



## Source routines

An introduction to a new approach to source routines

# Why user routines?

- Fluka offers plenty of built-in tools to define primary beams and estimate quantities
- Sometime these are not enough
- There is the need to write some dedicated code: a “User Routine”
- UR are beyond the scope of this course because of intrinsic difficulties
- Nevertheless, we have a started an effort to make URs more user-friendly
- We want to introduce here the first effort in this direction:  
a new format for the **source routine**
- Why the source routine first? Built-in options allow to sample from a limited number of distribution and not from histograms. This is an effort to overcome this limitation

# The “old” source routine

- Scary for beginners, limited documentation
- Use of **IMPLICIT** and **FORTRAN77** naming convention (see later)

```
* Statement function:
LISNUT (IJ) = INDEX ( PRNAME (IJ), 'NEUTRI' ) .GT. 0
-----
BASIC VERSION
-----
NOMORE = 0
-----
* First call initializations:
IF ( LFIRST ) THEN
* *** The following 3 cards are mandatory ***
TKESUM = ZERZER
LFIRST = .FALSE.
LUSSRC = .TRUE.
* *** User initialization ***
END IF
-----
* Push one source particle to the stack. Note that you could as well
* push many but this way we reserve a maximum amount of space in the
* stack for the secondaries to be generated
* Npflka is the stack counter: of course any time source is called it
* must be = 0
NPFLKA = NPFLKA + 1
* Wt is the weight of the particle
WTFLK ( NPFLKA ) = ONEONE
WEIPRI = WEIPRI + WTFLK ( NPFLKA )
* Particle type (1=proton.....). Ijbeam is the type set by the BEAM
* card
* (Radioactive) isotope:
IF ( IJBEAM .EQ. -2 .AND. LRDBEA ) THEN
IARES = IPROA
IZRES = IPROZ
IISRES = IPROM
CALL STISBM ( IARES, IZRES, IISRES )
IJHION = IPROM * 100000 + MOD ( IPROZ, 100 ) * 1000 + IPROA
IJHION = IJHION * 100 + KXHEAV
IONID = IJHION
CALL DCDION ( IONID )
CALL SETION ( IONID )
LFRPHN ( NPFLKA ) = .FALSE.
* Heavy ion:
ELSE IF ( IJBEAM .EQ. -2 ) THEN
IJHION = IPROM * 100000 + MOD ( IPROZ, 100 ) * 1000 + IPROA
IJHION = IJHION * 100 + KXHEAV
IONID = IJHION
CALL DCDION ( IONID )
CALL SETION ( IONID )
ILOFLK ( NPFLKA ) = IJHION
LFRPHN ( NPFLKA ) = .FALSE.
* Flag this is prompt radiation
LRADDC ( NPFLKA ) = .FALSE.
* Group number for "low" energy neutrons, set to 0 anyway
IGROUP ( NPFLKA ) = 0
* Parent radioactive isotope:
IRDAZM ( NPFLKA ) = 0
* Particle age (s)
AGESTK ( NPFLKA ) = +ZERZER
* Kinetic energy of the particle (GeV)
TKEFLK ( NPFLKA ) = SQRT ( PBEAM**2 + AM ( IONID )**2 )
&
- AM ( IONID )
* Particle momentum
PMOFLK ( NPFLKA ) = PBEAM
PMOFLK ( NPFLKA ) = SQRT ( TKEFLK ( NPFLKA ) * ( TKEFLK ( NPFLKA )
&
+ TWOTWO * AM ( IONID ) ) )
* Check if it is a neutrino, if so force the interaction
* (unless the relevant flag has been disabled)
IF ( LISNUT ( IJBEAM ) .AND. LNUFIN ) THEN
LFRPHN ( NPFLKA ) = .TRUE.
* Not a neutrino
ELSE
LFRPHN ( NPFLKA ) = .FALSE.
END IF
* From this point ....
* Particle generation (1 for primaries)
LOFLK ( NPFLKA ) = 1
* User dependent flag:
LOUSE ( NPFLKA ) = 0
* No channelling:
KCHFLK ( NPFLKA ) = 0
ECRFLK ( NPFLKA ) = ZERZER
* Extra infos:
INFSTK ( NPFLKA ) = 0
LNFSTK ( NPFLKA ) = 0
ANFSTK ( NPFLKA ) = ZERZER
* Parent variables:
IPRSTK ( NPFLKA ) = 0
EKPSTK ( NPFLKA ) = ZERZER
* User dependent spare variables:
DO 100 ISPR = 1, MKBMX1
SPAREK ( ISPR, NPFLKA ) = ZERZER
100 CONTINUE
* User dependent spare flags:
DO 200 ISPR = 1, MKBMX2
ISPARK ( ISPR, NPFLKA ) = 0
200 CONTINUE
* Save the track number of the stack particle:
ISPARK ( MKBMX2, NPFLKA ) = NPFLKA
NPARMA = NPARMA + 1
NUMPAR ( NPFLKA ) = NPARMA
NEVENT ( NPFLKA ) = 0
DFNEAR ( NPFLKA ) = +ZERZER
* ... to this point: don't change anything
AKNSHR ( NPFLKA ) = -TWO TWO
* Cosines (tx,ty,tz)
TXFLK ( NPFLKA ) = UBEAM
TYFLK ( NPFLKA ) = VBEAM
TZFLK ( NPFLKA ) = WBEAM
TZFLK ( NPFLKA ) = SQRT ( ONEONE - TXFLK ( NPFLKA )**2
&
- TYFLK ( NPFLKA )**2 )
* Polarization cosines:
TXPOL ( NPFLKA ) = -TWO TWO
TYPOL ( NPFLKA ) = +ZERZER
TZPOL ( NPFLKA ) = +ZERZER
* Particle coordinates
XFLK ( NPFLKA ) = XBEAM
YFLK ( NPFLKA ) = YBEAM
ZFLK ( NPFLKA ) = ZBEAM
* Calculate the total kinetic energy of the primaries: don't change
* (Radioactive) isotope:
IF ( IJBEAM .EQ. -2 .AND. LRDBEA ) THEN
* Heavy ion:
ELSE IF ( ILOFLK ( NPFLKA ) .EQ. -2 .OR.
&
ILOFLK ( NPFLKA ) .GT. 100000 ) THEN
TKESUM = TKESUM + TKEFLK ( NPFLKA ) * WTFLK ( NPFLKA )
* Standard particle:
ELSE IF ( ILOFLK ( NPFLKA ) .NE. 0 ) THEN
TKESUM = TKESUM + ( TKEFLK ( NPFLKA ) + AMDISC ( ILOFLK ( NPFLKA ) ) )
&
* WTFLK ( NPFLKA )
* RADDLY ( NPFLKA ) = ZERZER
* Here we ask for the region number of the hitting point.
* NREG ( NPFLKA ) = ...
* The following line makes the starting region search much more
* robust if particles are starting very close to a boundary:
CALL GEOCRS ( TXFLK ( NPFLKA ), TYFLK ( NPFLKA ), TZFLK ( NPFLKA ) )
CALL GEOREG ( XFLK ( NPFLKA ), YFLK ( NPFLKA ), ZFLK ( NPFLKA ),
&
NRGFLK ( NPFLKA ), IDISC )
* Do not change these cards:
CALL GEOHSM ( NHSPT ( NPFLKA ), 1, -11, MLATTC )
NLATTC ( NPFLKA ) = MLATTC
CMPATH ( NPFLKA ) = ZERZER
CALL SOEVSU
RETURN
* End of subroutine Source
END
```

# The “new” source routine

- To be distributed in the next release
- Simplified appearance
- Long & meaningful names for variables and routines
- Use of **implicit none** (see later)
- Abundant comments (removed in the snapshot)
- Variables for user’s usage clearly indicated
- Lines not to be edited are “hidden” in routines in the **source\_library.inc** library file
- Old source routines can still be used

Comments removed for clarity in this snapshot

```
double precision sample_gaussian_distribution
double precision sample_flat_distribution

nomore = 0

if ( lfirst ) then
  call initialization( lfirst )
end if

* Beginning of customizable code
* =====

particle_code = IJBEAM
momentum_energy = PBEAM
energy_logical_flag = .false.
particle_weight = ONEONE
divergence_x = DIVBM
divergence_y = DIVBM
gaussian_divergence_logical_flag = LDVGSS
coordinate_x = XBEAM
coordinate_y = YBEAM
coordinate_z = ZBEAM
direction_cosx = UBEAM
direction_cosy = VBEAM
direction_cosz = WBEAM
direction_flag = 0
polarization_cosx = -TWOTWO
polarization_cosy = ZERZER
polarization_cosz = ZERZER
particle_age = ZERZER
kshort_component = -TWOTWO
delayed_radioactive_decay = ZERZER

* End of customizable code - Do not change below
* =====

call set_internal_flags()
call set_beam_type( particle_code, ionid )
call set_particle_momentum_energy_weight( particle_code, ionid,
& momentum_energy, energy_logical_flag, particle_weight )
call set_particle_coordinates( coordinate_x, coordinate_y,
& coordinate_z )
call set_particle_direction( direction_cosx, direction_cosy,
& direction_cosz, direction_flag, divergence_x, divergence_y,
& gaussian_divergence_logical_flag )
call set_particle_polarization( polarization_cosx,
& polarization_cosy, polarization_cosz )
call set_particle_age( particle_code, particle_age,
& kshort_component, delayed_radioactive_decay )
call search_starting_region()

return
*=== End of subroutine Source =====*
end
-:--- source_layer_short.f Bot (76,0) (Fortran)
```

# The “new” source routine

- Without removing comments (notice the ratio code\_lines / comment\_lines)
- Note: the snapshot is not meant to be read
- A step by step look will follow

```
beam momentum. Other beam card parameters are obsolete.
Output variables:
nomore = if > 0 the run will be terminated
-----
include 'source_library.inc'
Source =====
subroutine SOURCE ( nomore )
use source_library
implicit none
logical lfirst
save lfirst
data lfirst / .true. /
integer nomore
integer particle_code, ionid
logical energy_logical_flag
double precision momentum_energy, particle_weight
logical gaussian_divergence_logical_flag
double precision divergence_x, divergence_y
double precision coordinate_x, coordinate_y, coordinate_z
integer direction_flag
double precision direction_cosx, direction_cosy, direction_cosz
double precision khort_component
double precision delayed_radioactive_decay
Function declarations
double precision sample_flat_momentum_energy
double precision sample_gaussian_momentum_energy
double precision sample_maxwell_boltzmann_energy
double precision sample_energy_from_histogram
double precision sample_gaussian_distribution
double precision sample_flat_distribution
nomore = 0
-----
double precision sample_gaussian_distribution
double precision sample_flat_distribution
nomore = 0
if ( lfirst ) then
call initialization( lfirst )
end if
Beginning of customizable code
=====
Accessing variables from the SOURCE card
Values set on the SOURCE card can be accessed with the following
variables:
Numerical values (double precision):
  WHASOU(1), WHASOU(2), ..., WHASOU(18)
SDUM text (8 character):
  SDUSOU
Particle type
-----
particle_code:
  FLUKA particle code of the primary
  See section 5.1 of the FLUKA manual for the list
  of particle code
Default:
  IJBEAM: Particle code of the primary defined on the
  BEAM card
particle_code = IJBEAM
Particle momentum / energy and weight
-----
momentum_energy:
  Set the momentum or the kinetic energy of the primary
energy_logical_flag:
  Possible values:
  .false. : The momentum_energy variable contains the momentum
            of the particle
  .true.  : The momentum_energy variable contains the kinetic
            energy of the particle
particle_weight:
  Sets the initial weight of the primary
Default:
  PBEAM: Particle momentum set on the BEAM card (even if
  energy was used on the BEAM card it)
momentum_energy = PBEAM
energy_logical_flag = .false.
particle_weight = ONEONE
Implemented samplings functions
-----
Implemented samplings functions
---
momentum_energy = ...
... = sample_flat_momentum_energy( [min], [max] )
... = sample_gaussian_momentum_energy( [mean], [fwhm] )
... = sample_maxwell_boltzmann_energy ( [temperature] )
temperature is given in GeV
... = sample_energy_from_histogram( [filename], [energy_unit] )
possible [energy_unit]: "TeV", "GeV", "MeV", "keV", "eV"
Implemented sampling subroutines
sample_exponential_energy_importance:
Input variables:
  - e_min, e_max, intensity_ratio = (int_e_min / int_e_max)
Output variables:
  - momentum_energy, particle_weight
call sample_exponential_energy_importance( [e_min], [e_max],
[intensity_ratio], momentum_energy, particle_weight)
&
Beam angular divergence
-----
divergence_x:
  Beam angular divergence in the X-Z plane
  (before applying beam direction)
divergence_y:
  Beam angular divergence in the Y-Z plane
gaussian_divergence_logical_flag:
  If .true., divergence values are taken as fwhm of a Gaussian
  distribution
  If .false., divergence values are taken as full angle of a flat
  distribution
Note:
  If one of the divergences is equal to zero, while the other is
  not then the sampling will fail and crash. If divergence in
  only one axis is desired, set the other to an infinitesimal
  one, like 1.0D-12.
Default:
  DIVBM: Divergence set on the BEAM card [in rad]
  LDVGS8: Flag indicating Gaussian divergence, set on the BEAM
  card
divergence_x = DIVBM
divergence_y = DIVBM
gaussian_divergence_logical_flag = LDVGS8
Beam starting position
-----
coordinate_x:
  The X coordinate of the beam's starting position
coordinate_y:
  The Y coordinate of the beam's starting position
coordinate_z:
  The Z coordinate of the beam's starting position
The Z coordinate of the beam's starting position
Defaults:
  XBEAM, YBEAM, ZBEAM: Coordinates set on the BEAMPOS card
coordinate_x = XBEAM
coordinate_y = YBEAM
coordinate_z = ZBEAM
Implemented samplings functions
---
coordinate_xyz = ...
... = sample_flat_distribution( [min], [max] )
... = sample_gaussian_distribution( [mean], [fwhm] )
Implemented sampling subroutines
---
sample_annular_location:
Input variables:
  - rmin, rmax [cm]
Output variables:
  - Two coordinates of the sampled location
call sample_annular_location( [rmin], [rmax], coordinate_x,
& coordinate_y )
Beam direction
-----
direction_cosx:
  Direction cosine of the beam with respect to the X-axis
direction_cosy:
  Direction cosine of the beam with respect to the Y-axis
direction_cosz:
  Direction cosine of the beam with respect to the Z-axis
direction_flag:
  Possible values:
  0: All 3 direction cosines are taken into account (values will
  be normalized)
  1: Only direction cosines with respect to the X- and Y-axis
  are taken into account, the third cosine is calculated
  with a positive sign
  2: Only direction cosines with respect to the X- and Y-axis
  are taken into account, the third cosine is calculated
  with a negative sign
Defaults:
  UBEAM, VBEAM, WBEAM: Direction cosines set on the BEAMPOS card
direction_cosx = UBEAM
direction_cosy = VBEAM
direction_cosz = WBEAM
direction_flag = 0
Implemented sampling subroutines
---
sample_isotropic_direction:
Input variables:
  - e_min, e_max, intensity_ratio = (int_e_min / int_e_max)
Output variables:
  - direction_cosx, direction_cosy, direction_cosz
  - direction_isotropic_direction( direction_cosx, direction_cosy,
  & direction_cosz)
Other changeable parameters
-----
For most of the uses none of these three should be changed from
the defaults
polarization_cos:
  The three inputs indicate the direction cosines of the particle
  polarization
particle_age:
  Set the starting age of the primary particle in seconds
khort_component:
  The Khort component of the K0/K0bar
delayed_radioactive_decay:
  the delay for the radioactive decay with respect to the
  standard primary zero time
polarization_cosx = -TWO TWO
polarization_cosy = ZERZER
polarization_cosz = ZERZER
particle_age = ZERZER
khort_component = -TWO TWO
delayed_radioactive_decay = ZERZER
End of customizable code - Do not change below
=====
call set_internal_flags()
call set_beam_type( particle_code, ionid )
call set_particle_momentum_energy_weight( particle_code, ionid,
& momentum_energy, energy_logical_flag, particle_weight )
call set_particle_coordinates( coordinate_x, coordinate_y,
& coordinate_z )
call set_particle_direction( direction_cosx, direction_cosy,
& direction_cosz, direction_flag, divergence_x, divergence_y,
& gaussian_divergence_logical_flag )
call set_particle_polarization( polarization_cosx,
& polarization_cosy, polarization_cosz )
call set_particle_age( particle_code, particle_age,
& khort_component, delayed_radioactive_decay )
call search_starting_region()
return
**** End of subroutine Source =====
end
```

# History of Fortran

- Fortran born in the early 1950s, and the first compiler was released in 1957

## Standards:

- Fortran 66 – The first standard
- Fortran 77 – Extension on Fortran 66
- Fortran 90 – Dynamic memory allocation / introduction of the *Free* format
- Fortran 95 – High performance Fortran specification
- Fortran 2003 – Object oriented programming
- Fortran 2008 / 2018 – Extensions of Fortran 2003

FLUKA is still mostly (if not fully) compatible Fortran 77

This doesn't mean that we can't use newer things in our user routines

# (Unexpected) Features and limitations of Fortran (77)


- Source file format
  - Fixed
  - Free
- Naming convention
- Subprograms
  - Functions
  - Subroutines
- Variable declaration
  - Implicit
  - Explicit

# Source file format

- Fortran 77 uses the *Fixed* file format (extensions: **.f** or **.for**):
  - Maximum 78 characters in one line
  - First 6 are reserved for special function:
    - If the first character is 'c' or '\*', then the line is a comment
    - If the 6<sup>th</sup> position is not empty, then the line is treated as a continuation of the previous one (Often the '&' character is used)
  - With the gfortran compiler it is possible to increase the maximum line length
    - In FLUKA 4 it is set to 132
- Fortran 90 introduced the *Free* format (extensions: **.f90**, [**.f95**, etc.]):
  - Code can start at the 1<sup>st</sup> position
- *Note*: It is not possible to mix both in the same source file  
Gfortran compiler expects the “correct” format based on the file extension.



# Naming convention

- Fortran 77 variable and (subprogram) names:
  - Limited to 6 alphanumerical characters
  - Have to start with a letter
  - Case insensitive
- Starting with Fortran 90 the variable names  Feature exploited in the new source routine
  - Can be up to 31 character long
  - Can contain letters, numbers and underscore ('\_')
  - Have to start with a letter
  - Case insensitive
- *Note:* Try to use descriptive names, to make code readable

# Subprograms

- Two types:
  - Function
    - Has a return value
    - Used in assignment: `variable = function(input_variable_1, ...)`
  - Subroutine
    - Doesn't have a return value
    - Accessible with the CALL statement: `call subroutine(input_variable_1, ...)`
- Passing variables
  - In Fortran you pass the variable, not the value of the variable. (Like passing a pointer in C)
  - This means the subprograms may irreversibly modify the value of the input variables
    - Desired behavior if you want to return multiple variables
    - Can lead to side effects

# Variable declaration

- Fortran by default uses *implicit declaration*, which means the type of the variable (integer, real, etc.) is determined by a preset rule.
- The default rule is:
  - If the variable starts with the letter I, J, K, L, M, or N it is an integer,
  - Otherwise it is a real (single precision float)
- In FLUKA however:
  - Variables with the 1<sup>st</sup> letter I, J, K, L, M, and N are still integers
  - But the others are double precision (floats)
- It is possible (and necessary) to overwrite this with *explicit declaration*, where you manually specify the type of the variable, like:

```
double precision my_intensity  
logical my_flag
```

# Variable declaration

- Biggest issue is that typos remain hidden:
  - If you have a typo in a variable name, the compiler won't raise an error
  - It is a different, but valid variable without a value
  - Using it in calculations will lead to unexpected results
- Other issue is the unexpected type conversion:
  - For example: Information is lost if you want to assign a double precision number to INTEGER
- Solution in the “new” source routine: **implicit none**
  - This statement disables the implicit declaration and every variable has to be manually declared
  - Exception: FLUKAs built in variables don't need to be declared in the source routine
    - (they will remain implicitly declared)
- Convention in the “new” source routine:
  - Variables with uppercase names: FLUKA variables
  - Variables with lowercase names: explicitly declared variables

# Numbers and Constants in User routines

- To keep the high accuracy of the calculation
  - Every variable containing a floating point number should have the type *double precision*
  - The assigned numbers should also be double precision:  
For example: `radius = 2.0D0`  
The 'D' character indicated, that this is number should be treated as double precision.  
If it is 'E' or missing, then the number will be single precision
- To simplify writing numbers FLUKA already defined many numbers as variables:
  - `ONEONE = 1.0D0`
  - `TWOTWO = 2.0D0`
  - `HLFHLE = 0.5D0`
  - `PIPIPI =  $\pi$  = 3.141592...`
  - `TWOPIP =  $2\pi$  = 6.283185...`Full list available in the `dblprc.inc` include file

# Source routine – initialization

```
if ( lfirst ) then  
    call initialization( lfirst )  
end if
```

- Initialization of internal variables
- Only performed the first time the routine is called

```
! beam momentum. Other beam card parameters are obsolete.  
! Output variables:  
! nmore = if > 0 the run will be terminated  
!-----  
! include "source_library.inc"  
!-----  
! Source routine SOURCE ( nmore )  
!-----  
! use source_library  
! explicit none  
! logical lfirst  
! save lfirst  
! data lfirst / .true. /  
! integer nmore  
! integer particle_code, ioid  
! logical energy_logical_flag  
! double precision momentum_energy, particle_weight  
! double precision divergence_x, divergence_y  
! logical gaussian_divergence_logical_flag  
! double precision divergence_x, divergence_y  
! double precision coordinate_x, coordinate_y, coordinate_z  
! integer direction_flag  
! double precision direction_cosx, direction_cosy, direction_cosz  
! double precision polarization_cox, polarization_coy  
! double precision polarization_coz  
! double precision particle_age  
! double precision kabut_component  
! double precision delay_radioactive_decay  
! Function declarations  
! double precision sample_flat_momentum_energy  
! double precision sample_gaussian_momentum_energy  
! double precision sample_maxwell_boltzmann_energy  
! double precision sample_energy_from_histogram  
! double precision sample_gaussian_distribution  
! double precision sample_flat_distribution  
!-----  
! nmore = 0  
!-----  
source_layer.f 118 (11,0) (Fortran)  
-----  
! double precision sample_gaussian_distribution  
! double precision sample_flat_distribution  
!-----  
! if ( lfirst ) then  
!     call initialization( lfirst )  
! end if  
!-----  
! Accessing variables from the SOURCE card  
!-----  
! Values set on the SOURCE card can be accessed with the following  
! variables:  
! Numerical values (double precision):  
!   *MADON(1), *MADON(2), ... *MADON(18)  
! *DUM test (8 characters)  
! *SOURCE  
! Particle type  
! *PARTICLE_CODE of the primary  
! See section 2.1 of the FLUKA manual for the list  
! of particle code  
! Default:  
! *PARTICLE_CODE of the primary defined on the  
! BEAM card  
! *PARTICLE_CODE = 12BEAM  
! Particle momentum / energy and weight  
!-----  
! momentum_energy  
! Set the momentum or the kinetic energy of the primary  
! energy_logical_flag  
! Possible values:  
! .false. : The momentum_energy variable contains the momentum  
! of the particle  
! .true. : The momentum_energy variable contains the kinetic  
! energy of the particle  
! particle_weight  
! Set the initial weight of the primary  
! Default:  
! *PARTICLE_MOMENTUM set on the BEAM card (even if  
! energy was used on the BEAM card !)  
! momentum_energy = *BEAM  
! momentum_energy_logical_flag = .false.  
! particle_weight = *DUMD  
!-----  
! Implemented sampling functions  
!-----  
! Sample momentum energy  
!-----  
! Sample Gaussian momentum energy (beam), (beam), (beam)  
!-----  
! Sample Maxwell Boltzmann energy (temperature)  
!-----  
! Sample energy from histogram (filename), (energy_unit)  
! possible (energy_unit) = "GeV", "MeV", "eV", "keV", "GeV"  
!-----  
! Implemented sampling subroutines  
!-----  
! Sample exponential energy importance:  
! Input variables:  
! *E_MIN, *E_MAX, intensity_ratio = (int_min / int_max)  
! Output variables:  
! *momentum_energy, particle_weight  
! call sample_exponential_energy_importance (e_min, e_max,  
! intensity_ratio, momentum_energy, particle_weight)  
!-----  
! Beam angular divergence  
! *divergence_x  
! Beam angular divergence in the X-Y plane  
! (before applying beam direction)  
! *divergence_y  
! Beam angular divergence in the Y-Z plane  
! *divergence_z  
! Gaussian divergence_logical_flag  
! If .true., divergence values are taken as full angle of a Gaussian  
! distribution  
! If .false., divergence values are taken as full angle of a flat  
! distribution  
! Note:  
! If one of the divergences is equal to zero, while the other is  
! not then the sampling will fail and crash. If divergence in  
! only one axis is desired, set the other to an infinitesimal  
! one, like 1.0E-12.  
! Default:  
! *DIVERGENCE set on the BEAM card (in rad)  
! *LOGDUM Flag indicating Gaussian divergence, set on the BEAM  
! card  
! *divergence_x = DIVERG  
! *divergence_y = DIVERG  
! *divergence_z = DIVERG  
! *gaussian_divergence_logical_flag = LOGDUM  
!-----  
! Beam starting position  
!-----  
! *coordinate_x  
! The X coordinate of the beam's starting position  
! *coordinate_y  
! The Y coordinate of the beam's starting position  
! *coordinate_z  
! The Z coordinate of the beam's starting position  
! Default:  
! *coordinate_x = 0.0  
! *coordinate_y = 0.0  
! *coordinate_z = 0.0  
!-----  
! Implemented sampling functions  
!-----  
source_layer.f 218 (122,0) (Fortran)
```

# Source routine – particle type

`particle_code = IJBEAM`

- By default the particle type given in the BEAM card is taken (**IJBEAM** variable)
- The particle type can be overridden in the source routine
- Possible application: beam made of more than one type particles
- Particle codes explained in Fluka manual section 5.1

```
! beam momentum. Other beam card parameters are obsolete.
!
! Output variables:
!
! nomore = if > 0 the run will be terminated
!
!-----
! include 'source_library.inc'
!-----
! Source =====
!-----
subroutine SOURCE ( nomore )
  use source_library
  implicit none
  logical ifirst
  save ifirst
  data ifirst / .true./
  integer nomore
  integer particle_code, ioid
  logical energy_logical_flag
  double precision momentum_energy, particle_weight
  double precision divergence_x, divergence_y
  logical gaussian_divergence_logical_flag
  double precision divergence_x, divergence_y
  double precision coordinate_x, coordinate_y, coordinate_z
  integer direction_flag
  double precision direction_cosx, direction_cosy, direction_cosz
  double precision polarization_cox, polarization_coy, polarization_coz
  double precision polarization_cox, polarization_coy
  double precision particle_age
  double precision kabut_component
  double precision damped_polarization_cox
!
! Function declarations
!
  double precision sample_flat_momentum_energy
  double precision sample_gaussian_momentum_energy
  double precision sample_maxwell_boltzmann_energy
  double precision sample_energy_from_histogram
  double precision sample_gaussian_distribution
  double precision sample_flat_distribution
!
  nomore = 0
!----- source_layer.f 118 (11,0) (Fortran)

!-----
double precision sample_gaussian_distribution
double precision sample_flat_distribution
!
! nomore = 0
!
! if ( ifirst ) then
!   call initialization( ifirst )
! end if
!-----
! Beginning of customizable code
!-----
! Accessing variables from the SOURCE card
!-----
! Values set on the SOURCE card can be accessed with the following
! variables:
!
! Numerical values (double precision):
!   MWD00(1), MWD00(2), ... MWD00(18)
!   SOW 'xxx' (8 character):
!
! Particle type
! particle_code
! FLUKA particle code of the primary
! See section 5.1 of the FLUKA manual for the list
! of particle code
! Default:
! IJBEAM: Particle code of the primary defined on the
! BEAM card
! particle_code = IJBEAM
!-----
! momentum_energy
! Set the momentum or the kinetic energy of the primary
! energy_logical_flag
! Possible values:
!   .false.: The momentum_energy variable contains the momentum
!           of the particle
!   .true.: The momentum_energy variable contains the kinetic
!           energy of the particle
! particle_weight
! Set the initial weight of the primary
! Default:
! PRBEAM: Particle momentum set on the BEAM card (even if
! energy was used on the BEAM card)
! momentum_energy = PRBEAM
! energy_logical_flag = .false.
! particle_weight = OMBEAM
!-----
! Implemented sampling functions
!-----
!
!-----
! Implemented sampling functions
!
! momentum_energy = ...
! ... = sample_flat_momentum_energy( msh, (msx) )
! ... = sample_gaussian_momentum_energy( beam, (fwhm) )
! ... = sample_maxwell_boltzmann_energy( (temperature) )
! temperature is given in GeV
! ... = sample_energy_from_histogram( (filename), (energy_unit) )
! possible (energy_unit) is "GeV", "MeV", "GeV", "MeV", "eV"
!-----
! Implemented sampling subroutines
!-----
! sample_exponential_energy_importance:
! Input variables:
!   - e_min, e_max, intensity_ratio = (int_e_min / int_e_max)
! Output variables:
!   - momentum_energy, particle_weight
! call sample_exponential_energy_importance( (e_min), (e_max),
! (intensity_ratio), momentum_energy, particle_weight )
!-----
! Beam angular divergence
! divergence_x
! Beam angular divergence in the X-Y plane
! (before applying beam direction)
! divergence_y
! Beam angular divergence in the Y-Z plane
! (before applying beam direction)
! If .true., divergence values are taken as FWHM of a Gaussian
! distribution
! If .false., divergence values are taken as full angle of a flat
! distribution
! Note:
! If one of the divergences is equal to zero, while the other is
! not then the sampling will fail and crash. If divergence in
! only one axis is desired, set the other to an infinitesimal
! one, like 1.0E-12.
! Default:
! DIVM: Divergence set on the BEAM card (in rad)
! DIVY: Flag indicating Gaussian divergence, set on the BEAM
! card
! divergence_x = DIVM
! divergence_y = DIVY
! gaussian_divergence_logical_flag = LOWDSS
!-----
! Beam starting position
! coordinate_x
! The X coordinate of the beam's starting position
! coordinate_y
! The Y coordinate of the beam's starting position
! coordinate_z
! The Z coordinate of the beam's starting position
!----- source_layer.f 438 (175,0) (Fortran)
```





# Source routine – particle momentum/energy

- Some predefined routines (4 functions and 1 subroutine) are already available

Flat spectrum

Gaussian spectrum

Maxwell-Boltzmann spectrum

Spectrum from histogram

Exponential spectrum

(via the change of  
particle's weight)

```
*      Implemented samplings functions
*      ---
*      momentum_energy = ...
*      ... = sample_flat_momentum_energy( [min], [max] )
*      ... = sample_gaussian_momentum_energy( [mean], [fwhm] )
*      ... = sample_maxwell_boltzmann_energy ( [temperature] )
*      temperature is given in GeV
*      ... = sample_energy_from_histogram( [filename], [energy_unit])
*      possible [energy_unit]s: "TeV", "GeV", "MeV", "keV", "eV"
*      Histogram file has to have 3 columns:
*      Emin (of the bin),
*      Emax (of the bin),
*      dN/dE (particle fluence in the bin)
*
*      Implemented sampling subroutines
*      ---
*      sample_exponential_energy_importance:
*      Input variables:
*      - e_min, e_max, intensity_ratio = (int_e_min / int_e_max)
*      Output variables:
*      - momentum_energy, particle_weight
*      call sample_exponential_energy_importance( [e_min], [e_max],
*      & [intensity_ratio], momentum_energy, particle_weight)
```

# Source routine – particle weight

`particle_weight = ONEONE`

- Allows to set the weight of the primary particles
- 99% of the times `weight=1` is ok
- Can be changed to distort the distribution of primaries (e.g. exponential distribution)
- Can be useful if dealing with more than one single type of primaries
- Not for a beginners' use, mentioned here for completeness

```
beam_something. Other beam card parameters are obsolete.
double precision sample_flat_distribution
...
Output variables:
...
nomore = if > 0 the run will be terminated
...
end if
include 'source_library.inc'
...
subroutine SOURCE ( nomore )
use source_library
implicit none
logical ifirst
save ifirst
data list / :row /
integer nomore
integer particle_code, ioxid
integer direction_flag
double precision direction_cosx, direction_cosy, direction_cosz
double precision momentum_energy, particle_weight
double precision divergence_x, divergence_y
double precision coordinate_x, coordinate_y, coordinate_z
integer direction_flag
double precision direction_cosx, direction_cosy, direction_cosz
double precision polarization_cox, polarization_coy, polarization_coz
double precision polarization_coxy
double precision particle_age
double precision kabut_component
double precision delay_radioactive_decay
...
Function declarations
double precision sample_flat_momentum_energy
double precision sample_gaussian_momentum_energy
double precision sample_maxwell_boltzmann_energy
double precision sample_energy_from_histogram
double precision sample_gaussian_distribution
double precision sample_flat_distribution
...
nomore = 0
source_layer.f 118 (71,0) (Fortran)

double precision sample_gaussian_distribution
double precision sample_flat_distribution
...
nomore = 0
if ( ifirst ) then
call initialization( ifirst )
end if
...
Accessing variables from the SOURCE card
...
Values set on the SOURCE card can be accessed with the following
variables:
...
MADNESS values (double precision):
... MADNESS(1), MADNESS(2), ... MADNESS(18)
SOURCE text (8 characters):
... SOURCE
...
Particle type
...
particle_code
...
FLUX: particle code of the primary
...
Default:
...
FLUX: Particle code of the primary defined on the
BEAM card
...
particle_code = 12BEAM
...
Particle momentum / energy and weight
...
momentum_energy
...
Set the momentum or the kinetic energy of the primary
energy_logical_flag
...
.false : The momentum_energy variable contains the momentum
of the particle
.true : The momentum_energy variable contains the kinetic
energy of the particle
particle_weight
...
Set the initial weight of the primary
energy_logical_flag = FLUX
energy_logical_flag = .false.
particle_weight = ONEONE
...
Implemented sampling functions
...
sample_exponential_energy_importance:
Input variables:
...
e_min, e_max, intensity_ratio = (int_e_min / int_e_max)
Output variables:
...
momentum_energy, particle_weight
call sample_exponential_energy_importance (e_min, e_max,
intensity_ratio, momentum_energy, particle_weight)
...
Beam angular divergence
divergence_x
...
Beam angular divergence in the X-Y plane
(before applying beam direction)
divergence_y
...
Beam angular divergence in the Y-Z plane
Gaussian divergence_logical_flag
...
If .false., divergence values are taken as full angle of a Gaussian
distribution
If .false., divergence values are taken as full angle of a flat
distribution
Note:
...
If one of the divergences is equal to zero, while the other is
not then the sampling will fail and crash. If divergence in
only one axis is nonzero, set the other to an infinitesimal
one, like 1.0E-12.
Default:
...
DIVERG: Divergence set on the BEAM card (in rad)
LAYOUT: Flag indicating Gaussian divergence, set on the BEAM
card
divergence_x = DIVERG
divergence_y = DIVERG
gaussian_divergence_logical_flag = LAYOUT
...
Beam starting position
coordinate_x
...
The X coordinate of the beam's starting position
coordinate_y
...
The Y coordinate of the beam's starting position
coordinate_z
...
The Z coordinate of the beam's starting position
source_layer.f 438 (175,0) (Fortran)
```

# Source routine – beam divergence

`divergence_x = DIVBM`



X-Z plane

`divergence_y = DIVBM`



Y-Z plane

`Gaussian_divergence_logical_flag = LDVGSS`

- By default, values are taken from the **BEAM** card
- Divergence values are taken
  - As Gaussian FWHM, if flag set **.true.**
  - As flat distribution full angle, if flag set **.false.**

```
! beam momentum. Other beam card parameters are obsolete.
!
! Output variables:
!
!   nmore = if > 0 the run will be terminated
!
!-----
! include "source_library.inc"
!-----
! Source =====
!
! subroutine SOURCE ( nmore )
! use source_library
! implicit none
! logical ifirst
! save ifirst
! data ifirst / .true./
!
! integer nmore
! integer particle_code, ioid
! double precision momentum_energy, particle_weight
! double precision divergence_x, divergence_y
! logical gaussian_divergence_logical_flag
! double precision coordinate_x, coordinate_y, coordinate_z
! integer direction_flag
! double precision direction_cosx, direction cosy, direction cosz
! double precision polarization_cosx, polarization cosy
! double precision polarization_cosz
! double precision particle_age
! double precision kabut_component
! double precision delay_radioactive_decay
!
! Function declarations
!
! double precision sample_flat_momentum_energy
! double precision sample_gaussian_momentum_energy
! double precision sample_maxwell_boltzmann_energy
! double precision sample_energy_from_histogram
! double precision sample_gaussian_distribution
! double precision sample_flat_distribution
!
! nmore = 0
!----- source_layer.f 118 (11,0) (Fortran)

! double precision sample_gaussian_distribution
! double precision sample_flat_distribution
!
! nmore = 0
!
! if ( ifirst ) then
!   call initialization( ifirst )
! end if
!
!-----
! Beginning of customizable code
!=====
!
! Accessing variables from the SOURCE card
!-----
! Values set on the SOURCE card can be accessed with the following
! variables:
!
! Numerical values (double precision):
!   * MANDOT(1), MANDOT(2), ... MANDOT(18)
!   * SLOW text (8 characters)
!   * SOURCE
!
! Particle type
!-----
! FLUX: particle code of the primary
! See section 3.1 of the FLUX manual for the list
! of particle code
!
! Default: Particle code of the primary defined on the
! BEAM card
!
! particle_code = 12BEAM
!
! Particle momentum / energy and weight
!-----
! FLUX: particle code of the primary
! momentum_energy
! Set the momentum or the kinetic energy of the primary
! energy_logical_flag
!
! Possible values:
!   .false.: The momentum_energy variable contains the momentum
!             of the particle
!   .true.: The momentum_energy variable contains the kinetic
!            energy of the particle
!
! particle_weight = 1
!
! FLUX: Particle momentum set on the BEAM card (even if
! energy was used on the BEAM card !)
```

# Source routine – beam starting position

`coordinate_x = XBEAM`

`coordinate_y = YBEAM`

`coordinate_z = ZBEAM`

- By default, values are taken from the **BEAMPOS** card
- Extended sources can be defined using different starting positions

```
! The X coordinate of the beam's starting position
! Default
! = 0.0, 0.0, intensity_ratio = (int_n_pos / int_n_max)
! Output variables:
! = direction_cosx, direction_cosy, direction_cosz
! = sample_integrator_direction, direction_cosx, direction_cosy, direction_cosz
!
!-----
! Other changeable parameters
! For most of the uses some of these three should be changed from
! the defaults
!-----
! polarization_cosx
! The three inputs indicate the direction cosines of the particle
! polarization
! particle_age
! Get the starting age of the primary particle in seconds
! labor_component
! The labor component of the R1/R2bar
! delayed_radioactive_decay
! The delay for the radioactive decay with respect to the
! standard primary zero time
!-----
! polarization_cosx = 0.0000
! polarization_cosy = 0.0000
! polarization_cosz = 0.0000
! particle_age = 0.0000
! labor_component = 0.0000
! delayed_radioactive_decay = 0.0000
!-----
! End of customizable code -- Do not change below
!-----
! call set_internal_flags()
! call set_beam_type( particle_code, isoid )
! call set_particle_momentum_energy_weight( particle_code, isoid,
! momentum_energy, energy_logical_flag, particle_weight )
! call set_particle_coordinates( coordinate_x, coordinate_y,
! coordinate_z )
! call set_particle_direction( direction_cosx, direction_cosy,
! direction_cosz, direction_flag, divergence_x, divergence_y,
! divergence_z, divergence_logical_flag )
! call set_particle_polarization( polarization_cosx,
! polarization_cosy, polarization_cosz )
! call set_particle_age( particle_age, particle_age,
! labor_component, delayed_radioactive_decay )
! call search_starting_region()
!-----
!*** End of subroutine Source *****
!
!
!
!----- source_layer.f  Bot (261,5) (Fortran)
```

`coordinate_x = XBEAM`  
`coordinate_y = YBEAM`  
`coordinate_z = ZBEAM`

4

`direction_cosx = 0.0000`  
`direction_cosy = 0.0000`  
`direction_cosz = 0.0000`

5

# Source routine – beam starting position

- Some predefined routines (2 functions and 1 subroutine) are already available

Implemented samplings functions

```
---  
coordinate_xyz = ...  
... = sample_flat_distribution( [min], [max] )  
... = sample_gaussian_distribution( [mean], [fwhm] )
```

Flat/Gaussian spatial distribution  
for the chosen coordinate

Implemented sampling subroutines

```
---  
sample_annular_distribution:  
  Input variables:  
    - rmin, rmax [cm]  
    - Coordinates of the center of the annular distribution  
  Output variables:  
    - Modified coordinates of the sampled location  
      (input values have been overwritten)  
  call sample_annular_distribution( [rmin], [rmax],  
  & coordinate_x, coordinate_y )
```

(x,y) coordinates of an annular distribution  
centered on the provided location



# Source routine – beam direction

`direction_cosx = UBEAM`

`direction_cosy = VBEAM`

`direction_cosz = WBEAM`

`direction_flag = 0`

- By default, values are taken from the **BEAM** card
- If `direction_flag = 1`:

`direction_cosz` is calculated from the other 2 and assumed positive

```

* The X coordinate of the beam's starting position
* Default:
* BEAM, YBEAM, ZBEAM: Coordinates set on the BEAM card
coordinates_x = BEAM
coordinates_y = YBEAM
coordinates_z = ZBEAM

* Implemented sampling functions
* ...
* ... = sample_flat_distribution( [min], [max] )
* ... = sample_gaussian_distribution( [mean], [stdev] )
* Implemented sampling subroutines
* ...
* sample_moments_location:
* Input variables:
*   = min, max (cm)
* Output variables:
*   = 2nd coordinates of the sampled location
* call sample_moments_location( [min], [max], coordinates_x,
*   & coordinates_y )

* Beam direction
* -----
* direction_cosx:
* Direction cosine of the beam with respect to the X-axis
* direction_cosy:
* Direction cosine of the beam with respect to the Y-axis
* direction_cosz:
* Direction cosine of the beam with respect to the Z-axis
* direction_flag:
* Possible values:
* 1) All 3 direction cosines are taken into account (values will
* be normalized)
* 2) Only direction cosines with respect to the X- and Y-axis
* are taken into account, the third cosine is calculated
* with a positive sign
* 3) Only direction cosines with respect to the X- and Y-axis
* are taken into account, the third cosine is calculated
* with a negative sign
* Default:
* direction_cosx = UBEAM
* direction_cosy = VBEAM
* direction_cosz = WBEAM
* direction_flag = 0

*
* sample_isotropic_direction:
* Input variables:
*   = min, max (cm)
* Output variables:
*   = 2nd coordinates of the sampled location
* call sample_isotropic_direction( [min], [max], coordinates_x,
*   & coordinates_y )

*
* Input variables:
*   = min, max, intensity_ratio = (int_min / int_max)
* Output variables:
*   = direction_cosx, direction_cosy, direction_cosz
*   = direction_flag
*   = direction_cosz

* Other changeable parameters
* For most of the uses some of these three should be changed from
* the default
* polarization_cosx:
* The three inputs indicate the direction cosines of the particle
* polarization
* particle_age:
* Get the starting age of the primary particle in seconds
* labor_component:
* The labor component of the R1/R2bar
* delayed_radioactive_decay:
* The delay for the radioactive decay with respect to the
* standard primary zero time
* polarization_cosz = *NONE
* polarization_cosy = *NONE
* polarization_cosx = *NONE
* particle_age = *NONE
* labor_component = *NONE
* delayed_radioactive_decay = *NONE

* End of customizable code - Do not change below
* =====
* call set_internal_flags()
* call set_beam_type( particle_code, isid )
* call set_particle_momentum_energy_weight( particle_code, isid,
*   & momentum_energy, energy_logical_flag, particle_weight )
* call set_particle_coordinates( coordinates_x, coordinates_y,
*   & coordinates_z )
* call set_particle_direction( direction_cosx, direction_cosy,
*   & direction_cosz, direction_flag, divergence_x, divergence_y,
*   & divergence_z, divergence_logical_flag )
* call set_particle_polarization( polarization_cosx,
*   & polarization_cosy, polarization_cosz )
* call set_particle_age( particle_code, particle_age )
* call labor_component, delayed_radioactive_decay )
* call search_starting_region()

*
* End of subroutine Source =====
*
*
* source_layer.f  421 (229,5)  (Fortran)

```

4

5





# Source routine – beam direction

- A predefined subroutine is already available

Isotropic direction

```
*      Implemented sampling subroutines
*      ---
*      sample_isotropic_direction:
*      Output variables:
*      - direction_cosx, direction_cosy, direction_cosz
*      call sample_isotropic_direction( direction_cosx, direction_cosy,
*      & direction_cosz)
```



# Source routine – lines not to be touched

- call set\_internal\_flags()
- call set\_beam\_type(...)
- call set\_particle\_momentum\_energy\_weight(...)
- call set\_particle\_coordinates(...)
- call set\_particle\_direction(...)
- call set\_particle\_polarization(...)
- call set\_particle\_age(...)
- call search\_starting\_region()



- These calls pass the provided inputs to Fluka
- Not to be touched for any reason

```

* The X coordinate of the beam's starting position
* Default:
* =====
* BEAM, YSDAM, XBEAM: Coordinates set on the BEAMPOD card
coordinates_x = BEAM
coordinates_y = YSDAM
coordinates_z = XBEAM
* =====
* Implemented sampling functions
* ---
* coordinate_type = ...
* ... = sample_flat_distribution( [min], [max] )
* ... = sample_gaussian_distribution( [mean], [stdev] )
* =====
* Implemented sampling subroutines
* ---
* sample_momentum_location:
* Input variables:
*   = minx, [max] [cm]
* Output variables:
*   = The coordinates of the sampled location
*   call sample_momentum_location( [minx], [maxx], coordinate_x,
*     coordinate_y )
* Beam direction
* =====
* direction_cosines
* Direction cosine of the beam with respect to the X-axis
* direction_cosx = BEAM
* Direction cosine of the beam with respect to the Y-axis
* direction_cosy = YSDAM
* Direction cosine of the beam with respect to the Z-axis
* direction_cosz = XBEAM
* Possible values:
* 1) All 3 direction cosines are taken into account (values will
* be normalized)
* 2) Only direction cosines with respect to the X- and Y-axis
* are taken into account, the third cosine is calculated
* with a positive sign
* 3) Only direction cosines with respect to the X- and Z-axis
* are taken into account, the third cosine is calculated
* with a negative sign
* Default:
* BEAM, YSDAM, XBEAM: Direction cosines set on the BEAMPOD card
direction_cosx = BEAM
direction_cosy = YSDAM
direction_cosz = XBEAM
* =====
* direction_flag = 0
* =====
* Implemented sampling subroutines
* ---
* sample_isotropic_direction:
* Input variables:
*   = [intensity_ratio] [int_max] / [int_min]
* Output variables:
*   = direction_cosx, direction_cosy, direction_cosz
*   = direction_flag
* Other choosable parameters
* For most of the uses some of these three should be changed from
* the default:
* polarization_cos
* The three inputs indicate the direction cosines of the particle
* polarization
* particle_age
* The starting age of the primary particle in seconds
* labor_component
* The labor component of the R1/R2bar
* delayed_radioactive_decay
* The delay for the radioactive decay with respect to the
* standard primary zero time
* polarization_cos = "NONE"
* polarization_cosy = BEERB
* polarization_cosz = BEERB
* particle_age = BEERB
* labor_component = "NONE"
* delayed_radioactive_decay = BEERB
* =====
* End of subroutine code - Do not change below
* =====
* call set_internal_flags()
* call set_beam_type( particle_code, ionid )
* call set_particle_momentum_energy_weight( particle_code, ionid,
* momentum_energy, energy_logical_flag, particle_weight )
* call set_particle_coordinates( coordinate_x, coordinate_y,
* coordinate_z )
* call set_particle_direction( direction_cosx, direction_cosy,
* direction_cosz, direction_flag, divergence_x, divergence_y,
* gaussian_divergence_logical_flag )
* call set_particle_polarization( polarization_cosx,
* polarization_cosy, polarization_cosz )
* call set_particle_age( particle_age, particle_age )
* call set_particle_labor( labor_component, delayed_radioactive_decay )
* call search_starting_region()
* =====
* End of subroutine BEERB
* =====

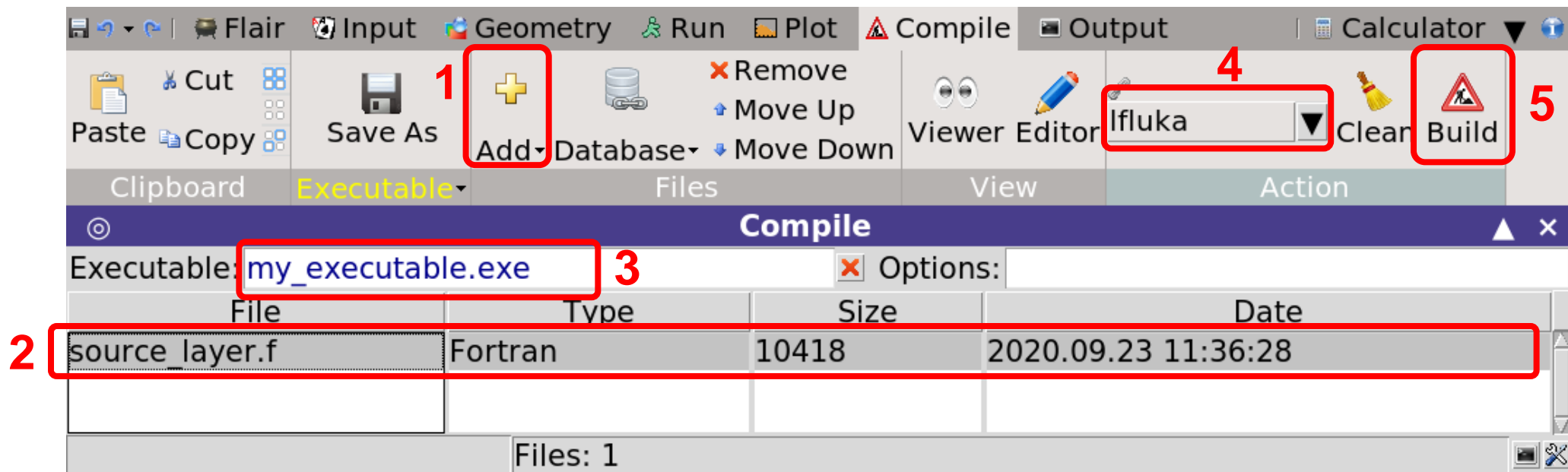
```

# Some predefined FLUKA random sampling routines

- Fluka offers some predefined routines for random sampling
- `my_variable = FLRNDM(XDUMMY)`  
Assigns a 64-bit random number in [0,1)
- `call FLNRRN (gauss1)`  
Returns a Gaussian distributed random number
- `call FLNRR2 (gauss1 , gauss2)`  
Returns two uncorrelated Gaussian distributed random numbers
- `call SFECFE (sint , cost)`  
Returns sine and cosine of a random azimuthal angle

# Compile

1. Add the routine
  2. Verify that it appears
  3. Insert the name of your executable
  4. Select the compiler
  5. Build your executable
- Warning: the library file (`source_library.inc`) must be in the same directory of the source file (`source_layer.f`)



# SOURCE card and passing parameters

- To invoke a source routine it is necessary to add a **SOURCE** card
- A **SOURCE** card can be empty or can be used to pass parameters to the routine
- Maximum 18 numerical values and 1 string can be passed
- **SOURCE** card and **BEAM** card can coexist
- Good practice advice:  
even if the beam energy/momentum is defined in the source routine,  
specify it in the **BEAM** card as it is used internally as default for some scoring

```
† SOURCE          #1: 7.          #2: 250.        #3: 12.5
                    sdum: linksour #4: 3.75       #5:             #6:
                    #7:             #8:             #9:
                    #10:            #11:            #12:
                    #13:            #14:            #15:
                    #16:            #17:            #18:
```

# Time to do some hands-on practice!

- We will now see together a small example of “new” source routine



