run_Number);
min -2 max 20 del 0.01
```

produces on execution the following:

```
out_file determnes the name of the output file created when refinement terminates. The OUT file comprises the INP file but with parameter values updated. `out_file` defaults to the name of the INP file but with an OUT extension. If `num_runs` is greater than 1 and `out_file` is not defined then no OUT file is saved. This can speed up certain refinements where an OUT file is not needed. `out_file` is an equation that needs to evaluate to a string; here are some examples:

```plaintext
out_file aac.out  ' This will throw an exception
out_file = aac.out;  ' This will throw an exception
out_file = "aac.out";
out_file = String(aac.out);
out_file = If(Get(r_wp) < 10, "aac.out", "");
out_file = If(Get(r_wp) < 10, Concat(String(INP_File), ".OUT"), "");
```

The standard macro `Save_Best` uses `out_file` as follows:

```plaintext
macro Save_Best {
  #if (Run_Number == 0)
```
prm Best_Rwp_ = 9999;
#else
  prm Best_Rwp_ = #include Best_Rwp_.txt;
#endif
out Best_Rwp_.txt Out(If(Get(r_wp) < Best_Rwp_, Get(r_wp), Best_Rwp_))
out_file = If(Get(r_wp) < Best_Rwp_, Concat(String(INP_File), ".OUT"), "");

**system_before_save_OUT** executes the system commands defined in $system_commands string just before the *.OUT file is updated. The system commands are executed from the directory of the INP file. $system_commands can comprise any operating system commands. The macro Backup_INP uses **system_before_save_OUT**; it’s defined in TOPAS.INC as:

```plaintext
macro Backup_INP {
  system_before_save_OUT {
    copy INP_File##.inp INP_File##.backup
  }
}
```

**system_after_save_OUT** executes the system commands defined in $system_commands string just after the *.OUT file is updated.

### 1.2.4.2 Reserved macro Names

The following are internally generated macros that can be used in INP files.

**ROOT**: Returns the root directory of the program.

**INP_File**: Returns the name of the current INP file being without a file path or extension.

**Run_Number**: Returns the current run numer.

**File_Can_Open($file)**: Returns a 1 if $file can be opened or 0 of it can’t be opened.

Running an INP file called AAC.INP from TC.EXE where AAC.INP comprises:

```plaintext
ROOT
INP_File
Run_Number
File_Can_Open(aac.xy)
```

and AAC.XY exists will produce in TC.LOG the following:

```plaintext
c:\topas-6\aac
0
1
```
1.2.4.3 The #list directive – creating arrays of macros

#list creates arrays of macros with a single implied argument than can be expanded depending on the value of the single implied argument. For example, the following creates three arrays of macros called File_Name, Temperature and Time.

```
#list File_Name & Temperature(, & la) Time {
    File0001.xy 300 0.0
    { File0002 .xy } 320 10.2 ' Line with curly brackets
    File0003.xy 340 21.0
    File0017.xy { 360 + la } 28.9 ' Line with curly brackets
    File0107.xy 380 101.2
}
```

An element of the array is invoked using the first argument of the macro. In the case of File_Name and Time the first argument is implied. In the case of Temperature the first argument is empty as it needs to be. When the macro is invoked the first argument is a # type equation that must equate to an integer. Here’s an example use of the File_Name macro in the above list:

```
xdd File_Name(Run_Number)
```

Curly brackets (as seen in the above list) can be used as delimiters within #list. The following:

```
File_Name(1)
Temperature(1,)
Temperature(3, Get(la) + 0.01)
```

will produce on expansion:

```
File0002 .xy
(320)
(360 + (Get(la) + 0.01))
```

Using curly brackets as delimiters allows for curly brackets themselves to be part of the macro body.

1.2.4.4 The File_Variable and File_Variables macro

The File_Variable macro can be used to run a series of runs with parameters changing in a user defined manner; the macro is defined in TOPAS.INC as follows:

```
macro File_Variable(c, x_start, dx) {
    #if (Run_Number == 0)
        #prm c = x_start;
    #else
        #prm c = #include c##.txt;
    #endif
    #prm c##_next = c + dx;
    out c##.txt Out(#out c##_next)
}
```

Using File_Variable as follows:

```
File_Variable(occ, 0, 0.1)
```
Will generate a file called occ.txt for each Run with values ranging from 0.1 to 1 in steps of 0.1. A #prm is defined each run with the corresponding values. #out can be used to place the #prm into the INP file, for example, the following:

```plaintext
iters 0
num_runs 11
File_Variable(occ, 0, 0.1)
macro Out_File { Occ##Run_Number##.Out }
out_file Out_File
system_after_save_OUT {
  if (Run_Number)
    type Out_File >> aac.out
  else
    type Out_File > aac.out
  endif
}
yobs_eqn !aac.xy = 1;
min 10 max 50 del 0.01
CuKα1(0.0001)
Out_X_Ycalc( occ##Run_Number##.xy )
STR(F_M_3_M)
scale @ 0.0014503208
Cubic(5.41)
site Ce1 occ Ce+4 = #out occ; beq 0.2028
site O1 x 0.25 y 0.25 z 0.25 occ O-2 1 beq 0.5959
```

will generate 11 *.XY files each generated with a different occupancy for the Ce1 site as determined by the occ #prm. The names of the files would be occ0.xy to occ10.xy. Additionally, using the `system_after_save_OUT` keyword the file AAC.OUT will contain a concatenation of all the *.OUT files.

To iterate over two variable, pa and pb say, then the `File_Variables` macro, defined in TOPAS.INC as:

```plaintext
macro File_Variables(a, a1, a2, da, b, b1, b2, db) {
  if (Run_Number == 0)
    prm a = a1;
    prm b = b1;
  else
    prm a = #include a##.txt;
    prm b = #include b##.txt;
  endif
  prm a##_next = If(b >= b2, a + da, a);
  prm b##_next = If(b >= b2, b1, b + db);
  out a##.txt Out(#out a##_next)
  out b##.txt Out(#out b##_next)
}
```

can be used as follows:
iters 0
num_runs 36
File_Variables(pa, 0, 1, 0.2, pb, 0, 1, 0.2)
prm !pa = #out pa; prm !pb = #out pb;
out papb.txt append
   out_record out_eqn = pa; out_fmt "(%1.1f, "
   out_record out_eqn = pb; out_fmt "%.1f) "
   #if (pb == 1) Out_String("\n") #endif

On running the above the PAPB.TXT file contains:

(0.0, 0.0) (0.0, 0.2) (0.0, 0.4) (0.0, 0.6) (0.0, 0.8) (0.0, 1.0)
(0.2, 0.0) (0.2, 0.2) (0.2, 0.4) (0.2, 0.6) (0.2, 0.8) (0.2, 1.0)
(0.4, 0.0) (0.4, 0.2) (0.4, 0.4) (0.4, 0.6) (0.4, 0.8) (0.4, 1.0)
(0.6, 0.0) (0.6, 0.2) (0.6, 0.4) (0.6, 0.6) (0.6, 0.8) (0.6, 1.0)
(0.8, 0.0) (0.8, 0.2) (0.8, 0.4) (0.8, 0.6) (0.8, 0.8) (0.8, 1.0)
(1.0, 0.0) (1.0, 0.2) (1.0, 0.4) (1.0, 0.6) (1.0, 0.8) (1.0, 1.0)

1.2.4.5 Equation String, Concat and Variable_Name_From_String functions

A distinction is made between parameter names and strings. Text occurring between double quotation marks or between brackets in the String function are deemed as String types. Note, the output from the following:

   prm aabb = 1.234;
   prm sasb = 4.321;
   prm sa = "aa";
   prm sb = "bb";
   prm sc = sb;
   prm = Variable_Name_From_String(Concat(sa, sc));  : 1.23400`
   prm = Variable_Name_From_String(Concat(String(sa), String(sb)));  : 4.32100`

The arguments of Concat are by default variable names and not strings. Here are some examples:

   prm = Concat("a", "b", "c"); ' result = "abc"
   prm = Concat(String(a), "b", "c"); ' result = "abc"
   prm p = "a"; prm = Concat(p, "b", "c"); ' result = "abc"

The Save_Best macro uses strings as follows:

macro Save_Best
{
   #if (Run_Number == 0)
      prm Best_Rwp_ = 9999;
   #else
      prm Best_Rwp_ = #include Best_Rwp_.txt;
   #endif
   out Best_Rwp_.txt Out(If(Get(r_wp) < Best_Rwp_, Get(r_wp), Best_Rwp_))
   out_file = If(Get(r_wp) < Best_Rwp_,Concat(String(INP_File),".OUT"), "");
}
1.2.4.6  dummy and dummy_prm keywords

The dummy keyword reads a word from the input stream. dummy_prm is similar except it reads parameter dependent text. For example, in the following the text in Red is loaded by dummy_prm and ignored by the Kernel.

```
load xo dummy_prm I
{
10 = 1/Max(0.00023, 0.0001); min 10 max = Val 2; @ 100
...
```

1.2.5  out_dependences and out_dependences_for

```
out_dependences $user_string
out_dependences_for $user_string $object_name
```

*out_dependences* outputs dependences for the most previously defined *prm* or *local* parameter. For example, the following:

```
iters 1
prm d 1
prm e 1
prm f 1
prm c = e + f;
prm b = d + e;
prm a = b + c;
out_dependences a_tag
penalty = a^2;
```

produces on refinement termination the following in standard output (fit window or DOS command line):

```
out_dependences a_tag prm_10
Object name followed by prm name
    prm_10   e
    prm_10   f
    prm_10   d
```

*out_dependents_for* is similar except that it names an object that is not a parameter, for example, the following lists independent refined parameters associated with the most recently defined rigid body.

```
rigid
...
out_dependents_for tag_1 rigid
```

There are many *$object_name’s* that are valid. Basically all parameters can be tagged, e.g.

```
x, y, z, occ, beq, u11, u22, u33, u12, u13, u23, a, b, c, al, be, ga, etc...
```

In addition the following non-parameters can be tagged:
1.2.6 The peaks buffer, speed and memory considerations

Anisotropic peak shapes result in the peaks buffer holding as many peaks as there are hkls. For problems with 100,000s of peaks the calculation time and storage of the peaks buffer can be prohibitive. This situation can be mitigated using the phase dependent keyword `peak_buffer_based_on`.

```
[str...] [hkl İs...] [xo_Is...] [d_Is...]
peak_buffer_based_on !E...
peak_buffer_based_on_tol !E
```

The normal means of determining the size of the peak buffer is over ruled when `peak_buffer_based_on` is defined. With `peak_buffer_based_on` peaks are grouped according to the `peak_buffer_based_on` criterion. For example, to insert a peak into the peak buffer at x-axis intervals of 1 then the following can be used:

```
peak_buffer_based_on = Xo;
peak_buffer_based_on_tol 1
```

Thus peaks with similar Xo’s, as defined by `peak_buffer_based_on_tol`, are grouped. Occasionally peaks that are a function of hkls have groups of hkls that are of the same peak shape and at a similar x-axis position. The following demonstrates how to group these peaks such that a single peak shape is calculated.

```
peak_buffer_based_on = Xo;
peak_buffer_based_on_tol .01
peak_buffer_based_on = sh;
peak_buffer_based_on_tol 1e-7
```

Where `sh` can be a spherical harmonics parameter or an equation describing hkI dependence or a `march_dollase` parameter. When more that one `peak_buffer_based_on` is defined then peak groups formed obey all of the `peak_buffer_based_on`'s.

`peak_buffer_based_on` disables the peak stretching procedures and any defined `aberration_range_change_allowed`. `peak_buffer_based_on` can be a function of the reserved parameters H, K, L, M, D_spacing, X, Xo, Th.

Depending on the problem at hand smaller values such as 1e-7 can significantly reduce the number of peaks stored in the peaks buffer (a factor of 15 at times) without significantly affecting Rwp. A negative value for `peak_buffer_based_on_tol` will force a calculation for each peak resulting in independent hkl peak shapes, for example:

```
peak_buffer_based_on 1
peak_buffer_based_on_tol -1
```

1.2.7 Threading

TOPAS is threaded to a large extent; this allows the utilization of multiple processors which results in faster program execution. The degree of speedup is computer and problem dependent. For non-
trivial problems the gain is 2 to 4 for a 4 core laptop PC with four i7 processors as seen in the following table.

<table>
<thead>
<tr>
<th>INP File</th>
<th>Comment</th>
<th>Time (secs)</th>
<th>Gain</th>
</tr>
</thead>
<tbody>
<tr>
<td>\t\alvo4a &quot;#define TEST_ #define USE_FT_ #define USE_SH_ &quot;</td>
<td>ft, spherical harmonics</td>
<td>1.70</td>
<td>3.97</td>
</tr>
<tr>
<td>\absorption-edge\A2O3-pam</td>
<td>Absorption edge, modify peak</td>
<td>1.46</td>
<td>3.87</td>
</tr>
<tr>
<td>\cime-z-auto &quot;#define TEST_2 _&quot;</td>
<td>no decompose</td>
<td>7.08</td>
<td>3.54</td>
</tr>
<tr>
<td>\tof\tof_bank2_2</td>
<td>user_defined_convolution</td>
<td>0.18</td>
<td>3.49</td>
</tr>
<tr>
<td>\wppm\cube-in-normal-1 &quot;#define TEST_ &quot;</td>
<td></td>
<td>2.34</td>
<td>3.44</td>
</tr>
<tr>
<td>\alvo4a &quot;#define TEST_ &quot;</td>
<td></td>
<td>1.22</td>
<td>3.24</td>
</tr>
<tr>
<td>\single\crystal\ae14-approx-a &quot;#define TEST_ &quot;</td>
<td></td>
<td>1.03</td>
<td>3.21</td>
</tr>
<tr>
<td>\mag\mag &quot;#define TEST_ &quot;</td>
<td></td>
<td>1.09</td>
<td>2.94</td>
</tr>
<tr>
<td>\single\crystal\pnn_0_2_adps</td>
<td>3970 parameters</td>
<td>6.32</td>
<td>2.75</td>
</tr>
<tr>
<td>\single\crystal\ae14-adsps</td>
<td></td>
<td>0.29</td>
<td>2.50</td>
</tr>
<tr>
<td>\l250</td>
<td></td>
<td>0.39</td>
<td>2.72</td>
</tr>
<tr>
<td>\tube-tails &quot;#define TEST_ &quot;</td>
<td></td>
<td>1.48</td>
<td>2.56</td>
</tr>
<tr>
<td>\bso4a &quot;#define TEST_ &quot;</td>
<td></td>
<td>2.29</td>
<td>2.50</td>
</tr>
<tr>
<td>\single\crystal\pnn_0_2_ &quot;#define TEST_1_ &quot;</td>
<td></td>
<td>3.23</td>
<td>2.48</td>
</tr>
<tr>
<td>\sp\serine__evans_N_ta_bang_rot_z</td>
<td>z-matrix</td>
<td>0.67</td>
<td>2.41</td>
</tr>
<tr>
<td>\ft\alvo4a &quot;#define TEST_ #define USE_FT_ &quot;</td>
<td>ft, no spherical harmonics</td>
<td>0.38</td>
<td>2.38</td>
</tr>
<tr>
<td>\ft\alvo4a &quot;#define TEST_ &quot;</td>
<td>no ft, no spherical harmonics</td>
<td>0.21</td>
<td>2.29</td>
</tr>
<tr>
<td>\single\crystal\ae14-adsps &quot;#define TEST_ &quot;</td>
<td>no approximate_A</td>
<td>0.57</td>
<td>2.29</td>
</tr>
<tr>
<td>\capillary-lpsd\capillary-simulated</td>
<td></td>
<td>0.10</td>
<td>2.28</td>
</tr>
<tr>
<td>\v2o3a</td>
<td></td>
<td>0.24</td>
<td>2.25</td>
</tr>
<tr>
<td>\peak-intensity-extraction\zhuleibai</td>
<td></td>
<td>0.16</td>
<td>2.20</td>
</tr>
<tr>
<td>\sp\serine__evans_N_ta_bang_rot_z</td>
<td></td>
<td>0.82</td>
<td>2.18</td>
</tr>
<tr>
<td>\voigt-approx\fit-obj.inp &quot;#define TEST_ &quot;</td>
<td></td>
<td>4.39</td>
<td>2.18</td>
</tr>
<tr>
<td>\zo2 &quot;#define TEST_ &quot;</td>
<td></td>
<td>1.87</td>
<td>2.05</td>
</tr>
<tr>
<td>\cime-z-auto &quot;#define TEST_1_ &quot;</td>
<td>decompose</td>
<td>6.95</td>
<td>2.00</td>
</tr>
<tr>
<td>\k-factor\k-factor</td>
<td></td>
<td>0.33</td>
<td>1.95</td>
</tr>
<tr>
<td>\single\crystal\pnn_0_2_ &quot;#define TEST_2_ &quot;</td>
<td>no approximate_A</td>
<td>25.68</td>
<td>1.92</td>
</tr>
<tr>
<td>\mag\\ohn\evans\\soe_fi_p1_to\fullpro\tric_0</td>
<td></td>
<td>4.00</td>
<td>1.90</td>
</tr>
<tr>
<td>\peak-intensity-extraction\pawley1 &quot;#define TEST_ &quot;</td>
<td></td>
<td>1.73</td>
<td>1.83</td>
</tr>
<tr>
<td>\cf\cf-1-a_y &quot;#define TEST_ &quot;</td>
<td>Charge Flipping</td>
<td>12.87</td>
<td>1.81</td>
</tr>
<tr>
<td>\ft\alvo4a &quot;#define TEST_ #define USE_FT_ &quot;</td>
<td>ft, no spherical harmonics</td>
<td>0.53</td>
<td>1.79</td>
</tr>
<tr>
<td>\sp\serine__evans_N_ta_bang_rot</td>
<td></td>
<td>0.74</td>
<td>1.71</td>
</tr>
<tr>
<td>\pvs &quot;#define TEST_ &quot;</td>
<td></td>
<td>1.22</td>
<td>1.70</td>
</tr>
<tr>
<td>\stacking-faults\kaolinite &quot;#define TEST_3_ #define Speed&quot;</td>
<td>with Speed</td>
<td>0.58</td>
<td>1.66</td>
</tr>
<tr>
<td>\stacking-faults\kaolinite &quot;#define TEST_1_ &quot;</td>
<td></td>
<td>3.10</td>
<td>1.60</td>
</tr>
<tr>
<td>\clay</td>
<td></td>
<td>0.75</td>
<td>1.53</td>
</tr>
<tr>
<td>\tof\tof_bank2_1 &quot;#define TEST_ &quot;</td>
<td></td>
<td>0.60</td>
<td>1.53</td>
</tr>
<tr>
<td>occ-merge &quot;#define TEST_&quot;</td>
<td>3.37</td>
<td>4.90</td>
<td>6.20</td>
</tr>
<tr>
<td>---------------------------</td>
<td>------</td>
<td>------</td>
<td>------</td>
</tr>
<tr>
<td>\simon\alan</td>
<td>Many small strs</td>
<td>0.73</td>
<td>0.97</td>
</tr>
</tbody>
</table>

Attention has been paid to reducing memory usage at the thread level. This is particularly apparent when using rigid bodies or occupancy merge where Version 6 uses far less memory than Version 5. Except for penalties all items are threaded and they include:

- Peak generation
- All convolutions
- All derivatives that are a function of Ycalc
- Equations that are a function of changing variables such as X, Th, D_spacing etc...
- Pawley refinement
- Structure refinement
- Charge Flipping
- Magnetic refinement
- Stacking faults
- PDF refinement
- Conjugate gradient solution method
- Indexing

1.2.7.1 Setting the maximum number of threads - MaxNumThreads.TXT

The maximum number threads available is used by default. The user can limit the maximum number of threads by editing the file MaxNumThreads.TXT. This file is read on program startup; it contains a single number, lets call it Max_Threads_File, which defines the maximum number of threads. Nonexistence of the file or a Max_Threads_File of zero results in the program using the maximum number of threads available. If Max_Threads_File is negative then the maximum number of threads is set to the following:

\[
\text{Max\_Number\_Threads} = \text{Max}(1, \text{Max\_Threads\_File} + \text{Max\_Threads\_Available});
\]

1.2.8 Using local to assist in using “for ... {}” loops

The following parameters have had their status changed from global to ‘local’ allowing their use in ‘for’ loops.

```plaintext
march_dollase $Name
spherical_harmonics_hkl $Name
sites_geometry $Name
sites_distance $Name
sites_angle $Name
sites_flatten $Name
```

To constrain the march_dollase parameter, as used in the PO macro, to the same value within a “for xdds { for strs { ... }}” construct across two or more structures then simply give them the same name, for example:

```plaintext
PO(pol, 0.8, , 1 0 4)
```
See examples po-constrained-create.inp and po-for.inp in the test_examples\po-constrained directory. Note also the use of “if Prm_Then(...) {...}” rather than “for strs 1 to 1 {...}” to facilitate the writing of the INP file.

The $Name in spherical_harmonics_hkl is ‘local’ but the spherical harmonics coefficients are global. In the following:

```
PO_Spherical_Harmonics(sh2, 8 load sh_Cij_prm {
  k00 sh2_c00 1.0000
  k41 sh2_c41 0.1000
  k61 sh2_c61 -0.2000
  k62 sh2_c62 0.3000
  k81 sh2_c81 -0.4000
}
```

the sh2 parameter is local to the str and the coefficients k00, k41 etc... are global. This allows the constraining of coefficients across different structures within “for strs”; see examples posh-constrained-create.inp and posh-for.inp in the test_examples\po-constrained directory:

1.2.9 Charge Flipping and neutron_data

The neutron_data keyword informs the charge flipping routine that neutron scattering lengths are to be used. It also results in the following default neutron flipping routine being used:

```
flip_equation =
  If(And(Get(density) < Get(threshold), Get(density) > 0.4
    Get(min_density)),
    -Get(density),
    Get(density)
 );
```

The flip_neutron equation defines the 0.4 in the above equation. For example, the following can be used to change the default to 0.5:

```
flip_neutron = 0.5;
```

The tangent formula is made less accurate due to the negative scattering of H atoms. However, if positive scattering lengths are dominant then the tangent formula can stabilize refinement. For example, try:

```
Tangent(.3, 30)
  tangent_scale_difference_by = Ramp(1, 0, Nc);
```

See test_examples\cf\neutron-cime\cf-neutron.inp

1.2.9.1 Powder data, the A matrix and the Tangent formula

In the case of charge-flipping from powder data TOPAS uses the diagonally normalized A-matrix cf_in_A_matrix (see example cf\cf-cime.inp), which we will call D, from a Pawley refinement (see example cf\cf-cime-pawley.inp) to modify normalized structure factors Eh calculated during the charge-flipping process; this produces better results than using reflections output in a SHELX format. Equation (1-3) shows how the structure factors are modified.
\[ E_{\text{h,calc,modified}} = E_{\text{h,calc}} \sqrt{\frac{I_{\text{obs,h,k}}}{I_{\text{calc,h,k}}}} \]  

where \( I_{\text{calc,h,k}} = \sum_k D_{h,k}^2 I_{\text{calc,k}} \) and \( I_{\text{obs,h,k}} = \sum_k D_{h,k}^2 I_{\text{obs,k}} \)

The subscripts \( h \) and \( k \) correspond to reflections \( h \) and \( k \) respectively and the summation in \( k \) is over all reflections. \( I_{\text{calc,k}} \) and \( I_{\text{obs,k}} \) corresponds to observed and calculated intensities. Equation (1) modifies the calculated intensities to include intensities from overlapping peaks. When there’s no overlap \( D_{h,k} = 1 \) and \( D_{h,k} = 0 \) and the calculated intensities as well as \( E_h \) are not modified. When using the direct-methods tangent formula within the charge-flipping process as described by Coelho (2007), the \( D \) matrix is also used to modify \( E_h \) values used in triple phase relationships as shown in equation (2).

\[ E_{\text{obs,h,modified}} = E_{\text{obs,h}} q_h + E_{\text{calc,h}} (1-q_h) \]  

where \( q_h = \sum_k D_{h,k}^2 \)

\( E_{\text{obs,h}} \) and \( E_{\text{calc,h}} \) corresponds to tangent formula \( E_h \) values calculated from the observed and calculated intensities respectively. \( E_{\text{calc,h}} \) is typically not used in the tangent formula, however, intensities used for determining \( E_{\text{obs,h}} \) can be grossly in error due to peak overlap. Equation (2) therefore influences triple phase relationships by weighing \( E_{\text{obs,h}} \) by \( q_h \); when there’s no overlap \( q_h = 1 \) resulting in no modification. When there’s significant overlap then \( q_h \) is small and the influence of triple phase relationships using the \( h \) reflection is reduced. Equation (2) also includes a \( (1-q_h) \) portion of \( E_{\text{calc,h}} \) thus stating that when there’s significant overlap the calculated \( E_h \) is to be more trustworthy than the observed \( E_h \). Equation (1-4) corrects for errors in \( E_h \) when \( I_{\text{obs}} \) is similar to \( I_{\text{calc}} \); this assists in reducing the goodness of fit value thus enhancing the chances of solving the structure.

1.2.10 Error determination using SVD

Errors have previously been determined from a covariance matrix obtained by LU decomposition. Version 6 uses Singular Value Decomposition (SVD) with resulting errors typically smaller for strongly correlated parameters. Additionally SVD errors more closely resemble those obtained by the bootstrap method. bootstrap_errors are potentially more accurate as it considers parameter limits; for example the fact that intensities are positive is not considered by matrix inversion.

The keyword use_LU_for_errors can be used to force the use of LU decomposition. The three means of determining errors are demonstrated in a Pawley refinement of Y2O23 in the following INP files:

Test_examples/svd-errors/y2o3a-lu.inp  
Test_examples/svd-errors/y2o3a-svd.inp  
Test_examples/svd-errors/y2o3a-boot.inp

Comparisons showing the similarity between the bootstrap and SVD errors are seen in the following INP snippets:

<table>
<thead>
<tr>
<th>Method</th>
<th>Model Parameters</th>
<th>Coordinates</th>
</tr>
</thead>
<tbody>
<tr>
<td>LU</td>
<td>Full Axial Model(12, 15, 12, 5.1, @ 8.56191^-0.06668)</td>
<td>8.56191^-0.06668</td>
</tr>
<tr>
<td>SVD</td>
<td>Full Axial Model(12, 15, 12, 5.1, @ 8.56190^-0.05045)</td>
<td>8.56190^-0.05045</td>
</tr>
<tr>
<td>Boot</td>
<td>Full Axial Model(12, 15, 12, 5.1, @ 8.56191^-0.04988)</td>
<td>8.56191^-0.04988</td>
</tr>
</tbody>
</table>
Note the very large errors obtained by LU-decomposition for intensities that are 100% correlated.

1.2.11 Error Propagation using `prm_with_error`

Fixed parameter errors determined outside of refinement can be included and propagated to dependent parameters using the keyword `prm_with_error`. For example, consider the INP snippet (see test_example\prm-with-error.inp):

```plaintext
xo_1s
  xo 0 I = 10 t i;
  prm i  9.99999_0.00065 min 1e-6
  prm_with_error !t  1_0.33
  prm t_squared = t^2; : 1.00000_0.66000
```

Here the parameter is defined using `prm_with_error` and with an error of 0.33; this error is then used to determine errors for dependent parameters, such as `t_squared`, that are a function of `t`.

1.2.12 New Keywords

Typically keywords can be placed anywhere within an allowed scope. Due to the increasing number of keywords Version 6 introduces delimited keyword blocks which at present only applies to the `generate_stack_sequences` block. Keywords that are dependents of `generate_stack_sequences` can only be used within the `generate_stack_sequences` block. This avoids keyword name clashes and allows for keywords to be more readable. For example, `generate_stack_sequences` has the dependents of `ta`, `tb` and `tz` which are also keywords used by the `translate` keyword dependents. Use...
of these keywords however is possible as \textit{generate_stack_sequences} uses opening and closing curly brack delimiters; as follows:

\begin{verbatim}
generate_stack_sequences {
    ' generate_stack_sequences dependents placed here
    gauss_fwhm 0.02  ' non-generate_stack_sequences dependent
}
\end{verbatim}

Non-dependent keywords of \textit{generate_stack_sequences} can still be placed within the \textit{generate_stack_sequences} block.

New Version 6 keywords are as follows:

\begin{verbatim}
a_add
b_add
flip_neutron
generate_stack_sequences
height
layers_tol
match_transition_matrix_stats
n
number_of_sequences
number_of_stacks_per_sequence
num_runs
num_unique_vx_vy
n_avg
out_dependencies
out_dependents_for
out_file
pdf_convolute
pdf_data
pdf_scale_simple
peak_buffer_based_on
peak_buffer_based_on_tol
prm_with_error
save_sequences
save_sequences_as_strs
system_after_save_OUT
system_before_save_OUT
to
transition
use_layer
use_LU_for_errors
z_add
convolute_X_recal
pdf_for_pairs
pdf_gauss_fwhm
pdf_info
pdf_only_eq_0
pdf_ymin_on_ymax
\end{verbatim}
1.2.13 New Functions

Gamma_\(P(a, x)\): Returns the incomplete Gamma function \(P(a, x)\)

Gamma_\(Q(a, x)\): Returns the incomplete Gamma function \(Q(a, x) = 1 - P(a, x)\)

1.3 Stacking faults

```plaintext
[generate_stack_sequences] {
    [number_of_sequences !E]
    [number_of_stacks_per_sequence !E]
    [save_sequences $file]
    [save_sequences_as_strs $file]
    [layers_tol !E]
    [n_avg !E]
    [num_unique_vx_vy !N]
    [match_transition_matrix_stats { ... }]
    [transition $transition_name]...
        [use_layer $layer]
        [height E]
        [n !N]
    [to $to_transition_name !E]...
        [ta E] [tb E] [tz E]
        [a_add E] [b_add E] [z_add E]
}
' Get(generated_c)
```

Examples:

```plaintext
test_examples\stacking_faults\fit-1.inp
fit-2.inp
fit-3.inp

test_examples\stacking-faults\Rietveld-Generate\Rietveld-Generate\Create-Sequences.inp
Rietveld-Generate\Rietveld-Generate.inp
Rietveld-Generate\Fit-to-Rietveld-Generated.INP
Rietveld-Generate\Rietveld-Generated-200-2000.xy
Rietveld-Generate\strs-200-2000.txt
```

Stacking fault generation and refinement can now be performed at speeds that make routine analysis possible.

`generate_stack_sequences` generates sequences of stacks from the transition matrix described by the transition keywords. The opening and closing braces of {} corresponds to a block where keywords local to `generate_stack_sequences` can be used. Outside of the braces the `generate_stack_sequences` can’t be used. Get(generated_c) is updated after generation of the sequences with the average thickness of the sequences. It can be used to set the c lattice parameter.
num_unique_vx_vy: On termination of refinement the number of unique \{ sx, sy \} stacking vector coordinates is reported for all layer types.

transition: defines a ‘from’ transition with the name $transition_name. The transition uses the layer defined in use_layer.

to: defines the to-transition. $to_transition_name must be a defined $transition_name.

n returns the number of transitions generated for the corresponding to to-transition.

height: can be used instead of z_add keywords.

ta, tb: defines the stacking vector x and y coordinates in terms of the crystallographic \(a\) and \(b\) axes.

a_add, b_add: defines the stacking vector x and y coordinates relative to the previous stacking vector in terms of the crystallographic \(a\) and \(b\) axes.

tz: defines stacking vector z coordinate along the crystallographic \(c\) axis in Å.

add_z: defines stacking vector z coordinate along the crystallographic \(c\) axis in Å relative to the previous stacking vector.

1.3.1 Generating the same stacking sequences each run

To generate the same set of stacking sequences each run the random number generated can be seeded with a constant seed using seed, for example:

    seed #number

#number is a constant integer. Each #number generates its own unique set of random numbers. Generating identical sets of stacking sequences is useful when changes in \(R_{wp}\) that excludes stacking sequence variation is required.

1.3.2 The SF_Smooth macro

Stacking faulted calculated patterns can contain ripples when the peak shapes are small or when there’s too few layers stacked. The SF_Smooth macro, defined in TOPAS.INC smooths out these ripples such that small supercells can approximate large supercells; this crease computation speed and reduces memory usage. All stacking fault examples uses SF_Smooth; typical usage is as follows:

    SF_smooth(@, 1, 1)

The refined parameter adjust the width of a Gaussian convolution that is dependent on hkl's and the intensities of the reflections. The last argument (the ‘1’) can be used to adjusted the tolerance of a peak_buffer_based_on keyword used in SF_Smooth. The definition is as follows:

    peak_buffer_based_on = idl;
    peak_buffer_based_on_tol = Max(0.01 idl, Peak_Calculation_Step 0.5 s);

Reducing s increases the number of peaks in the peaks buffer and increases the accuracy of the calculated pattern. Typically s=1 is sufficient.
1.3.3  Fitting to DIFFaX test diamond data

Fit-1.INP uses `generate_stack_sequences` to fit to data generated from the DIFFaX suite (Treacy, 1991); INP segment that generates the sequences looks like:

```plaintext
generate_stack_sequences {
    number_of_sequences Nseqs 200
    number_of_stacks_per_sequence Nv 200
    num_unique_vx_vy 6
    Transition(1, lpc)
        to 1 = pa;   a_add = 2/3;  b_add = 1/3;   n !n1 349984
        to 2 = 1-pa; a_add = 0;    b_add = 0;     n !n2 149781
    Transition(2, lpc)
        to 1 = 1-pa; a_add = 0;    b_add = 0;     n !n3 149781
        to 2 = pa;   a_add = -2/3; b_add = -1/3;  n !n4 350254
}
```

The generated probability parameter `pa` can be determined using the `n` values as follows:

```plaintext
prm !pa_gen = (n1+n3)/(n1+n2+n3+n4); : 0.699974874
```

The fit to the DIFFaX data looks like:

![Fit-1.INP, Fitting to DIFFaX test diamond data](image)

1.3.4  Stacking faults from layers of different layer heights

Layers of different thicknesses can be modelled accurately and refinement fast. Here’s a fit to simulated data (FIT-2.INP) for two different layer heights of 5 and 6Å.
1.3.5 Rietveld-Generated example

The files in the test_examples\stacking-faults\Rietveld-Generate\ directory can be used to create a stacking faulted test pattern using Rietveld refinement. The test pattern can then be fitted to. Create-Sequences.inp created the INP format stacking sequences and places the result int the file strs-200-2000.txt. The file Rietveld-Generate.inp can then be used to create the test pattern Rietveld-Generated-200-2000.xy. This test pattern can be fitted to using the file Fit-to-Rietveld-Generated.INP; this file uses generate_stack_sequences. It demonstrates the accuracy and speed of the stacking fault averaging fitting procedure. The fit to the Rietveld generated stacking faulted pattern looks like:

1.3.6 Refining on layer heights

Layer heights can be refined by refining on parameters that are a function of the add_z or height keywords. The Fit-3.INP example refines on 3 height parameters as well as the z fractional atomic coordinates of the sites that comprise the layers. It also lists six types of transitions which operate on three unique layer types. The transitions point to the layer types using the use_layer keyword. The c lattice parameter is defined and refined using the following:

```plaintext
prm qq 0 c = Get(generated_c) + 0.0001 qq; : 1828.085117
```

Get(generated_c) is also used to initialize the z fractional coordinates of the sites as follows:
The fit to the test data looks like:

![Fit-3.INP, Refining on stacking vector and structural parameters](image_url)

1.4 PDF refinement

PDF patterns are treated as an ‘xDD’. Many keywords used for xdds can also be used in PDF refinement. PDF patterns can be refined simultaneously with other types of xdd patterns comprising x-ray dependent or x-ray independent phases or peaks phases. Penalties, restraints and in particular keywords such as rigid, atomic_interaction, sites_geometry, sites_distance etc... can all be used. Data structure items frequently used in PDF refinement are:

```plaintext
xdd...
  pdf_data
  *[scale_phase_X E]...
  *[fit_obj]...
  [start_X #] [finish_X #]
  *[rebin_with_dx_of !E] [rebin_start_x_at !E]
  *[weighting !E]
  *[Tpdf_convolute]...
str...
  *[scale_phase_X E]...
  [scale E]
  [view_structure]
  [rigid]...
  [occ_merge $sites]...
  [pdf_scale_simple]
  *[pdf_zero E]
  [pdf_ymin_on_ymax 0.001]
  [pdf_info]
  *[Tpdf_convolute]...
  [pdf_for_pairs $sites_1 $sites_2]...
    [pdf_only_eq_0]
    *[pdf_gauss_fwhm E]
```
where

\[ T_{pdf\_convolute} \]

\[ [pdf\_convolute \ E] \]

\[ [min\_X \ !E] \]

\[ [max\_X \ !E] \]

\[ [convolute\_X\_recal \ !E] \]

Examples files in directory test_examples\pdf

beq-2.inp
beq-2-create.inp
beq-3.inp
beq-3-create.inp
pdf-1.inp
df-2.inp
alvo4\structure-solution-create.inp
alvo4\structure-solution.inp
alvo4\rigid.inp
occ-merge-pbso4\create.inp
occ-merge-pbso4\occ-merge-test.inp
occ-merge-pbso4\occ-merge.inp

Data files in directory test_examples\pdf

beq-2.xy
beq-3.xy
alvo4\alvo4.xy
occ-merge-pbso4\pbso4.xy

Keywords with '*' next to them can be a function of the x-axis reserved parameter name X; for PDF data X corresponds to r.

\textbf{pdf\_data} tells the program that the data set is of G(r) type.

Let’s write G(r) as follows:

\[ G(r) = s1 \frac{S(r)}{r} - s2 \]

where

r corresponds to the X-axis

s1 and s2 are constants

S(r) are the pairs

\textbf{pdf\_scale\_simple} tells the program to calculate S(r)/(Np r) only.

\textbf{pdf\_ymin\_on\_ymax} defines the minimum/maximum value for the Gaussians in regards to the x-axis range calculated for the Gaussians; the default value of 0.001 in typically sufficient.
*pdf_for_pairs* can be used to select site pairs using the site name; for example:

```
pdf_for_pairs "V* Al* !O2" *
```

The ‘!’ character excludes the sequence from the wild card string. Multiple *pdf_for_pairs* can be defined.

*pdf_only_eq_0* informs the parent *pdf_for_pairs* that only equivalent position 0 is to be considered.

*pdf_gauss_fwhm* is used to write the width equation for the pairs selected by *pdf_for_pairs*. If all of the pairs possible are described by *pdf_for_pairs* then the associated *beq’s* are not used and they become redundant. The user is informed of unused *beq’s*. Consider the following abbreviated INP segment:

```
site Al1 ... beq 1
site O1 ... beq 1
pdf_for_pairs Al1 Al1 pdf_only_eq_0 pdf_gauss_fwhm 0.1 ' Line A
pdf_for_pairs Al1 O1 pdf_only_eq_0 pdf_gauss_fwhm 0.2 ' Line B
pdf_for_pairs Al1 O1 pdf_gauss_fwhm 0.3 ' Line C
```

There are a number of types of interactions with FWHMs as follows:

- Al1↔O1 interactions for equivalent position 0 described using Line B
- Al1↔O1 interactions excluding equivalent position zero described using Line C
- O1↔O1 interactions described using *beq’s*

*pdf_info* displays the interactions used in matrix form; for the above INP segment we get:

```
pdf_info
{
- = No pdf_for_pairs defined hence *beq’s* used
 0 = pdf_for_pairs defined with *pdf_only_eq_0*
 1 = pdf_for_pairs defined without *pdf_only_eq_0*
 2 = two pdf_for_pairs defined, one with *pdf_only_eq_0* and one without *pdf_only_eq_0*

       Al1  -2
       O1   2-
}
```

The matrix is shown in blue. *pdf_for_pairs* together with *beq’s* defaults offers great flexibility in describing peak widths. See PDF-1.INP, PDF-2.INP, BEQ-3.INP.

*scale_phase_X* can be used to describe Gaussian dampening, for example:

```
prm damp_fwhm 50 min 1e-6 max 200
prm damp = Gauss(0, damp_fwhm);
scale_phase_X = damp;
```
1.4.1 Inter and Intra molecule FWHMs

Assigning different interaction types for molecules is done using `pdf_for_pairs`. For example, to set the bond lengths for the atom Al1 of AlVO4 (see PDF-2.INP) to a different FWHM for equivalent position 0 the following could be used:

```plaintext
prm intra_molec 0.01 min 1e-6
pdf_for_pairs Al1 "01 02 03 04 05 06" pdf_only_eq_0
    pdf_gauss_fwhm = intra_molec;
```

The calculated pattern from PDF-2.INP therefore becomes:

Notice the 6 spikes; they correspond to the Al1 bonds which has narrow FWHMs. If we then wanted Al1 bonds that are not equivalent position 0 to be different to the `beq`'s values then we could use:

```plaintext
prm inter_molec 0.1 min 1e-6
prm intra_molec 0.01 min 1e-6
pdf_for_pairs Al1 "01 02 03 04 05 06" pdf_only_eq_0
    pdf_gauss_fwhm = intra_molec;
pdf_for_pairs Al1 "01 02 03 04 05 06"
    pdf_gauss_fwhm = inter_molec;
```

This gives the following calculated pattern where we see the different bonds of Al1.

The corresponding output from `pdf_info` is:

```plaintext
pdf_info
{
```

26
A11 ------222222------
A12 -------------------
A13 -------------------
V1 -------------------
V2 -------------------
V3 -------------------
O1 2-------------------
O2 2-------------------
O3 2-------------------
O4 2-------------------
O5 2-------------------
O6 2-------------------
O7 -------------------
O8 -------------------
O9 -------------------
O10 -------------------
O11 -------------------
O12 -------------------
}

An exception is thrown if the same interaction is referenced in more than one pdf_for_pairs, for example, the following will throw an exception as A11→O1 is referenced twice:

```
pdf_for_pairs A11 "O1 O2 O3 O4 O5 O6" pdf_only_eq_0 ...
pdf_for_pairs A11 O1 pdf_only_eq_0 ...
```

The following will not throw an exception:

```
pdf_for_pairs A11 "O1 O2 O3 O4 O5 O6" pdf_only_eq_0 ...
pdf_for_pairs A11 O1 ...
```

### 1.4.2 Instrument Sinc function Sinc-1.INP

In Sinc-1.PDF pdf_convolute is used at the xdd level to convolute a Sinc function into pdf type phases as follows:

```
pdf_convolute = Sin(Qmax X+q3)/If(Abs(X)<0.5 Step_Size, If(X<0,-q2,q2), X);
min_X = -conv_max;
max_X = conv_max;
```

Sinc-1.INP also uses an xo_Is phase defined as:

```
xo_Is
NoThDependence(0.0001)
xo 10 I @ 100
peak_type pv
```
pdf_convolute operates on pdf type phases only; hence the xo_ls phase is untouched. Note the phase dependent use of an emission profile as defined in the NoThDependence macro. Multiple pdf_convolute’s can be described at the global, xdd, str and pdf_for_pairs levels. Note, use of pdf_convolute as a dependent of pdf_for_pairs is slower than at the other levels; thus where possible use pdf_convolute at non-pdf_for_pairs levels.

1.4.3 Weighting of PDF and 2-Theta type data

PDF and 2-Theta data can be of very different intensities; the xdd_sum keyword can assist in modifying the weighing of data in order to give the patterns approximately similar weights. For example:

```plaintext
xdd file1.xy
    xdd_sum !sum1 = Abs(Yobs);
    weighting = 1/sum2;
xDD file2.xy
    xdd_sum !sum2 = Abs(Yobs);
    weighting = 1/sum2;
```

1.4.4 Test_examples\pdf\BEQ-2.INP

Use Test_examples\pdf\BEQ-2-create.INP to generate a simulated pattern for BEQ-2.INP

BEQ-2.INP

- Comprise the structure of AIVO4
- 3 types of beq parameters
- beq is a function of X (ie. X corresponds to the X-axis which is r) and hence peak widths are a function of X.
- demonstrates the use of pdf_zero
- demonstrates the use of rebin_with_dx_of and rebin_start_x_at

1.4.5 Test_examples\pdf\BEQ-3.INP

Use Test_examples\pdf\BEQ-3-create.INP to generate a simulated pattern for BEQ-3.INP

BEQ-3.INP

- Demonstrates the use of pdf_for_pairs

1.4.6 Speeding up refinement with rebin_with_dx_of

The step size in PDF data must be of equal size. Also, the start of the x-axis needs to be an integral multiple of the step size. Increasing the step size in the data speeds up refinement; see BEQ-2.INP. The step size can be increased using:

```plaintext
macro Rebin_Step   { 0.015 }
rebin_with_dx_of Rebin_Step rebin_start_x_at Rebin_Step
```
Rebinning is akin to collecting the data at a larger step size. All data is included with counts after rebinning being equal to counts before rebinning. esds associated with the data are also rebinned.

rebin_start_x_at can be used to place the start of the data at an integral multiple of the step size. In BEQ-2.INP parameters such as scale are written in terms of the rebin step size to reflect the fact that the scaling of the data is changed due to rebinning.

1.4.7 Refining on beq parameters

Modify the BB macro so that it comprises:

```
macro BB { } ' beq, Insert/remove !
```

Gives an Rwp plot of:

This type of convergence is indicative of derivatives being calculated correctly. Convergence for coordinates, occupancies, lattice parameters and pdf_zero are similar.

1.4.8 Structure Solution, Simulated Annealing

test_examples\pdf\alvo4\structure-solution-create.inp creates a simulated pattern for structure-solution.inp. It’s a simulated annealing refinement with all coordinates starting at zero with anti-bump penalties applied using:

```
AI_Anti_Bump(O*, O*, 2.4, 1, 5)
AI_Anti_Bump(Al*, O*, 1.6, 1, 5)
AI_Anti_Bump(Al*, Al*, 2.8, 1, 5)
```

The correct solution is found as seen in the following:
The range of convergence however for coordinates are smaller than with reciprocal space as in normal Rietveld refinement. This is because the coordinates in the PDF case change peak positions rather that peak intensities with the former having a narrow range of convergence. It may be possible to increase the range of convergence for PDF by increasing the peak widths but this comes at the expense of resolution and it may result in an even smaller range of convergence.

1.4.9 Rigid bodies with PDF data

test_examples\pdf\alvo4\rigid.inp operates on the simulated data created by structure-solution-create.inp. It demonstrates the use of rigid bodies with PDF data.

1.4.10 Occupancy merging with PDF data

test_examples\pdf\occ-merge-PbSO4\occ-merge.inp operates on simulated data created by create.inp. It demonstrates the use of occ_merge with PDF data.

1.4.11 Equivalence of pdf_gauss_fwhm and beq when there’s one atom type

test_examples\pdf\si1.inp comprises an option to use beq or pdf_gauss_fwhm as follows:

For the beq case we have:

\[
\text{beq} = \text{width};
\]

and for pdf_gauss_fwhm we have:

\[
\text{pdf}_\text{gauss}_\text{fwhm} = \sqrt{\text{width} \cdot \text{Ln}(2) / \pi^2};
\]

The above cases are equivalent when there’s one atom type.
2 NEW GUI FUNCTIONALITY

2.1 TOF x-axis can be displayed in d-spacing, Q and tof

The x-axis of TOF data can be displayed as either tof, d-spacing or Q by cycling the x-axis button:

2.2 Displaying many files at once

See files in the directory test_examples\3d\.

2.2.1 Surface plots – 2D with offsets

Scans can be displayed and offset from one another using the icon, for example:

The Quickzoom window is operational in all 2D-offset plots.
Pressing the **Middle Mouse Button** and moving the mouse changes the x and y offsets. This movement greatly assists in determining the curvature of the surface. The QuickZoom display is not offset allowing for two views of the same data.

### 2.2.2 Inserting peaks and identifying scans

Peak can be inserted by pressing the **Ctrl-Key** and clicking the RMB. When the Ctrl key is pressed a solid circle is displayed on the scan closest to the mouse. The circle is coloured to match the scan lines and in addition the closest scan is displayed with a thickened line. Displayed at the bottom of the plot is the name of the scan as seen by the arrow below. Peaks as well as excluded regions move with the offsets.

When the Ctrl-Key is pressed the x and y axis values displayed on the status line are offset to match the closest scan. Similarly when “For LAM Cursor” option is selected the LAM cursor is changed to match the axis of the closest scan.

### 2.2.3 2D-offset Surface plots

2D-offset plot can be displayed as a 3D-Surface, for example:
These plots can be manipulated in real time; the 871 file `test_examples\je-para\d8_02999.raw` with over 4 million data points can be easily manipulated:

Pressing the Shift key whilst performing a Zoom (forming a box using the mouse) zooms into a region. Zooming in this manner deselects scans for display. An unzoom is performed by performing an Unzoom whilst holding down the Shift key. Colour schemes can be changed by using the Colours options:

Contour-Orange-15 for looks like:
2.2.4 2D-offset Planview plots

Moving the y-offset such that it's at a maximum automatically produces a Planview; a Kaleidoscope colour scheme gives:

The Standard colour scheme gives:

Zooming gives:
Planview can also have x-axis offsets with line scans overlain:

These line scans can include the calculated and/or difference patterns as well as patterns for individual phases. Beneath the displayed line scans are their shadows. Colours are blended across scans as well as across the x-axis to sharpen images.

### 2.2.5 OpenGL Surface plots

OpenGL surface plots can be displayed alongside 2D-offset plots:
The scans displayed in the chart area are displayed to the right as a surface plot. Use RMB on the surface plot for options; these are:

- Use the Mouse Wheel to scroll the x-axis from either the 2D or 3D plots.
- RMB-Pressed and moving zooms
- Pressing ‘x’ whilst rotating allows rotation around an axis vertical to the screen.
- Pressing ‘y’ whilst rotating allows rotation around an axis horizontal to the screen.
- Pressing ‘z’ whilst rotating allows rotation around an axis perpendicular to the screen.
- Pressing the Mouse Wheel button (as opposed to rotating the mouse wheel) moves the object and hence the centre of rotation.
- When the Mouse is close to the Left or Right borders of the OpenGL window then rotation is around an axis perpendicular to the computer screen. Very useful for positioning 3D objects.

Opening the OpenGL Text Dialog and clicking on the 3D surface writes text into the Text Dialog; this text comprises the names of the two files bordering the polygon that has been clicked and the average x and y values of the polygon, for example:

2.2.6 **OpenGL – Weighted difference for colours**

The RMB “Weight difference for colours” option displays colours corresponding to the

\[
\text{WtDiff} = \frac{\text{Abs}(Yobs-Ycalc)}{\text{Weighting}}
\]
2.3 Normalizing scans to the maximum scan value within a Scan Window

Displayed scans can be normalized using the option “Yobs Normalize” which is activated using the RMB on the Scan window. Normalizing scales displayed scans such that the maximum values of the displayed data are all equal. Normalizing is temporary and can be toggled on/off by executing the “Yobs Normalize”. The following shows scans normalized with all the peaks on the right having the same height.

References