CrysFML: A crystallographic library in modern Fortran

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Content of the talk

→ Scientific Computing: Why Fortran?

→ Crystallographic computing: CrysFML





Programming paradigms for scientific applications (I)

→ Procedural, imperative structured programming (PP)

Pascal, C, Fortran 77, ...

→ Module-Oriented Programming (MOP)

Fortran 95, ADA95, Modula-2, ... Fortran 2003

→ Object oriented programming (OOP)

C++, Java, Smalltalk, Eiffel, ADA95, ...

Fortran 2003





Programming paradigms for scientific applications (II). Why Fortran?

Some reasons for developing in modern Fortran

- → Simplicity and clarity of the syntax and the new facilities for global array manipulation. This is important for the common scientist that may write programs occasionally. This makes programming in **Fortran** more natural and problem solving oriented.
- → Availability of many OOP features in modern **Fortran**: user-defined types, encapsulation, overload of procedures and functions. The lacking features (e.g. direct inheritance and class methods) are of less importance for scientific computing than those already available (all of them are available in **Fortran 2003**).





Programming paradigms for scientific applications (III). Why Fortran?

Some reasons for developing in modern Fortran

- → The powerful implicit interface provided by encapsulating all functions and subroutines in *modules*, allowing to catch many errors at compile time, if one uses the *intent* attribute for procedure arguments. We may consider that Module Oriented Programming as an alternative/complement to OOP.
- → Efficiency of the generated executable codes compared to C/C++ programs of similar complexity.
- → Compatibility with legacy code and availability of a huge amount of free mathematical subroutines and functions. Re-usability of procedures written in **Fortran 77** was already a reality.





Programming paradigms for scientific applications (IV). Why Fortran?

Some reasons for developing in modern Fortran

- → The new standard (published in November 2004):
 Fortran 2003 contains all necessary features to perform pure OOP
- → John Reid, WG5 Convener: *The new features of Fortran 2003*, PDF document available directly from the Internet: ftp://ftp.nag.co.uk/sc22wg5/N1551-N1600/N1579.pdf
- → To our knowledge **Fortran 2003** exist partially in NagF95, G95, in the new Lahey compiler for .NET, ...





Existing Crystallographic Libraries (CCSL)

CCSL (Crystallographic Cambridge Subroutines Library)

- J. Brown, J.C. Matthewmann (W.I.F. David for powder diffraction)
- ⇒ The most complete set of procedures for crystallographic calculations. Well documented.
- ⇒ Written in Fortran 77 and with single crystal work in mind. Profuse use of commons. Difficult to adapt to modern programming techniques.





Existing Crystallographic Libraries (cctbx, Clipper)

Computational Crystallography Toolbox (cctbx)

R.W. Grosse-Kunstleve, P.D. Adams....

Clipper

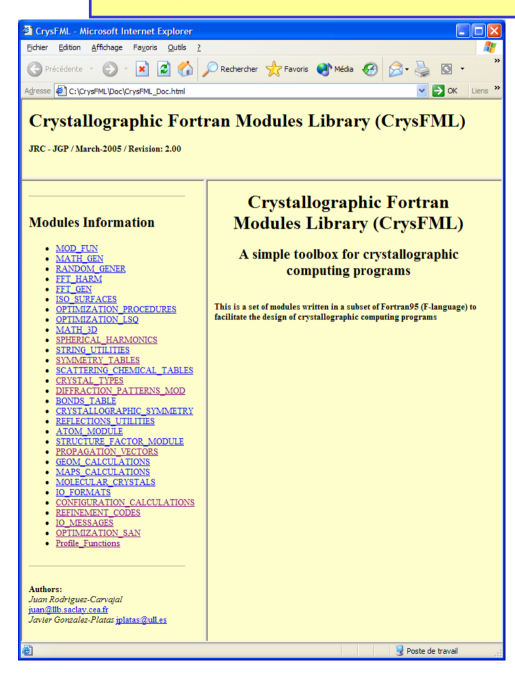
Kevin Cowtan

⇒ Written in C++ and handled using Python scripts.





CrysFML: a collection of F-modules for crystallography



"Crystallographic Fortran
Modules Library (CrysFML). A
simple toolbox for crystallographic
computing programs"
Commission on Crystallographic
Computing, IUCr
Newsletter No.1, pp 50-58,

January 2003.

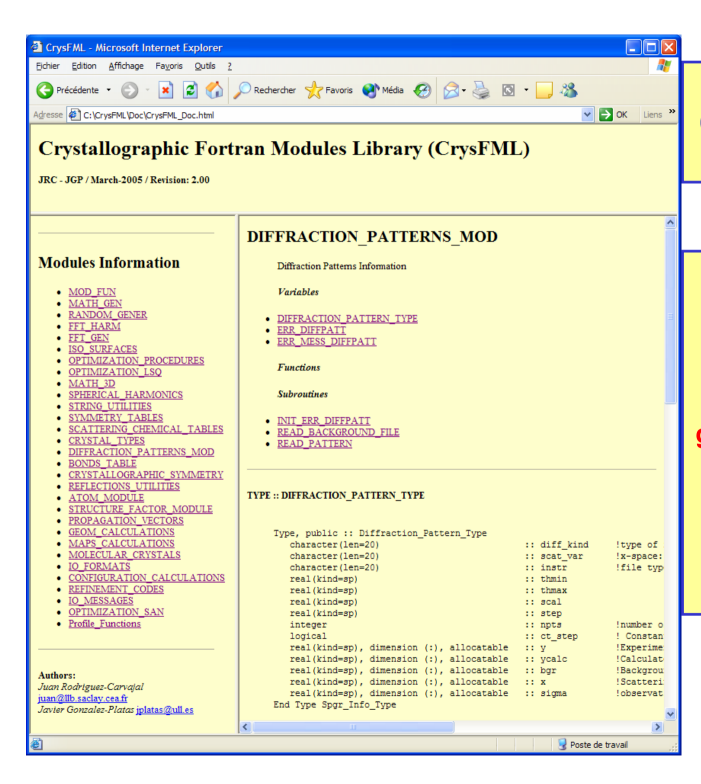
There are many other modules that are not ready for distribution: Magnetism,

Cost_functions

Instrument descriptions (Four circles + large PSD)

Refinement codes for molecular crystals





CrysFML info

In some cases the information about a particular procedure doesn't appear, then goto the source code!

The reason: I didn't obey my own rules for documentation!



Developers of CrysFML/WinPLOTR/FullProf

Juan Rodríguez-Carvajal (LLB, France)

CrysFML, FullProf, Baslreps, Simbo, Enermag, Polar3D,...

Javier González-Platas (ULL, Tenerife, Spain)

CrysFML, GUIs, GFourier, EdPCR

- **⇒** Contributors:
- ⇒ Thierry Roisnel (LCSIM, Rennes, France)

WinPLOTR

⇒ Carlos Frontera (ICMAB, Barcelona, Spain)

Polarized neutrons, Flipping ratio data handling

⇒ Marc Janoschek (PSI, Villigen, Switzerland)

Polarized neutrons, 3D-Polarimetry

⇒ Laurent Chapon & Aziz Daoud-Aladine (ISIS, U.K.)

T.O.F. powder diffraction, WCrysFGL, Fp_Studio Incommensurate crystal structures









Developers of ...

We are not professional programmers!

- ⇒ Juan Rodríguez-Carvajal (LLB, CEA-CNRS, France) Structural, electronic and magnetic properties of oxides and intermetallics. Modeling of magnetic structures
- ⇒ Javier González-Platas (ULL, Tenerife, Spain) Crystal structure determination of organic natural compounds. Teaching in Physics.
- ⇒ Thierry Roisnel (LCSIM, Rennes, France) Crystal structure determination of cluster compounds Single Crystal X-ray diffraction service (U. of Rennes)
- ⇒ Carlos Frontera (ICMAB, Barcelona, Spain) Magnetic properties of oxides
- ⇒ Aziz Daoud-Aladine (ISIS, UK)
 Charge, spin and orbital ordering in manganites (Co-resp. SXD)
- ⇒ Laurent Chapon (ISIS, UK)

 Thermo-electrics, multi-ferroics, ... (Co-resp. GEM)
- ⇒ Marc Janoschek (PSI, Villigen, Switzerland)
 Polarized neutrons instrumentation, Mu-PAD









Scope of CrysFML

We have developed a set of Fortran 95 modules, Crystallographic Fortran Modules Library (CrysFML), that may be used (in the Fortran 95 sense) in crystallographic and diffraction computing programs.

- ⇒ Modern array syntax and new features of Fortran 95 are used through the modules. In fact the whole library is written in **F-language**, a strict subset of Fortran 95 for which free compilers are available.
- ⇒ We take advantage of all object oriented programming (OOP) techniques already available in Fortran: user-defined types, encapsulation, overload (polymorphism) of procedures and functions. The lacking features (e.g. inheritance and class methods) will be easily implemented as soon as Fortran 2003 compilers become available.
- ⇒ Main programs using the adequate modules may perform more or less complicated calculations with only few lines of code.





F-language (strict subset of Fortran 95)

All free F-compilers can be downloaded from the site:

ftp://ftp.swcp.com/~walt/pub/F

See also:

http://www.fortran.com/fortran/Imagine1





Free Fortran 95 compiler G95: strong development

All implementations of the G95-compiler (based in gcc) can be downloaded from the G95 home page:

http://www.g95.org

Platforms: Linux, Windows, Mac OS, Solaris, OpenBSD, etc...





Present status of CrysFML

- ⇒ The present **CrysFML** contains general and specific Mathematical modules (FFTs, geometrical calculations, optimizers, matrix operations). Procedures for reading files of many different formats, string utilities for handling free format, generation and reading of CIF files.
- ⇒ Modules for generating space groups from their Hermann-Mauguin or Hall symbols. Generic space groups with non-conventional lattice centring vectors can also be built using user-defined generators.
- ⇒ Reflection handling modules, including propagation vectors, may be used for generating reflections in selected regions of reciprocal space and for calculating structure factors.
- ⇒ The documentation is written within the source code using special comment symbols. A document, in HTML format, containing the description of all modules and procedures can be generated using a Fortran program (get_doc).







Present status of CrysFML

- ⇒ At present there is no formal way of distributing **CrysFML**, I can send copies (of the most stable modules) by e-mail to everyone wishing to use it.
- ⇒ There are parts of the library that are not completely developed so be patient and comprehensive.
- ⇒ The library is distributed with a set of working examples so that the user can mimic in order to create his (her) own programs.





Programs using CrysFML (I)

FullProf: Crystal and magnetic structure refinement, powder/single crystals, polarised neutrons, constant wavelength, TOF, energy dispersive, multiple patterns.

FOURIER, **GFOURIER** and **EdPCR**. These programs work on Windows and Linux and are already distributed from the LLB Web site.

BasIREPS: Program for calculating basis functions of irreducible representations of space groups. This program is useful for determining magnetic structures and phonon symmetry analysis.

SIMBO: Program for the analysis of the magnetic topology of an arbitrary crystal structure. Generates a formal description of the Fourier transform of the exchange interactions to be used by other programs.





Programs using CrysFML (II)

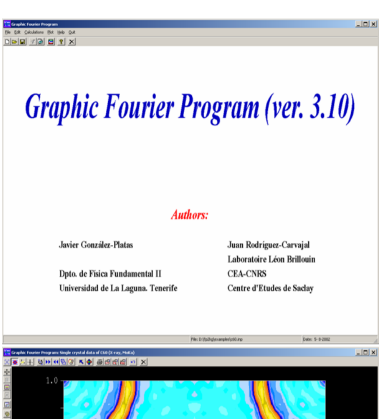
ENERMAG: Program to analyse the classical magnetic energy as a function of the exchange interactions and the point in the Brillouin Zone. This program can be used to generate theoretical magnetic phase diagrams in the J-space in order to get insight into the experimentally determined magnetic structures.

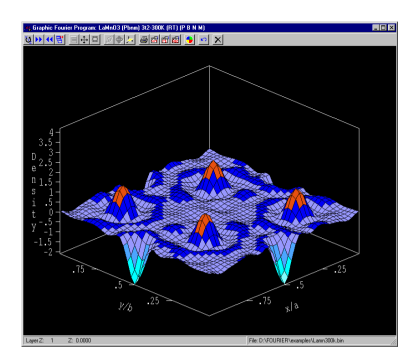
SIMILAR: Program to make conversion of settings for describing crystallographic structures. It determines automatically the splitting of Wyckoff positions on going from a space group to one of their subgroups. Calculate all the *translationengleiche* subgroups of a space group, co-set decompositions, etc.

DATARED: Program for data reduction of single crystal data. It handles twinning and incommensurate magnetic and crystal structures. Prepares files to be read by *FullProf* when using single crystals.



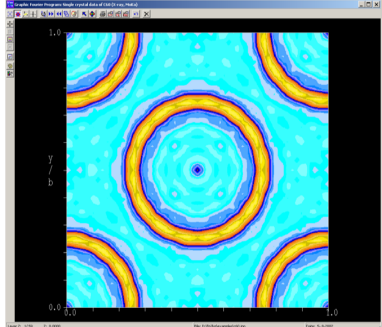






The programs Gfourier and Fourier are based in CrysFML

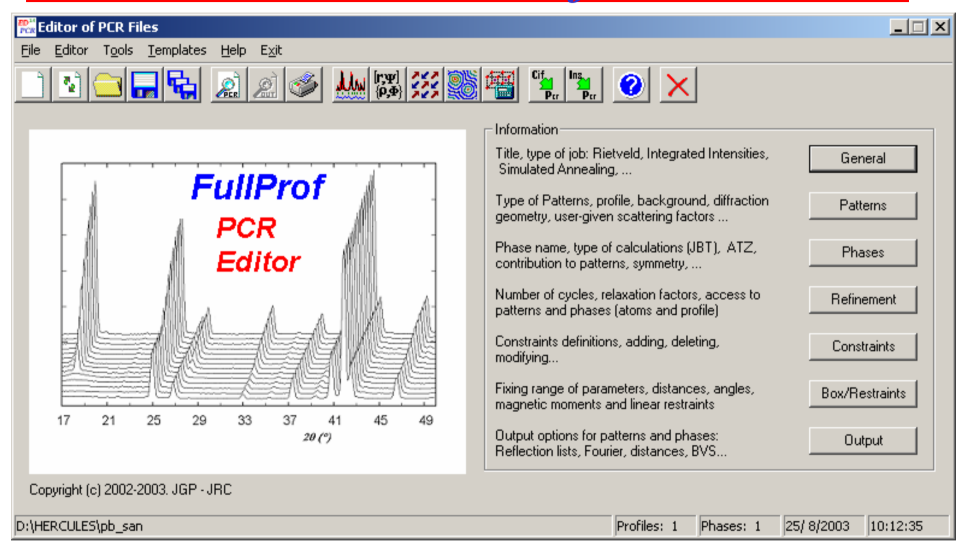
Graphic utilities: Winteracterhttp://www.winteracter.com







A GUI for FullProf: EdPCR



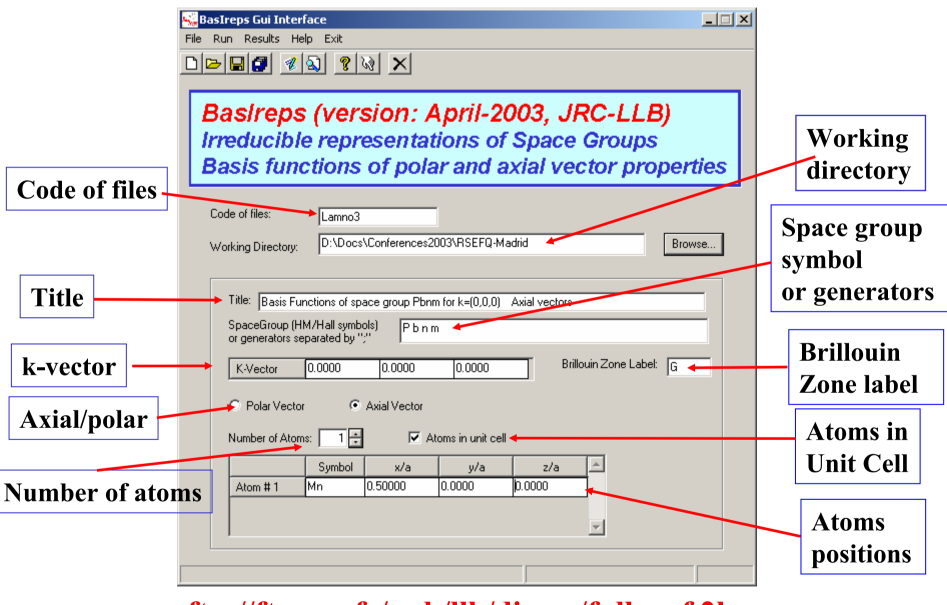
GUI using Winteracter: http://www.winteracter.com







GUI for BasIreps



Siena 2005:
Crystallographic Computing Computi

ftp://ftp.cea.fr/pub/llb/divers/fullprof.2k





Example of *BasIreps* output: *.bsr

PROPAGATION VECTOR GROUP INFORMATION _____ => The input propagation vector is: K=(0.5000 0.5000 0.5000) => K .. IS NOT .. equivalent to -K => The operators following the k-vectors constitute the co-set decomposition G[Gk] The list of equivalent k-vectors are also given on the right of operators. => The star of K is formed by the following 2 vectors: k 1 = (0.5000 0.5000 0.5000)Op: (1) x, y, zOp: (3) x,-y,-z-> (0.5000 -0.5000 -0.5000) -> (-0.5000 -0.5000 0.5000) Op: (4) -x+1/2, -y, z+1/2Op: (7) -x+1/2, y, -z+1/2-> (-0.5000 0.5000 -0.5000) Op: (10) y+3/4,-x+1/4,-z+3/4-> (-0.5000 0.5000 -0.5000) Op: (13) -y+3/4, -x+1/4, z+3/4-> (-0.5000 -0.5000 0.5000) -> (0.5000 -0.5000 -0.5000) Op: (14) -y+3/4,x+3/4,-z+1/4Op: (16) y+3/4,x+3/4,z+1/4-> (0.5000 0.5000 0.5000) Eqv. -K: k 2 = (0.5000 - 0.5000 0.5000)Op: (2) -y+1/4, x+3/4, z+1/4Op: (5) y+1/4,x+3/4,-z+1/4-> (0.5000 0.5000 -0.5000) Op: (6) y+1/4,-x+1/4,z+3/4-> (-0.5000 0.5000 0.5000) Op: $(8) -y+1/4, -x+1/4, -z+3/4 \rightarrow (-0.5000 -0.5000 -0.5000)$ Op: (9) -x, -y, -z-> (-0.5000 -0.5000 -0.5000) Op: (11) -x,y,z-> (-0.5000 0.5000 0.5000) Op: (12) x+1/2, y, -z+1/2-> (0.5000 0.5000 -0.5000) -> (0.5000 -0.5000 0.5000) Op: (15) x+1/2,-y,z+1/2=> G k has the following symmetry operators: 1 SYMM(1) = x,y,z2 SYMM(3) = x, -y, -z4) = -x+1/2, -y, z+1/23 SYMM(

 $\frac{4}{\text{mission on CrystalloS YMM}} \left(\frac{1}{\text{mission on CrystalloS YMM}} \right) \right) = -x+1/2, y, -z+1/2$



Example of *BasIreps* output: *.bsr

```
=> Number of elements of G k:
=> Number of irreducible representations of G_k:
=> Dimensions: 2 2
=> Symmetry elements of G k and ireps:
   Symmetry elements reduced to the standard form (positive translations < 1)
   The matrices of IRreps have been multiplied by the appropriate phase factor
-> SYMM K(2): -x+1/2, -y, z+1/2 : 2 (0, 0, z) -> h4
   Phase factor for correcting input data: 0.0000
   Matrix of IRrep(1):
                    0 - i
   Matrix of IRrep(2):
-> SYMM_K(8): y+3/4,x+3/4,z+1/4 : m(x,x,z) -> h37
   Phase factor for correcting input data: 1.5000
   Matrix of IRrep(1):
   Matrix of IRrep(2):
                       -i
```





Example of *BasIreps* output: *.bsr

=> Basis functions of Representation IRrep(1) of dimension 2 contained 3 times in GAMMA -x+1/2,-y,z-1/2y+3/4,-x+1/4,-z+3/4-y+1/4,x+1/4,-z+3/4SYMM x, y, z Atoms: Cu 1 Cu 4 Cu 3 1:Re (0) (0 -1 0) 0) (0) (Im (0) (-1 0) (0) (0) 2:Re (0) (0) (0) (0) 0) (-1 0) (0) (0) Im (3:Re (1) (0) (-1) (1) Im (0) (1) (0) (0) -1 0) (0) (0) (0) 4:Re (-1 Im (0)(-1 0) (0) (0) 1 0) (0) (0) (0 5:Re (0) 0) (0) (0) (Im (-1 0) 6:Re (1) (0) (0) (0) 0) (-1) (-1) (-1) Im (---- LINEAR COMBINATIONS of Basis Functions: coefficients u,v,w,p,q General expressions of the Fourier coefficients Sk(i) i=1,2,...nat SYMM x,y,z Atom: Cu 1 0.0000 0.0000 0.5000 Sk(1): (u-p,v+q,w+r)SYMM -x+1/2, -y, z-1/2Atom: Cu 2 0.5000 0.0000 0.0000 Sk(2): i.(-u-p,-v+q,w-r) SYMM y+3/4,-x+1/4,-z+3/4Atom: Cu 3 0.7500 0.2500 0.2500 Sk(3): (v,-u,-w)+i.(q,p,-r)SYMM -y+1/4,x+1/4,-z+3/40.2500 Atom: Cu 4 0.2500

2005 SK(4); (v), -u, w)+i.(-q, -p, -r)

Programming with CrysFML using it nearly as a black-box

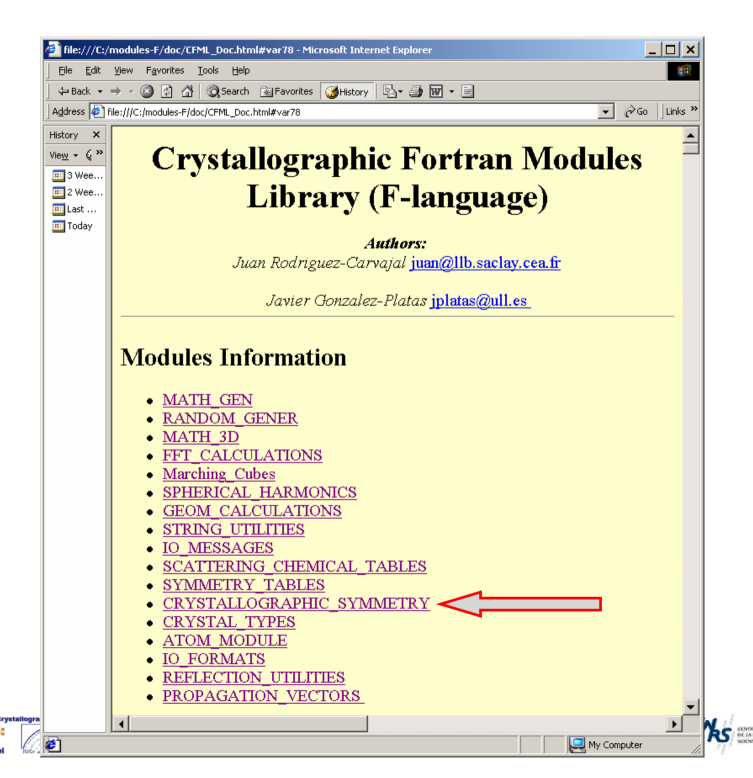
A Fortran 95/2003 compiler is needed (G95 is free!)

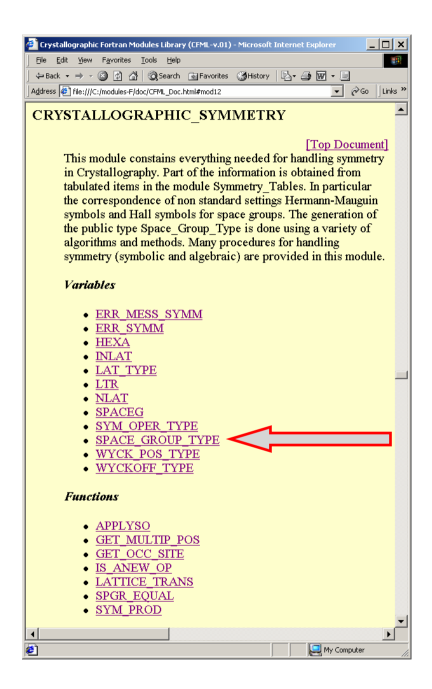
Learn the main structure types and procedures existing in the modules of the library by reading the documentation

Write a main program, using the modules of the library, for a particular purpose









• AXES_ROTATION • DECODMATMAG • GET_CRYSTAL_SYSTEM

• GET_CENTRING_VECTORS

• GET_LATTICE_TYPE

• GET_LAUE_NUM

• GET_LAUE_PG

GET_LAUE_STR

• GET_POINTGROUP_NUM

• GET POINTGROUP STR

• GET SO FROM FIX

• GET SO FROM GENER

GET_SO_FROM_HALL

• GET SO FROM HMS

• GET_SPG_FROM_GENER

• GET STABILIZER

• GET STRING RESOLV

• GET SYMEL

GET SYMKOV

GET_SYMSYMB

GET_SYMSYMB2

GET_WYCKOFF_FILE

• <u>INIT_ERR_SYMM</u>

• INVERSE_SYMM

LATSYM

• READ MSYMM

READ_XSYM

SEARCHOP

• <u>SET_SPACEGROUP</u>

SETTING_CHANGE

SYM_B_RELATIONS

SYMMETRY_SYMBOL

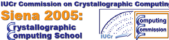
SYM_PROD_str

WRITE_SPACEG

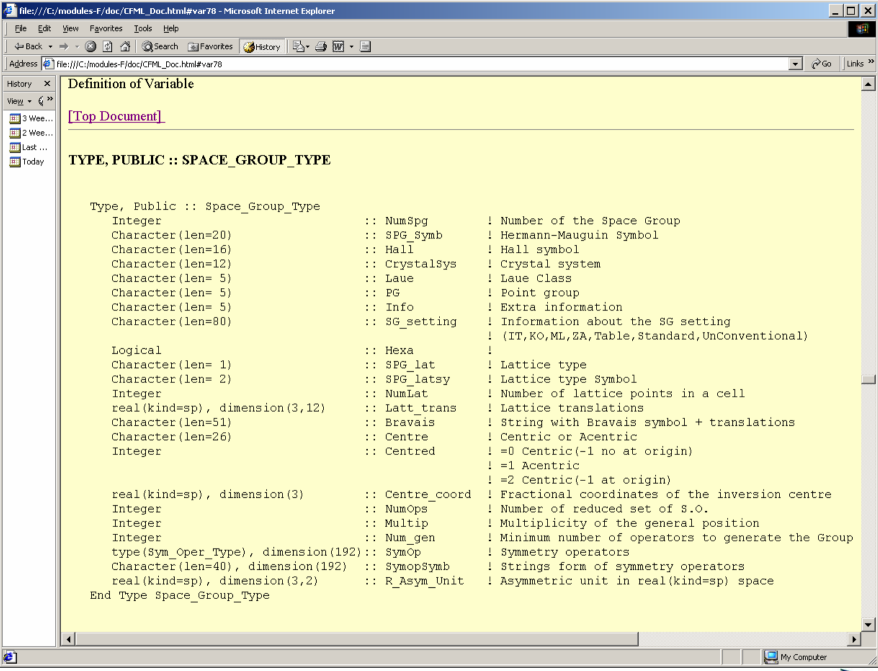
WRITE_SYM

WRITE_WYCKOFF

Example using the HTML automatically generated documentation









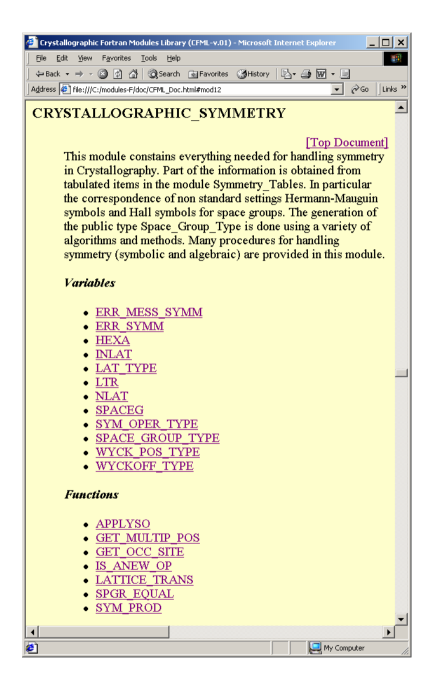


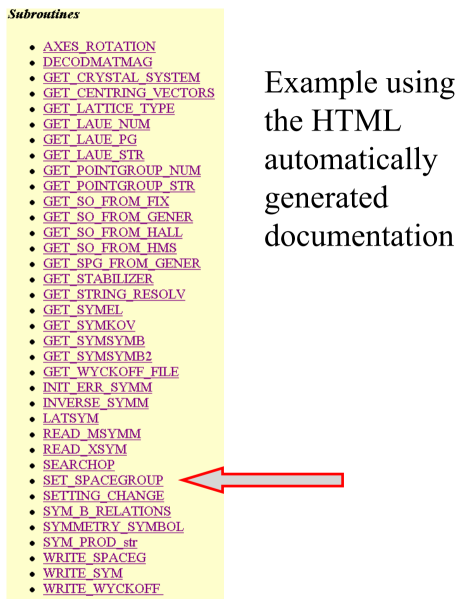
Space Group Type in CrysFML

```
Type :: Space Group Type
                                           :: NumSpg
                                                            ! Number of the Space Group
      Integer
      Character (len=20)
                                           :: SPG Symb
                                                            ! Hermann-Mauguin Symbol
      Character (len=16)
                                          :: Hall
                                                            ! Hall symbol
      Character (len=12)
                                          :: CrystalSys
                                                            ! Crvstal system
      Character(len= 5)
                                                            ! Laue Class
                                          :: Laue
      Character (len= 5)
                                          :: PG
                                                            ! Point group
      Character(len= 5)
                                          :: Info
                                                            ! Extra information
      Character (len=80)
                                          :: SG setting
                                                            ! Information about the SG setting
                                                            ! (IT, KO, ML, ZA, Table, Standard, UnConventional)
      Logical
                                          :: Hexa
      Character (len= 1)
                                          :: SPG lat
                                                            ! Lattice type
      Character (len= 2)
                                          :: SPG latsv
                                                            ! Lattice type Symbol
      Integer
                                          :: NumLat
                                                            ! Number of lattice points in a cell
     real(kind=sp), dimension(3,12)
                                                            ! Lattice translations
                                           :: Latt trans
      Character (len=51)
                                          :: Bravais
                                                            ! String with Bravais symbol + translations
      Character (len=26)
                                                            ! Centric or Acentric
                                          :: Centre
      Integer
                                           :: Centred
                                                            ! = 0 Centric(-1 no at origin)
                                                            ! =1 Acentric
                                                            ! =2 Centric(-1 at origin)
                                                           ! Fractional coordinates of the inversion centre
      real(kind=sp), dimension(3)
                                           :: Centre coord
                                                            ! Number of reduced set of S.O.
      Integer
                                          :: NumOps
                                           :: Multip
                                                            ! Multiplicity of the general position
      Integer
                                                            ! Minimum numb. of oper. to generate the group
      Integer
                                          :: Num gen
     type(Sym Oper Type), dimension(192):: SymOp
                                                            ! Symmetry operators
     Character (len=40), dimension (192)
                                                            ! Strings form of symmetry operators
                                          :: SymopSymb
     type(wyckoff type)
                                                            ! Wyckoff Information
                                           :: Wyckoff
                                                            ! Asymmetric unit in real space
      real(kind=sp), dimension(3,2)
                                          :: R Asym Unit
  End Type Space Group Type
```



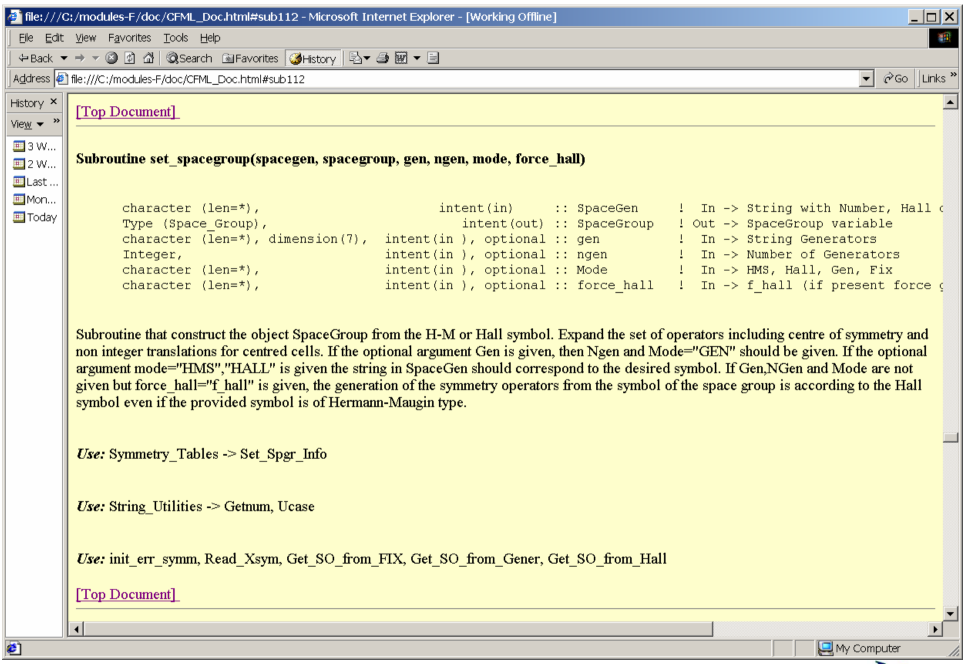
















The procedure Set_SpaceGroup

Header of the subroutine Set_Spacegroup. Only two arguments are needed in the most simple cases.

The string Spacegen may contain the Hermann-Mauguin (H-M) symbol, the Hall symbol or, simply, the number of the space group.

The object Spacegroup is provided by a call to the subroutine.





The procedure Set_SpaceGroup

One can make a call to the subroutine as follows:

! Declarations omitted

```
Ngen=3
Gen(1)="y,-x, z"
Gen(2)="-x,-y,-z"
Gen(3)="x+1/2, y+1/2, -z"
Call Set_Spacegroup(Spacegen,Spacegroup,Gen,Ngen,"GEN")
```

On output the object Spacegroup of type Space_Group_type is filled with all possible information obtained from the list of the given generators.





The procedure Write_SpaceGroup

```
Example of simple program using CrysFML
Program Get SPG info
  use crystallographic symmetry, only:
 space group type, set spacegroup, write spacegroup
  character(len=20)
                     :: spg symb
  type(space group type) :: SPG
  do
    write(unit=*,fmt="(a)",advance="no") &
    " => Please enter a space group (H-M/Hall/number): "
    read(unit=*,fmt="(a)") spg symb
    if(len trim(spg symb) == 0) exit
    call set spacegroup(spg symb,SPG)
    call write_spacegroup(SPG,full=.true.)
  end do
  stop
End Program Get SPG info
```





The procedure Write_SpaceGroup

The argument full in procedure Write_SpaceGroup means that all detailed information in asked to be output in the screen. One may change the instruction to write directly to an already opened file. For instance writing:

Call Write_SpaceGroup(SPG,iunit=3,full=.true.)

directs the output to the file connected with logical unit 3





Output of the small program: Get_SPG_info

```
DOS-shell - simple-a
                                                                                   _ | D | X |
       Information on Space Group:
=> Number of Space group: 15
=> Hermann-Mauguin Symbol: C 2/C
              Hall Symbol: -C 2yc
     Table Setting Choice: b1
             Setting Type: IT (Generated from Hermann-Mauguin symbol)
           Crystal System: Monoclinic
               Laue Class: 2/m
              Point Group: 2/m
          Bravais Lattice: C
           Lattice Sumbol: mC
=> Reduced Number of $.0.:
     General multiplicity:
           Centrosymmetry: Centric (-1 at origin)
-> Generators (exc. -1&L):
          Asymmetric unit: 0.000 <= x <= 0.500
                             0.000 <= y <= 0.500
                             0.000 <= z <= 0.500
=> Centring vectors: 1
=> Latt( 1): ( 1/2, 1/2, 0 )
=> List of all Symmetry Operators and Symmetry Symbols
=> SYMM( 1): x, y, z
                                                 Symbol: 1
=> SYMM( 2): -x,y,-z+1/2
                                                 Symbol: 2 0, y, 1/4
=> SYMM( 3): -x,-y,-z
                                                 Sumbol: -1 0.0.0
=> SYMM(
          4): x,-y,z+1/2
                                                 Symbol: c \times 0, z
=> SYMM(
          5): x+1/2, y+1/2, z
                                                 Symbol: t(1/2,1/2,0)
=> SYMM(
          6): -x+1/2, y+1/2, -z+1/2
                                                 Sumbol: 2 (0,1/2,0) 1/4,y,1/4
=> SYMM( 7): -x+1/2,-y+1/2,-z
                                                 Symbol: -1 1/4,1/4,0
\Rightarrow SYMM( 8): x+1/2,-q+1/2,z+1/2
                                                 Symbol: n (1/2,0,1/2) \times 1/4,z
=> Special Wyckoff Positions for C 2/C
   Multp
             Site
                         Representative Coordinates (centring translations excluded)
                             0, y, 1/4
                                                 0, -y, 3/4
                            1/4.1/4.1/2
                                                 3/4.1/4.0
                            1/4.1/4.0
                                                 3/4.1/4.1/2
                                                 0,1/2,1/2
                             0.1/2.0
                             0,0,0
                                                 0,0,1/2
=> Please enter a space group (H-M/Hall/number):
```







Another small program: test_subgroup

```
Program test subgroups
 use crystallographic symmetry, only: get T SubGroups, space group type, &
            set spacegroup, write spaceg, Lattice trans, similar transf SG
 use math 3d, only: determ a
 character(len=20)
                      :: spq symb
 type(space group type) :: SpG, SpGn
 type(space group type), dimension(48) :: Subgroup
 real, dimension (3,3) :: trans
 real, dimension (3) :: orig
 integer :: nsg, i, j, ng, 1
 real
         :: det
   read(unit=*,fmt="(a)") spq symb
   if(len trim(spq symb) == 0) exit
   call set spacegroup(spg symb,SPG)
                                        !Constructing the space group SPG
   write(unit=*,fmt="(a)",advance="no") " => Please enter a transformation matrix: "
   read(unit=*,fmt=*) (trans(i,:),i=1,3)
   det=determ a(trans)
   det=abs(det)
   write(unit=*,fmt="(a)",advance="no") " => Please enter the new origin: "
   read(unit=*,fmt=*) oriq
                                            !Construct the subgroup of SPG that is compatible
   call similar transf SG(trans, orig, SpG, SpGn)
                                            !with the transformation matrix and change of origin
   call write spaceq(SPGn, full=.true.)
                                            !give above
  !Determine all subgroups of the new space group
   call qet T SubGroups(SPGn, SubGroup, nsq)
   write(unit=*,fmt="(/,a,/)") " => LIST of Translationengleische Subgroups: "
   do i=1, nsq
     j=SPGn%Multip/SubGroup(i)%multip
     ng=SubGroup(i)%numops
     write(unit=*,fmt="(4a,i2,30a)") " => ", SubGroup(i)%Spq Symb, SubGroup(i)%hall,&
       "Index: [",j,"] -> { ", (trim(SubGroup(i)%SymopSymb(1))//" : ",l=1,nq-1),&
       end do
 end do
 stop
```

End Program test subgroups

Output of the small program: test_subgroup

```
Information on Space Group:
   Number of Space group: 176
=> Hermann-Mauguin Symbol: P 63/M
             Hall Symbol: -P 6c
=>
    Table Setting Choice:
=>
            Setting Type: a'=a, b'=b, c'=c -> Origin: (0,0,0)
=>
=> LIST of Translationengleiche Subgroups:
                                      Index: [2] -> \{x,y,z:...\},
=> P 63
                      P 6c
                                                                      Acentric
                                      Index: [ 1] -> { x,y,z : ...}, Centric
=> P 63/M
                      -P 6c
                                      Index: [ 4] -> { x,y,z : ...}, Acentric
=> P 3
                     P 3
                                      Index: [ 2] -> { x,y,z : ...}, Centric
=> P - 3
                    -P 3
=> P 1 1 21
                                      Index: [ 6] -> { x,y,z : ...}, Acentric
                     P 2c
                                      Index: [ 3] -> { x,y,z : ...}, Centric
=> P 1 1 21/M
                      -P 2c
                                      Index: [6] -> \{x,y,z\},
                                                                      Centric
=> P -1
                      -P 1
=> unknown
                                      Index: [ 2] -> { x,y,z : ...}, Acentric
                      P -6c
                                      Index: [ 6]
                                                   -> { x,y,z : ...}, Acentric
=> unknown
                       P-2c
=> Please enter a space group (H-M/Hall/number):
```





Another Example: Check_Group

```
Program Check Group
 use crystallographic symmetry, only: Space Group Type, set spacegroup
 use reflections utilities, only: Hkl Absent
 use Symmetry Tables, only: spgr info, Set Spgr Info
  ..... ! Read reflections, apply criterion of "goodness" for checking,
          ! set indices i1,i2 for search in space group tables ...
  ..... ! omitted for simplicity
 call Set Spgr Info()
 m=0
  do group: do i=i1,i2
   hms=adjust1(spgr info(i)%HM)
    hall=spgr info(i)%hall
   if( hms(1:1) /= "P" .and. .not. check cent ) cycle do group ! Skip centred groups
   call set spacegroup(hall,Spacegroup,Force Hall="y")
    do j=1,nhkl
       if(good(j) == 0) cycle !Skip reflections that are not good (overlap) for checking
       absent=Hkl Absent(hkl(:,j), Spacegroup)
       if (absent .and. intensity(j) > threshold) cycle do group !Group not allowed
    end do
   ! Passing here means that all reflections are allowed in the group -> Possible group!
   num group(m)=i
 end do do group
 write(unit=*,fmt=*) " => LIST OF POSSIBLE SPACE GROUPS, a total of ",m," groups are possible"
 write(unit=*,fmt=*) "
 write(unit=*,fmt=*) "
                                                                     Hall Symbol"
                                            Hermann-Mauguin Symbol
 write(unit=*,fmt=*) "
 do i=1,m
    j=num group(i)
   hms=adjust1(spgr info(j)%HM)
   hall=spgr info(j)%hall
   numg=spgr info(j)%N
   write(unit=*,fmt="(i10,4a)") numg,"
                                                           ",hms,"
                                                                            ",hall
  end do
```









Check_Group output (1)

PROGRAM CHECK_GROUP: attempt to select the possible space groups from an experimental Powder Diffraction Pattern

Author: J.Rodriguez-Carvajal (version 0.01, based on CrysFML)

Conditions:

Input hkl-file : testgal.hkl
Crystal System : Tetragonal

Check centred cells?: Y

Maximum angle : 20.0000 Number of FWHMs : 2.0000 Threshold in % : 0.1000

=> List of read reflections:

h	k	1	Intensity	Sigma	2theta	FWHM Good?
1	0	1	0.0000	0.0000	3.2518	0.0093 1
1	1	0	3.4230	0.4030	3.6146	0.0113 1
0	0	2	0.5280	0.2050	4.0212	0.0091 1
1	1	1	1.8570	0.3130	4.1363	0.0111 1
	•					
2	2	2	3562.2319	38.4840	8.2781	0.0138 1
2	1	3	5.4550	0.3910	8.3152	0.0118 0
3	1	1	23.2680	0.1620	8.3347	0.0124 0
c Computing	0	4	0.0000	0.0000	8.4448	0.0102×11

Check_Group output (2)

=> Number of good reflections : 94

Maximum intensity : 3562.2319
Minimum (for observed) : 3.5622

Number of Space Group tested: 85

=> LIST OF POSSIBLE SPACE GROUPS, a total of 24 groups are possible

Number(IT)	Hermann-Mauguin Symbol	Hall	Symbol
75	P 4	P	4
76	P 41	P	4w
77	P 42	P	4c
78	P 43	P	4cw
81	P -4	P	-4
83	P 4/M	-P	4
84	P 42/M	-P	4c
89	P 4 2 2	P	4 2
90	P 4 21 2	P	4ab 2ab
91	P 41 2 2	P	4w 2c
92	P 41 21 2	P	4abw 2nw
113	P -4 21 M	P	-4 2ab
114	P -4 21 C	P	-4 2n
115	P -4 M 2	P	-4 -2





```
Program Calc structure factors
use crystallographic symmetry, only: space group type, Write SpaceGroup
use Atom Module,
                               only: Atoms List Type, Write Atoms List
                               only: Crystal Cell Type, Write Crystal Cell
use crystal types,
use Reflections Utilities,
                               only: Reflection Type, Hkl Uni, get maxnumref
use IO Formats,
                               only: Readn set Xtal Structure, err mess form, err form
use Structure Factor Module, only: Structure Factors, Write Structure Factors
type (space group type) :: SpG
type (Atoms list Type)
                          :: A
type (Crystal Cell Type) :: Cell
type (Reflection Type),allocatable, dimension(:) :: hkl
                          :: filcod !Name of the input file
character(len=256)
real
                          :: stlmax !Maximum Sin(Theta)/Lambda
integer
                          :: MaxNumRef, Num, lun=1
   do
    write(unit=*,fmt="(a)") " => Code of the file xx.cif (give xx):
    read(unit=*,fmt="(a)") filcod
    if(len trim(filcod) == 0) exit
    write(unit=*,fmt="(a)") " => Maximum sinTheta/Lambda: "
    read(unit=*,fmt=*) stlmax
    open(unit=lun,file=trim(filcod)//".sfa", status="replace",action="write")
    call Readn set Xtal Structure(trim(filcod)//".cif",Cell,SpG,A,Mode="CIF")
    If(err form) then
      write(unit=*,fmt="(a)") trim(err mess form)
       exit
     else
      call Write Crystal Cell(Cell,lun)
      call Write SpaceGroup(SpG,lun)
      call Write Atoms List(A,lun=lun)
      MaxNumRef = get maxnumref(stlmax,Cell%CellVol,mult=SpG%Multip)
       if(allocated(hkl)) deallocate(hkl); allocate (hkl(MaxNumRef))
      call Hkl Uni(Cell,Spg,.true.,0.0,stlmax,"s",Num,hkl)
      call Structure_Factors(A,SpG,Num,hkl,mode="NUC")
      call Write Structure Factors(lun, Num, hkl, mode="NUC")
    end if
    close(unit=lun)
    end do
End Program Calc_structure factors
```

```
call Readn_set_Xtal_Structure(trim(filcod)//".cif",&
                              Cell,SpG,A,Mode="CIF")
```

Reads a CIF file and sets up the objects:

Cell: contains everything related to metrics

SpG: contains everything related to symmetry

A: contains everything concerned with atoms in the asymmetric unit

End Program Calc_structure_factors





```
Program Calc structure factors
use crystallographic symmetry, only: space group type, Write SpaceGroup
use Atom Module,
                              only: Atoms List Type, Write Atoms List
                               only: Crystal Cell Type, Write Crystal Cell
use crystal types,
use Reflections Utilities,
                               only: Reflection Type, Hkl Uni, get maxnumref
use IO Formats,
                               only: Readn set Xtal Structure, err mess form, err form
use Structure Factor Module, only: Structure Factors, Write Structure Factors
 type (space group type) :: SpG
type (Atoms list Type)
                          :: A
type (Crystal Cell Type) :: Cell
type (Reflection Type),allocatable, dimension(:) :: hkl
                          :: filcod !Name of the input file
 character(len=256)
real
                          :: stlmax !Maximum Sin(Theta)/Lambda
 integer
                          :: MaxNumRef, Num, lun=1
    do
     write(unit=*,fmt="(a)") " => Code of the file xx.cif (give xx): "
     read(unit=*,fmt="(a)") filcod
     if(len trim(filcod) == 0) exit
    write(unit=*,fmt="(a)") " => Maximum sinTheta/Lambda: "
     read(unit=*,fmt=*) stlmax
     open(unit=lun,file=trim(filcod)//".sfa", status="replace",action="write")
     call Readn set Xtal Structure(trim(filcod)//".cif",Cell,SpG,A,Mode="CIF")
     If(err form) then
      write(unit=*,fmt="(a)") trim(err mess form)
      exit
     else
       call Write Crystal Cell(Cell,lun)
       call Write SpaceGroup(SpG,lun)
      call Write Atoms List(A,lun=lun)
      MaxNumRef = get maxnumref(stlmax,Cell%CellVol,mult=SpG%Multip)
      if(allocated(hkl)) deallocate(hkl); allocate (hkl(MaxNumRef))
      call Hkl_Uni(Cell,Spg,.true.,0.0,stlmax,"s",Num,hkl)
       call Structure Factors(A,SpG,Num,hkl,mode="NUC")
       call Write Structure Factors(lun,Num,hkl,mode="NUC")
     end if
     close(unit=lun)
    end do
End Program Calc_structure factors
```

```
Program Calc structure factors
use crystallographic symmetry, only: space group type, Write SpaceG
use Atom Module,
                              only: Atoms List Type, Write Atoms List
                               only: Crystal Cell Type, Write Crystal Cell
use crystal types,
use Reflections Utilities,
                               only: Reflection Type, Hkl Uni, get maxnumref
use IO Formats,
                               only: Readn set Xtal Structure, err mess form, err form
use Structure Factor Module, only: Structure Factors, Write Structure Factors
type (space group type) :: SpG
type (Atoms list Type) :: A
type (Crystal Cell Type) :: Cell
type (Reflection Type),allocatable, dimension(:) :: hkl
                          :: filcod !Name of the input file
 character(len=256)
real
                          :: stlmax !Maximum Sin(Theta)/Lambda
 integer
                          :: MaxNumRef, Num, lun=1
    do
     write(unit=*,fmt="(a)") " => Code of the file xx.cif (give xx): "
     read(unit=*,fmt="(a)") filcod
     if(len trim(filcod) == 0) exit
    write(unit=*,fmt="(a)") " => Maximum sinTheta/Lambda: "
     read(unit=*,fmt=*) stlmax
     open(unit=lun,file=trim(filcod)//".sfa", status="replace",action="write")
     call Readn set Xtal Structure(trim(filcod)//".cif",Cell,SpG,A,Mode="CIF")
If(err form) then
      write(unit=*,fmt="(a)") trim(err mess form)
       exit
     else
       call Write Crystal Cell(Cell,lun)
       call Write SpaceG(SpG,lun)
       call Write Atoms List(A,lun=lun)
      MaxNumRef = get maxnumref(stlmax,Cell%CellVol,mult=SpG%Multip)
       if(allocated(hkl)) deallocate(hkl); allocate (hkl(MaxNumRef))
      call Hkl_Uni(Cell,Spg,.true.,0.0,stlmax,"s",Num,hkl)
      call Structure Factors(A,SpG,Num,hkl,mode="NUC")
      call Write Structure Factors(lun,Num,hkl,mode="NUC")
     end if
     close(unit=lun)
    end do
End Program Calc_structure factors
```

Hkl_Uni: Generates unique reflections in a $\sin\theta/\lambda$ range (constructs, partially, the array of hkl objects)

Structure_Factors: Completes the construction of the array of hkl objects

Write_Structure_Factors: Writes the results in a file

End Program Calc structure factors





```
Program Calc structure factors
use crystallographic symmetry, only: space group type, Write SpaceG
use Atom Module,
                               only: Atoms List Type, Write Atoms List
                               only: Crystal Cell Type, Write Crystal Cell
use crystal types,
use Reflections Utilities,
                               only: Reflection Type, Hkl Uni, get maxnumref
use IO Formats,
                               only: Readn set Xtal Structure, err mess form, err form
use Structure Factor Module, only: Structure Factors, Write Structure Factors
type (space group type) :: SpG
type (Atoms list Type) :: A
type (Crystal Cell Type) :: Cell
type (Reflection Type),allocatable, dimension(:) :: hkl
 character(len=256)
                          :: filcod !Name of the input file
real
                          :: stlmax !Maximum Sin(Theta)/Lambda
 integer
                          :: MaxNumRef, Num, lun=1
    do
     write(unit=*,fmt="(a)") " => Code of the file xx.cif (give xx): "
     read(unit=*,fmt="(a)") filcod
     if(len trim(filcod) == 0) exit
    write(unit=*,fmt="(a)") " => Maximum sinTheta/Lambda: "
     read(unit=*,fmt=*) stlmax
     open(unit=lun,file=trim(filcod)//".sfa", status="replace",action="write")
     call Readn set Xtal Structure(trim(filcod)//".cif",Cell,SpG,A,Mode="CIF")
     If(err form) then
      write(unit=*,fmt="(a)") trim(err mess form)
       exit
     else
       call Write Crystal Cell(Cell,lun)
       call Write SpaceG(SpG,lun)
       call Write Atoms List(A,lun=lun)
      MaxNumRef = get maxnumref(stlmax,Cell%CellVol,mult=SpG%Multip)
       if(allocated(hkl)) deallocate(hkl); allocate (hkl(MaxNumRef))
      call Hkl_Uni(Cell,Spg,.true.,0.0,stlmax,"s",Num,hkl)
       call Structure_Factors(A,SpG,Num,hkl,mode="NUC")
       call Write Structure Factors(lun, Num, hkl, mode="NUC")
     end if
     close(unit=lun)
    end do
End Program Calc_structure_factors
```

Installing and compiling CrysFML using G95 in Windows

G95 in Windows using MinGW

Copy the file g95-MinGW.exe in a temporary directory and double-click on it: select the installation folder and say "yes" to all questions! (e.g. c:\G95, ... warning! do not use "Program Files")

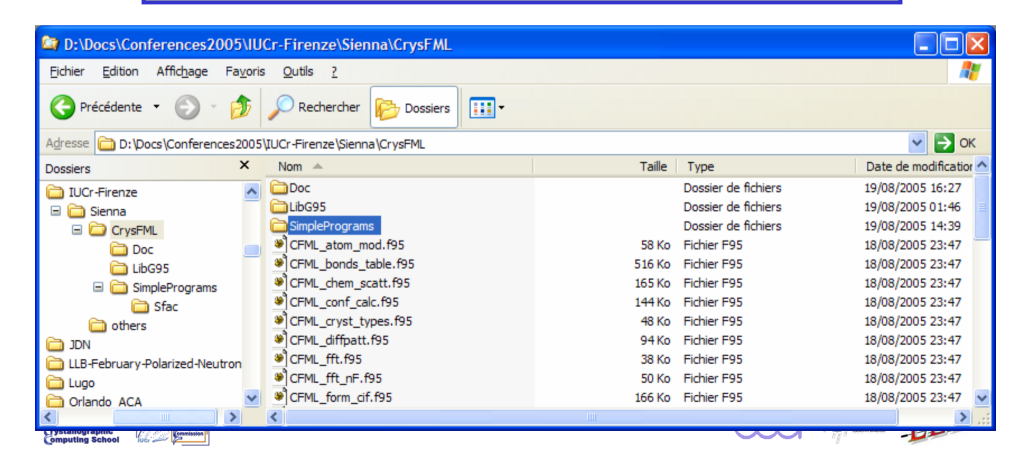
- -Create a directory called "CrysFML" (e.g. c:\CrysFML)
- -Copy the file CrysFML_G95.zip in and extract all files respecting the directory structure
- -Compile and build the library running the file "crysfml_g95.bat"





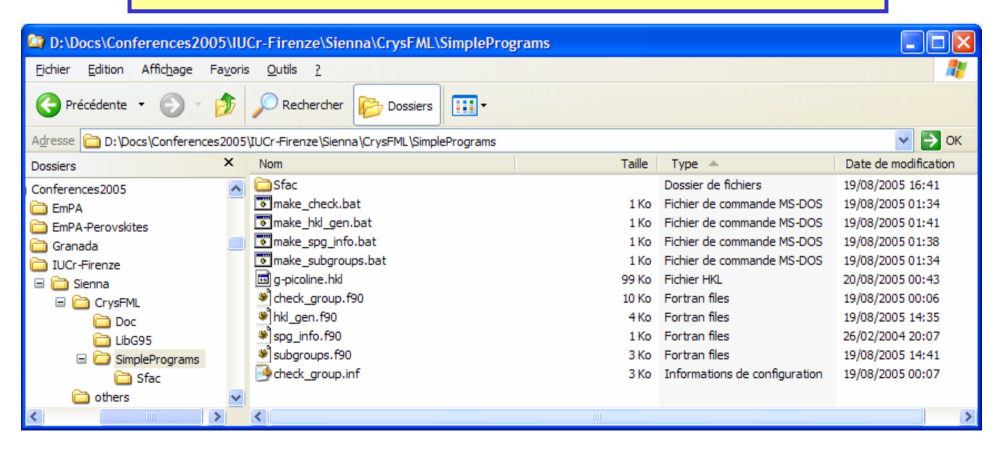
The content of the CrysFML folder and sub-folders

All CrysFML files start with the prefix "CFML_" and have extension .f95



Content of the "SimplePrograms" folder

Four main programs and make*.bat files, one *.hkl file coming from FullProf, a *.inf file and a subfolder called "Sfac"



Content of the "Sfac" folder

Source code files:

```
There are two modules:
```

"observed_reflections" in file "observ.f90"

And

"cost functions" in file "cost functions.f90"

Three main programs:

"Calc_structure_factors" in "sfac_test.f90"

"Optimizing_structure" in "Optim_Sfac.f90"

"Optimizing_structure" in "Opt_restraints.f90"







Input files for CrysFML (CIF and CFL)

```
NiFePO5
Title
                               alpha
                b
                                       beta
        а
                         C
                                              gamma
      7.1882
               6.3924
                       7.4847 90.000 90.000
                                              90.000
Cell
     Space Group
Spgr Pnma
                                               Spin Charge
               х
                        У
                                          occ
             0.0000 0.0000 0.0000 0.74
                                          0.5
                                                2.0
Atom
    Νi
        NI
                                                     2.0
                                                5.0 3.0
Atom Fe
        FE 0.1443 0.2500
                            0.7074 0.63 0.5
             0.3718 0.2500
                            0.1424 0.79 0.5
                                                0.0 5.0
    P
         Р
Atom
    01
             0.3988 0.2500
                           0.64585 0.71 0.5
                                                0.0 - 2.0
Atom
           0.19415 0.2500
Atom
    02 \quad 0
                           0.0253 0.70 0.5
                                                0.0 - 2.0
Atom O3 O
           0.0437
                     0.2500 0.4728 0.83 0.5
                                                0.0 - 2.0
Atom O4 O
           0.3678
                     0.0566 0.2633 0.77 1.0
                                                0.0 - 2.0
! Codes for refinement
Vary xyz 0 1 0 1
!Fix x Fe y O4
!Equal y_Fe y_P 1.00
HKL-OBS mfe.hkl
            1.5
MIN-DSPACING
OPTIMIZE Fobs-Fcal 1.0
SIM ANN
       Name of the cost function
CostNam FobsFcal
         T ini
                  anneal
                             num temps
                    0.95
           8.0
                               90
TemParM
         Nalgor Nconf nm cycl num therm
                                           accept
                         90
                                             0.01
Algor T
         Value of Seed (if SeedVAL = 0, random seed)
SeedVAL
           0
         Treatment of initial configuration
InitCON
         RAN
```

