

```
#####  
### FullProf-generated CIF output file (version: February 2008) ###  
### Template of CIF submission form for structure report ###  
#####
```

```
# This file has been generated using FullProf.2k taking one example of  
# structure report provided by Acta Cryst. It is given as a 'template' with  
# filled structural items. Many other items are left unfilled and it is the  
# responsibility of the user to properly fill or suppress them. In principle  
# all question marks '?' should be replaced by the appropriate text or  
# numerical value depending on the kind of CIF item.  
# See the document: cif_core.dic (URL: http://www.iucr.org) for details.
```

```
# Please notify any error or suggestion to:  
# Juan Rodriguez-Carvajal (jrc@ill.eu)  
# Improvements will be progressively added as needed.
```

```
#=====
```

data_global

```
#=====
```

PROCESSING SUMMARY (IUCr Office Use Only)

```
_journal_data_validation_number    ?  
  
_journal_date_recd_electronic      ?  
_journal_date_to_coeditor          ?  
_journal_date_from_coeditor        ?  
_journal_date_accepted              ?  
_journal_date_printers_first       ?  
_journal_date_printers_final       ?  
_journal_date_proofs_out           ?  
_journal_date_proofs_in            ?  
_journal_coeditor_name              ?  
_journal_coeditor_code              ?  
_journal_coeditor_notes  
; ?  
;  
_journal_techeditor_code            ?  
_journal_techeditor_notes  
; ?  
;  
_journal_coden_ASTM                 ?  
_journal_name_full                   ?  
_journal_year                         ?  
_journal_volume                       ?  
_journal_issue                         ?  
_journal_page_first                  ?  
_journal_page_last                   ?  
_journal_paper_category              ?  
_journal_suppl_publ_number           ?  
_journal_suppl_publ_pages            ?
```

#=====

1. SUBMISSION DETAILS

_publ_contact_author_name ? # Name of author for correspondence
_publ_contact_author_address # Address of author for correspondence
; ?
;
_publ_contact_author_email ?
_publ_contact_author_fax ?
_publ_contact_author_phone ?

_publ_contact_letter
; ?
;

_publ_requested_journal ?
_publ_requested_coeditor_name ?
_publ_requested_category ? # Acta C: one of CI/CM/CO/FI/FM/FO

Definition of non standard CIF items (Reliability indices used in FULLPROF)

loop_
_publ_manuscript_incl_extra_item
_publ_manuscript_incl_extra_info
_publ_manuscript_incl_extra_defn
Name Explanation Standard?

'_pd_proc_ls_prof_cR_factor' 'Prof. R-factor CORRECTED for background' no
'_pd_proc_ls_prof_cwR_factor' 'wProf.R-factor CORRECTED for background' no
'_pd_proc_ls_prof_cwR_expected' 'wProf.Expected CORRECTED for background' no
'_pd_proc_ls_prof_chi2' 'Chi-square for all considered points' no
'_pd_proc_ls_prof_echi2' 'Chi-2 for points with Bragg contribution' no

#=====

3. TITLE AND AUTHOR LIST

_publ_section_title
; 'NaCo1'
;
_publ_section_title_footnote
;
;

The loop structure below should contain the names and addresses of all
authors, in the required order of publication. Repeat as necessary.

loop_
_publ_author_name
_publ_author_footnote
_publ_author_address
? #<--'Last name, first name'
; ?

;
; ?
;

#=====

4. TEXT

_publ_section_synopsis

; ?

;

_publ_section_abstract

; ?

;

_publ_section_comment

; ?

;

_publ_section_exptl_prep # Details of the preparation of the sample(s)
should be given here.

; ?

;

_publ_section_exptl_refinement

; ?

;

_publ_section_references

; ?

;

_publ_section_figure_captions

; ?

;

_publ_section_acknowledgements

; ?

;

#=====

#=====

If more than one structure is reported, the remaining sections should be
completed per structure. For each data set, replace the '?' in the
data_? line below by a unique identifier.

data_Co10Ge3O16

#=====

5. CHEMICAL DATA

_chemical_name_systematic

; ?

;

_chemical_name_common ?

_chemical_formula_moiety ?

_chemical_formula_structural ?

_chemical_formula_analytical ?

_chemical_formula_iupac ?
_chemical_formula_sum ?
_chemical_formula_weight ?
_chemical_melting_point ?
_chemical_compound_source ? # for minerals and
natural products

loop_
_atom_type_symbol
_atom_type_scatter_length_neutron
_atom_type_scatter_source
Fe 0.94500 V.F._Sears_Neutron_News_3_26_(1992)
Ge 0.81850 V.F._Sears_Neutron_News_3_26_(1992)
O 0.58030 V.F._Sears_Neutron_News_3_26_(1992)

#=====

6. POWDER SPECIMEN AND CRYSTAL DATA

_symmetry_cell_setting Monoclinic
_symmetry_space_group_name_H-M 'C 1 2/c 1'
_symmetry_space_group_name_Hall '-C 2yc'

loop_
_symmetry_equiv_pos_as_xyz #<--must include 'x,y,z'
'x,y,z'
'-x,y,-z+1/2'
'-x,-y,-z'
'x,-y,z+1/2'
'x+1/2,y+1/2,z'
'-x+1/2,y+1/2,-z+1/2'
'-x+1/2,-y+1/2,-z'
'x+1/2,-y+1/2,z+1/2'

_cell_length_a 9.78383(5)
_cell_length_b 9.13493(5)
_cell_length_c 5.19289(3)
_cell_angle_alpha 90.00000
_cell_angle_beta 101.7500(4)
_cell_angle_gamma 90.00000
_cell_volume 454.387(4)
_cell_formula_units_Z ?
_cell_measurement_temperature ?
_cell_special_details

; ?

;

The next three fields give the specimen dimensions in mm. The equatorial
plane contains the incident and diffracted beam.

_pd_spec_size_axial ? # perpendicular to
equatorial plane
_pd_spec_size_equat ? # parallel to
scattering vector
in transmission

```
_pd_spec_size_thick      ?    # parallel to
                          # scattering vector
                          # in reflection
```

The next five fields are character fields that describe the specimen.

```
_pd_spec_mounting      # This field should be
                        # used to give details of the
                        # container.
```

```
; ?
;
```

```
_pd_spec_mount_mode    ?    # options are 'reflection'
                        # or 'transmission'
```

```
_pd_spec_shape        ?    # options are 'cylinder'
                        # 'flat_sheet' or 'irregular'
```

```
_pd_char_particle_morphology ?
```

```
_pd_char_colour       ?    # use ICDD colour descriptions
```

The following three fields describe the preparation of the specimen.

The cooling rate is in K/min. The pressure at which the sample was

prepared is in kPa. The temperature of preparation is in K.

```
_pd_prep_cool_rate     ?
```

```
_pd_prep_pressure     ?
```

```
_pd_prep_temperature  ?
```

The next four fields are normally only needed for transmission experiments.

```
_exptl_absorpt_coefficient_mu ?
```

```
_exptl_absorpt_correction_type ?
```

```
_exptl_absorpt_process_details ?
```

```
_exptl_absorpt_correction_T_min ?
```

```
_exptl_absorpt_correction_T_max ?
```

```
#=====
```

7. EXPERIMENTAL DATA

```
_exptl_special_details
```

```
; ?
;
```

The following item is used to identify the equipment used to record

the powder pattern when the diffractogram was measured at a laboratory

other than the authors' home institution, e.g. when neutron or synchrotron

radiation is used.

```
_pd_instr_location
```

```
; ?
;
```

```
_pd_calibration_special_details # description of the method used
                                # to calibrate the instrument
```

```
; ?
;
```

_diffrn_ambient_temperature ?
_diffrn_source 'nuclear reactor'
_diffrn_radiation_type 'Constant Wavelength Neutron Diffraction'
_diffrn_radiation_wavelength 2.53600
_diffrn_source_type ? # Put here the diffractometer and site

_diffrn_radiation_monochromator ?
_diffrn_measurement_device_type ?
_diffrn_measurement_method ?
_diffrn_detector_area_resol_mean ? # Not in version 2.0.1
_diffrn_detector ?
_diffrn_detector_type ? # make or model of detector
_pd_meas_scan_method ? # options are 'step', 'cont',
'tof', 'fixed' or
'disp' (= dispersive)
_pd_meas_special_details
; ?
;

The following four items give details of the measured (not processed)
powder pattern. Angles are in degrees.

_pd_meas_number_of_points 3020
_pd_meas_2theta_range_min 0.95000
_pd_meas_2theta_range_max 151.89999
_pd_meas_2theta_range_inc 0.050017

#=====

8. REFINEMENT DATA

_refine_special_details
; ?
;

Use the next field to give any special details about the fitting of the
powder pattern.

_pd_proc_ls_special_details
; ?
;

The next three items are given as text.

_pd_proc_ls_profile_function ?
_pd_proc_ls_background_function ?
_pd_proc_ls_pref_orient_corr
; ?
;

The following profile R-factors are NOT CORRECTED for background
The sum is extended to all non-excluded points.
These are the current CIF standard

_pd_proc_ls_prof_R_factor 3.8219
_pd_proc_ls_prof_wR_factor 5.1178
_pd_proc_ls_prof_wR_expected 2.2903

The following profile R-factors are CORRECTED for background
The sum is extended to all non-excluded points.
These items are not in the current CIF standard, but are defined above

_pd_proc_ls_prof_cR_factor 7.4371
_pd_proc_ls_prof_cwR_factor 7.4087
_pd_proc_ls_prof_cwR_expected 3.3155

The following items are not in the CIF standard, but are defined above

_pd_proc_ls_prof_chi2 4.9932
_pd_proc_ls_prof_echi2 5.3327

Items related to LS refinement

_refine_ls_R_I_factor 1.2079
_refine_ls_number_reflns 106
_refine_ls_number_parameters 94
_refine_ls_number_restraints 0

The following four items apply to angular dispersive measurements.
2theta minimum, maximum and increment (in degrees) are for the
intensities used in the refinement.

_pd_proc_2theta_range_min 0.8918
_pd_proc_2theta_range_max 151.8418
_pd_proc_2theta_range_inc 0.050017
_pd_proc_wavelength 2.536000

_pd_block_diffraction_id ? # The id used for the block containing
the powder pattern profile (section 11)

Give appropriate details in the next two text fields.

_pd_proc_info_excluded_regions ?
_pd_proc_info_data_reduction ?

The following items are used to identify the programs used.

_computing_data_collection ?
_computing_structure_solution ?
_computing_structure_refinement FULLPROF
_computing_molecular_graphics ?
_computing_publication_material ?

#=====

9. ATOMIC COORDINATES AND DISPLACEMENT PARAMETERS

```

loop_
  _atom_site_label
  _atom_site_fract_x
  _atom_site_fract_y
  _atom_site_fract_z
  _atom_site_U_iso_or_equiv
  _atom_site_occupancy
  _atom_site_adp_type          # Not in version 2.0.1
  _atom_site_type_symbol
Fe1 0.00000 0.9086(3) 0.25000 0.0020(7) 1.00000 Uiso Fe
Fe2 0.00000 0.2706(2) 0.25000 0.0029(8) 1.00000 Uiso Fe
Ge1 0.30032(16) 0.0924(2) 0.2159(3) 0.0039(6) 1.00000 Uiso Ge
O1 0.1188(2) 0.0905(3) 0.1403(5) 0.0056(7) 1.00000 Uiso O
O2 0.3841(3) 0.2406(3) 0.3807(6) 0.0017(8) 1.00000 Uiso O
O3 0.3589(2) 0.0657(3) 0.9130(4) 0.0016(8) 1.00000 Uiso O

```

Note: if the displacement parameters were refined anisotropically
the U matrices should be given as for single-crystal studies.

#=====

10. DISTANCES AND ANGLES / MOLECULAR GEOMETRY

_geom_special_details ?

```

loop_
  _geom_bond_atom_site_label_1
  _geom_bond_atom_site_label_2
  _geom_bond_site_symmetry_1
  _geom_bond_site_symmetry_2
  _geom_bond_distance
  _geom_bond_publ_flag
? ? ? ? ? ?

```

```

loop_
  _geom_contact_atom_site_label_1
  _geom_contact_atom_site_label_2
  _geom_contact_distance
  _geom_contact_site_symmetry_1
  _geom_contact_site_symmetry_2
  _geom_contact_publ_flag
? ? ? ? ? ?

```

```

loop_
  _geom_angle_atom_site_label_1
  _geom_angle_atom_site_label_2
  _geom_angle_atom_site_label_3
  _geom_angle_site_symmetry_1
  _geom_angle_site_symmetry_2
  _geom_angle_site_symmetry_3
  _geom_angle
  _geom_angle_publ_flag
? ? ? ? ? ? ? ?

```

```
loop_
_geom_torsion_atom_site_label_1
_geom_torsion_atom_site_label_2
_geom_torsion_atom_site_label_3
_geom_torsion_atom_site_label_4
_geom_torsion_site_symmetry_1
_geom_torsion_site_symmetry_2
_geom_torsion_site_symmetry_3
_geom_torsion_site_symmetry_4
_geom_torsion
_geom_torsion_publ_flag
? ? ? ? ? ? ? ? ? ?
```

```
loop_
_geom_hbond_atom_site_label_D
_geom_hbond_atom_site_label_H
_geom_hbond_atom_site_label_A
_geom_hbond_site_symmetry_D
_geom_hbond_site_symmetry_H
_geom_hbond_site_symmetry_A
_geom_hbond_distance_DH
_geom_hbond_distance_HA
_geom_hbond_distance_DA
_geom_hbond_angle_DHA
_geom_hbond_publ_flag
? ? ? ? ? ? ? ? ? ?
```

```
#=====
```

```
#=====
```

```
# Additional structures (last six sections and associated data_? identifiers)
# may be added at this point.
```

```
#=====
```

```
data_Co10Ge3O16
```

```
#=====
```

```
# 5. CHEMICAL DATA
```

```
_chemical_name_systematic
; ?
;
_chemical_name_common ?
_chemical_formula_moiety ?
_chemical_formula_structural ?
_chemical_formula_analytical ?
_chemical_formula_iupac ?
_chemical_formula_sum ?
_chemical_formula_weight ?
_chemical_melting_point ?
_chemical_compound_source ? # for minerals and
# natural products
```

```
loop_
  _atom_type_symbol
  _atom_type_scatter_length_neutron
  _atom_type_scatter_source
```

```
#=====
```

6. POWDER SPECIMEN AND CRYSTAL DATA

```
_symmetry_cell_setting      Cubic
_symmetry_space_group_name_H-M  'F d -3 m'
_symmetry_space_group_name_Hall  '-F 4vw 2vw 3'
```

```
loop_
  _symmetry_equiv_pos_as_xyz  #<--must include 'x,y,z'
  'x,y,z'
  'x,-y+1/4,-z+1/4'
  '-x+1/4,y,-z+1/4'
  '-x+1/4,-y+1/4,z'
  'y,z,x'
  '-y+1/4,-z+1/4,x'
  'y,-z+1/4,-x+1/4'
  '-y+1/4,z,-x+1/4'
  'z,x,y'
  '-z+1/4,x,-y+1/4'
  '-z+1/4,-x+1/4,y'
  'z,-x+1/4,-y+1/4'
  'y,x,z'
  '-y+1/4,x,-z+1/4'
  'y,-x+1/4,-z+1/4'
  '-y+1/4,-x+1/4,z'
  'z,y,x'
  '-z+1/4,-y+1/4,x'
  '-z+1/4,y,-x+1/4'
  'z,-y+1/4,-x+1/4'
  'x,z,y'
  'x,-z+1/4,-y+1/4'
  '-x+1/4,-z+1/4,y'
  '-x+1/4,z,-y+1/4'
  '-x,-y,-z'
  '-x,y+3/4,z+3/4'
  'x+3/4,-y,z+3/4'
  'x+3/4,y+3/4,-z'
  '-y,-z,-x'
  'y+3/4,z+3/4,-x'
  '-y,z+3/4,x+3/4'
  'y+3/4,-z,x+3/4'
  '-z,-x,-y'
  'z+3/4,-x,y+3/4'
  'z+3/4,x+3/4,-y'
  '-z,x+3/4,y+3/4'
  '-y,-x,-z'
  'y+3/4,-x,z+3/4'
  '-y,x+3/4,z+3/4'
```

'y+3/4,x+3/4,-z'
'-z,-y,-x'
'z+3/4,y+3/4,-x'
'z+3/4,-y,x+3/4'
'-z,y+3/4,x+3/4'
'-x,-z,-y'
'-x,z+3/4,y+3/4'
'x+3/4,z+3/4,-y'
'x+3/4,-z,y+3/4'
'x+1/2,y+1/2,z'
'x+1/2,-y+3/4,-z+1/4'
'-x+3/4,y+1/2,-z+1/4'
'-x+3/4,-y+3/4,z'
'y+1/2,z+1/2,x'
'-y+3/4,-z+3/4,x'
'y+1/2,-z+3/4,-x+1/4'
'-y+3/4,z+1/2,-x+1/4'
'z+1/2,x+1/2,y'
'-z+3/4,x+1/2,-y+1/4'
'-z+3/4,-x+3/4,y'
'z+1/2,-x+3/4,-y+1/4'
'y+1/2,x+1/2,z'
'-y+3/4,x+1/2,-z+1/4'
'y+1/2,-x+3/4,-z+1/4'
'-y+3/4,-x+3/4,z'
'z+1/2,y+1/2,x'
'-z+3/4,-y+3/4,x'
'-z+3/4,y+1/2,-x+1/4'
'z+1/2,-y+3/4,-x+1/4'
'x+1/2,z+1/2,y'
'x+1/2,-z+3/4,-y+1/4'
'-x+3/4,-z+3/4,y'
'-x+3/4,z+1/2,-y+1/4'
'-x+1/2,-y+1/2,-z'
'-x+1/2,y+1/4,z+3/4'
'x+1/4,-y+1/2,z+3/4'
'x+1/4,y+1/4,-z'
'-y+1/2,-z+1/2,-x'
'y+1/4,z+1/4,-x'
'-y+1/2,z+1/4,x+3/4'
'y+1/4,-z+1/2,x+3/4'
'-z+1/2,-x+1/2,-y'
'z+1/4,-x+1/2,y+3/4'
'z+1/4,x+1/4,-y'
'-z+1/2,x+1/4,y+3/4'
'-y+1/2,-x+1/2,-z'
'y+1/4,-x+1/2,z+3/4'
'-y+1/2,x+1/4,z+3/4'
'y+1/4,x+1/4,-z'
'-z+1/2,-y+1/2,-x'
'z+1/4,y+1/4,-x'
'z+1/4,-y+1/2,x+3/4'
'-z+1/2,y+1/4,x+3/4'
'-x+1/2,-z+1/2,-y'

'-x+1/2,z+1/4,y+3/4'
'x+1/4,z+1/4,-y'
'x+1/4,-z+1/2,y+3/4'
'x+1/2,y,z+1/2'
'x+1/2,-y+1/4,-z+3/4'
'-x+3/4,y,-z+3/4'
'-x+3/4,-y+1/4,z+1/2'
'y+1/2,z,x+1/2'
'-y+3/4,-z+1/4,x+1/2'
'y+1/2,-z+1/4,-x+3/4'
'-y+3/4,z,-x+3/4'
'z+1/2,x,y+1/2'
'-z+3/4,x,-y+3/4'
'-z+3/4,-x+1/4,y+1/2'
'z+1/2,-x+1/4,-y+3/4'
'y+1/2,x,z+1/2'
'-y+3/4,x,-z+3/4'
'y+1/2,-x+1/4,-z+3/4'
'-y+3/4,-x+1/4,z+1/2'
'z+1/2,y,x+1/2'
'-z+3/4,-y+1/4,x+1/2'
'-z+3/4,y,-x+3/4'
'z+1/2,-y+1/4,-x+3/4'
'x+1/2,z,y+1/2'
'x+1/2,-z+1/4,-y+3/4'
'-x+3/4,-z+1/4,y+1/2'
'-x+3/4,z,-y+3/4'
'-x+1/2,-y,-z+1/2'
'-x+1/2,y+3/4,z+1/4'
'x+1/4,-y,z+1/4'
'x+1/4,y+3/4,-z+1/2'
'-y+1/2,-z,-x+1/2'
'y+1/4,z+3/4,-x+1/2'
'-y+1/2,z+3/4,x+1/4'
'y+1/4,-z,x+1/4'
'-z+1/2,-x,-y+1/2'
'z+1/4,-x,y+1/4'
'z+1/4,x+3/4,-y+1/2'
'-z+1/2,x+3/4,y+1/4'
'-y+1/2,-x,-z+1/2'
'y+1/4,-x,z+1/4'
'-y+1/2,x+3/4,z+1/4'
'y+1/4,x+3/4,-z+1/2'
'-z+1/2,-y,-x+1/2'
'z+1/4,y+3/4,-x+1/2'
'z+1/4,-y,x+1/4'
'-z+1/2,y+3/4,x+1/4'
'-x+1/2,-z,-y+1/2'
'-x+1/2,z+3/4,y+1/4'
'x+1/4,z+3/4,-y+1/2'
'x+1/4,-z,y+1/4'
'x,y+1/2,z+1/2'
'x,-y+3/4,-z+3/4'
'-x+1/4,y+1/2,-z+3/4'

'-x+1/4,-y+3/4,z+1/2'
'y,z+1/2,x+1/2'
'-y+1/4,-z+3/4,x+1/2'
'y,-z+3/4,-x+3/4'
'-y+1/4,z+1/2,-x+3/4'
'z,x+1/2,y+1/2'
'-z+1/4,x+1/2,-y+3/4'
'-z+1/4,-x+3/4,y+1/2'
'z,-x+3/4,-y+3/4'
'y,x+1/2,z+1/2'
'-y+1/4,x+1/2,-z+3/4'
'y,-x+3/4,-z+3/4'
'-y+1/4,-x+3/4,z+1/2'
'z,y+1/2,x+1/2'
'-z+1/4,-y+3/4,x+1/2'
'-z+1/4,y+1/2,-x+3/4'
'z,-y+3/4,-x+3/4'
'x,z+1/2,y+1/2'
'x,-z+3/4,-y+3/4'
'-x+1/4,-z+3/4,y+1/2'
'-x+1/4,z+1/2,-y+3/4'
'-x,-y+1/2,-z+1/2'
'-x,y+1/4,z+1/4'
'x+3/4,-y+1/2,z+1/4'
'x+3/4,y+1/4,-z+1/2'
'-y,-z+1/2,-x+1/2'
'y+3/4,z+1/4,-x+1/2'
'-y,z+1/4,x+1/4'
'y+3/4,-z+1/2,x+1/4'
'-z,-x+1/2,-y+1/2'
'z+3/4,-x+1/2,y+1/4'
'z+3/4,x+1/4,-y+1/2'
'-z,x+1/4,y+1/4'
'-y,-x+1/2,-z+1/2'
'y+3/4,-x+1/2,z+1/4'
'-y,x+1/4,z+1/4'
'y+3/4,x+1/4,-z+1/2'
'-z,-y+1/2,-x+1/2'
'z+3/4,y+1/4,-x+1/2'
'z+3/4,-y+1/2,x+1/4'
'-z,y+1/4,x+1/4'
'-x,-z+1/2,-y+1/2'
'-x,z+1/4,y+1/4'
'x+3/4,z+1/4,-y+1/2'
'x+3/4,-z+1/2,y+1/4'

_cell_length_a	8.39960(10)
_cell_length_b	8.39960(10)
_cell_length_c	8.39960(10)
_cell_angle_alpha	90.00000
_cell_angle_beta	90.00000
_cell_angle_gamma	90.00000
_cell_volume	592.618(13)
_cell_formula_units_Z	?

```
_cell_measurement_temperature    ?
_cell_special_details
; ?
;
# The next three fields give the specimen dimensions in mm. The equatorial
# plane contains the incident and diffracted beam.
```

```
_pd_spec_size_axial      ?    # perpendicular to
                          # equatorial plane
_pd_spec_size_equat     ?    # parallel to
                          # scattering vector
                          # in transmission
_pd_spec_size_thick     ?    # parallel to
                          # scattering vector
                          # in reflection
```

```
# The next five fields are character fields that describe the specimen.
```

```
_pd_spec_mounting      # This field should be
                        # used to give details of the
                        # container.
; ?
;
_pd_spec_mount_mode    ?    # options are 'reflection'
                        # or 'transmission'
_pd_spec_shape        ?    # options are 'cylinder'
                        # 'flat_sheet' or 'irregular'
_pd_char_particle_morphology ?
_pd_char_colour       ?    # use ICDD colour descriptions
```

```
# The following three fields describe the preparation of the specimen.
# The cooling rate is in K/min. The pressure at which the sample was
# prepared is in kPa. The temperature of preparation is in K.
```

```
_pd_prep_cool_rate     ?
_pd_prep_pressure      ?
_pd_prep_temperature   ?
```

```
# The next four fields are normally only needed for transmission experiments.
```

```
_exptl_absorpt_coefficient_mu ?
_exptl_absorpt_correction_type ?
_exptl_absorpt_process_details ?
_exptl_absorpt_correction_T_min ?
_exptl_absorpt_correction_T_max ?
```

```
#=====
```

7. EXPERIMENTAL DATA

```
_exptl_special_details
; ?
;
```

The following item is used to identify the equipment used to record
the powder pattern when the diffractogram was measured at a laboratory
other than the authors' home institution, e.g. when neutron or synchrotron
radiation is used.

```
_pd_instr_location  
; ?  
;  
_pd_calibration_special_details      # description of the method used  
                                     # to calibrate the instrument  
; ?  
;
```

```
_diffrn_ambient_temperature      ?  
_diffrn_source                    'nuclear reactor'  
_diffrn_radiation_type            'Constant Wavelength Neutron Diffraction'  
_diffrn_radiation_wavelength      2.53600  
_diffrn_source_type               ? # Put here the diffractometer and site
```

```
_diffrn_radiation_monochromator   ?  
_diffrn_measurement_device_type   ?  
_diffrn_measurement_method        ?  
_diffrn_detector_area_resol_mean  ? # Not in version 2.0.1  
_diffrn_detector                  ?  
_diffrn_detector_type              ? # make or model of detector  
_pd_meas_scan_method               ? # options are 'step', 'cont',  
                                     # 'tof', 'fixed' or  
                                     # 'disp' (= dispersive)  
_pd_meas_special_details  
; ?  
;
```

The following four items give details of the measured (not processed)
powder pattern. Angles are in degrees.

```
_pd_meas_number_of_points         0  
_pd_meas_2theta_range_min         0.95000  
_pd_meas_2theta_range_max         151.89999  
_pd_meas_2theta_range_inc         0.050017
```

#=====

8. REFINEMENT DATA

```
_refine_special_details  
; ?  
;
```

Use the next field to give any special details about the fitting of the
powder pattern.

```
_pd_proc_ls_special_details  
; ?  
;
```

The next three items are given as text.

_pd_proc_ls_profile_function ?
_pd_proc_ls_background_function ?
_pd_proc_ls_pref_orient_corr
; ?
;

The following profile R-factors are NOT CORRECTED for background
The sum is extended to all non-excluded points.
These are the current CIF standard

_pd_proc_ls_prof_R_factor 3.8219
_pd_proc_ls_prof_wR_factor 5.1178
_pd_proc_ls_prof_wR_expected 2.2903

The following profile R-factors are CORRECTED for background
The sum is extended to all non-excluded points.
These items are not in the current CIF standard, but are defined above

_pd_proc_ls_prof_cR_factor 7.4371
_pd_proc_ls_prof_cwR_factor 7.4087
_pd_proc_ls_prof_cwR_expected 3.3155

The following items are not in the CIF standard, but are defined above

_pd_proc_ls_prof_chi2 4.9932
_pd_proc_ls_prof_echi2 5.3327

Items related to LS refinement

_refine_ls_R_I_factor 1.3413
_refine_ls_number_reflns 14
_refine_ls_number_parameters 94
_refine_ls_number_restraints 0

The following four items apply to angular dispersive measurements.
2theta minimum, maximum and increment (in degrees) are for the
intensities used in the refinement.

_pd_proc_2theta_range_min 0.8918
_pd_proc_2theta_range_max 151.8418
_pd_proc_2theta_range_inc 0.050017
_pd_proc_wavelength 2.536000

_pd_block_diffraction_id ? # The id used for the block containing
the powder pattern profile (section 11)

Give appropriate details in the next two text fields.

_pd_proc_info_excluded_regions ?
_pd_proc_info_data_reduction ?

The following items are used to identify the programs used.

_computing_data_collection ?
_computing_structure_solution ?
_computing_structure_refinement FULLPROF
_computing_molecular_graphics ?
_computing_publication_material ?

#=====

9. ATOMIC COORDINATES AND DISPLACEMENT PARAMETERS

loop_
_atom_site_label
_atom_site_fract_x
_atom_site_fract_y
_atom_site_fract_z
_atom_site_U_iso_or_equiv
_atom_site_occupancy
_atom_site_adp_type # Not in version 2.0.1
_atom_site_type_symbol
Ge1 0.12500 0.12500 0.12500 0.00458 1.00000 Uani Ge
Fe1 0.50000 0.50000 0.50000 0.00608 0.99988 Uani Fe
O1 0.24660 0.24660 0.24660 0.00618 0.99994 Uani O

loop_
_atom_site_aniso_label
_atom_site_aniso_U_11
_atom_site_aniso_U_22
_atom_site_aniso_U_33
_atom_site_aniso_U_12
_atom_site_aniso_U_13
_atom_site_aniso_U_23
_atom_site_aniso_type_symbol
Ge1 0.00458 0.00458 0.00458 0.00000 0.00000 0.00000 Ge
Fe1 0.00608 0.00608 0.00608 0.00039 0.00039 0.00039 Fe
O1 0.00618 0.00618 0.00618 -0.00150 -0.00150 -0.00150 O

Note: if the displacement parameters were refined anisotropically
the U matrices should be given as for single-crystal studies.

#=====

10. DISTANCES AND ANGLES / MOLECULAR GEOMETRY

_geom_special_details ?

loop_
_geom_bond_atom_site_label_1
_geom_bond_atom_site_label_2
_geom_bond_site_symmetry_1
_geom_bond_site_symmetry_2
_geom_bond_distance
_geom_bond_publ_flag

? ? ? ? ? ?

loop_
_geom_contact_atom_site_label_1
_geom_contact_atom_site_label_2
_geom_contact_distance
_geom_contact_site_symmetry_1
_geom_contact_site_symmetry_2
_geom_contact_publ_flag
? ? ? ? ? ?

loop_
_geom_angle_atom_site_label_1
_geom_angle_atom_site_label_2
_geom_angle_atom_site_label_3
_geom_angle_site_symmetry_1
_geom_angle_site_symmetry_2
_geom_angle_site_symmetry_3
_geom_angle
_geom_angle_publ_flag
? ? ? ? ? ? ? ?

loop_
_geom_torsion_atom_site_label_1
_geom_torsion_atom_site_label_2
_geom_torsion_atom_site_label_3
_geom_torsion_atom_site_label_4
_geom_torsion_site_symmetry_1
_geom_torsion_site_symmetry_2
_geom_torsion_site_symmetry_3
_geom_torsion_site_symmetry_4
_geom_torsion
_geom_torsion_publ_flag
? ? ? ? ? ? ? ? ? ?

loop_
_geom_hbond_atom_site_label_D
_geom_hbond_atom_site_label_H
_geom_hbond_atom_site_label_A
_geom_hbond_site_symmetry_D
_geom_hbond_site_symmetry_H
_geom_hbond_site_symmetry_A
_geom_hbond_distance_DH
_geom_hbond_distance_HA
_geom_hbond_distance_DA
_geom_hbond_angle_DHA
_geom_hbond_publ_flag
? ? ? ? ? ? ? ? ? ? ?

#=====

#=====

Additional structures (last six sections and associated data_? identifiers)
may be added at this point.

#=====

The following lines are used to test the character set of files sent by
network email or other means. They are not part of the CIF data set.
abcdefghijklmnopqrstuvwxyzABCDEFGHIJKLMNOPQRSTUVWXYZ0123456789
!@#\$%^&*()_+{ }:"~<>?|\=-[];'`.,/