

Explain the following sample codes:

- (1) `udm_int_sample1.f90`
- (2) `udm_int_sample2.f90`
- (3) `udm_part_sample1.f90`
- (4) `udm_part_sample2.f90`
- (5) `udm_Manager.f90`

You can create your own user code by modifying the parts described below.

File name rules

The file name defining **interactions** should be as follows

```
udm_int_<Your File Name>.f90
```

The file name defining **particles** should be as follows

```
udm_part_<Your File Name>.f90
```

[Note]: You can name files other than these by writing directly to the makefile.

Description of the ***interaction file*** (*udm_int_*.f90*)

Two functions/subroutines that users primarily modify:

```
function Xsec_per_atom(...)
```

[Role]: Defines the total cross-section of the interaction.

```
subroutine generate_final_state
```

[Role]: Sampling final state energies, scattering angles, etc.

Description of the ***particle file*** (*udm_part_*.f90*)

Three functions/subroutines that users primarily modify:

```
function mass()
```

[Role]: Define the mass of a particle.

```
function lifetime()
```

[Role]: Define the lifetime.

```
subroutine decay
```

[Role]: Define the decay pattern.

(1) udm_int_sample1.f90

An example of an interaction in which a user-defined particle (X) is emitted by an incident electron or positron.



```
1  ! Interaction Template Version = 1.0
2
3  !=====
4  module udm_int_sample_1
5  !=====
6
7  use udm_Parameter
8  use udm_Utility
9  implicit none
10 private ! Functions and variables are set to private by default.
11 public :: caller ! The 'caller' subroutine should be
12
13 !-----
14 ! Default variables
15 !-----
16 character(len=99), parameter :: Name = "my_interaction_1" !
17 double precision, allocatable, save :: Parameters(:) ! Parameters
18 integer, parameter :: num_initial = 2 ! The number of incident
19 integer, save :: kf_initial(num_initial) = (/ 11, -11 /) ! The k
20 !-----
21 ! User variables
22 !-----
23 ! integer i,j
24 ! double precision x,y
25 integer kf_X
26 contains
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330 !=====
331 end module udm_int_sample_1
332 !=====
333
334
```

Defines the **module name**. This is used in udm_Manager.f90 described below.

Define a **Name** for this interaction. This is used in the input file.

The number of incident particles causing this interaction.

kf-codes of the incident particles causing this interaction. In this case, the electron (11) and positron (-11)

(1) udm_int_sample1.f90

```
39  !=====
40  subroutine initialize
41  ! This subroutine is called only once at the beginning of the calculation.
42  !=====
43  kf_X=Parameters(1) ! kf-codes (particle ID) of X (outgoing particle).
44  end subroutine initialize
45
```

Parameters entered in the [user defined interaction] section of the input file are automatically assigned to the array `Parameters(i)` in the source code.

(Example of input file)

```
[ user defined interaction ]
n_int = 2
```

\$ Name	Bias	Parameters
my_interaction_1	1	900000
my_interaction_3	100	900000 1.1 2.2

This value is available as `Parameters(1)` in the source code whose `Name` is `my_interaction_1`.

`Parameters(1)`

`Parameters(2)`

`Parameters(3)`

(In `my_interaction_3`)

(1) `udm_int_sample1.f90`

Define the total cross section per atom in the function `Xsec_per_atom`. Units are in barn.

```
48  !=====
49  double precision function Xsec_per_atom(Kin,Z,A)
50  ! Integrated cross section per an atom.
51  ! Unit: barn (10^-24 cm2)
52  implicit none
53  double precision Kin ! Kinetic energy of incident particle [MeV]
54  integer Z ! Atomic number of target atom
55  integer A ! Mass number of target atom
56  !-----
57  ! [Variables available in this function]
58  ! udm_kf_incident: The kf-codes (particle IDs) of the incident particles.
59  !-----
60
61  if(Kin < 100.0) then
62    Xsec_per_atom=0.0
63    return
64  endif
65
66  if (udm_kf_incident == 11) then
67    Xsec_per_atom=1e-6*Z
68  else if(udm_kf_incident == -11) then
69    Xsec_per_atom=2e-6*Z
70  else
71    print*, "error"
72    stop
73  endif
74
75  return
76  end
```

The kf-code of the incident particle is assigned to `udm_kf_incident` in the code.

In this case, the cross section is 0 barn if the incident particle kinetic energy (= Kin) is less than 100 MeV.

($10^{-6} * Z$) barn, for electron.
($2 * 10^{-6} * Z$) barn, for positron.

(1) udm_int_sample1.f90

First half

```
112 !=====
113 subroutine generate_final_state
114 !=====
115 ! Subroutine to determine final state info
116 ! In this example, the X and e+ momenta of
117 ! [Variables available in this subroutine]
118 ! udm_Kin : Kinetic Energy of the
119 ! udm_Kin : Kinetic Energy of the
120 ! udm_Kin : Kinetic Energy of the
121 ! udm_Kin : Kinetic Energy of the
122 integer Z_A_hit(2), Z, A
123 integer i, n_sampling_max
124 double precision Kout, Kout_min, Kout_max
125 double precision P, R
126 double precision m_X, E_X, p_X, theta_X
127 double precision m_e, E_e, p_e, theta_e
128 !-----
129 ! You may use Z and A for final state varia
130 ! Z and A of a target material are automati
131 Z_A_hit=get_hit_nuclide_Z_A(udm_Kin)
132 Z=Z_A_hit(1)
133 A=Z_A_hit(2)
134 !-----
135 ! Obtain interacted atom
136 ! information (Z, A) if needed
137 !=====
138 !=====
139 !=====
140 ! Range of sampling variable
141 Kout_min=0.0d0
142 Kout_max=udm_Kin-get_mass(kf_X)
143 !-----
144 ! Start sampling
145 n_sampling_max = 100000
146 !-----
147 ! The distribution of the emission energy (Kout) of
148 ! the particle X is defined by another function.
149 Kout=get_random(Kout_min, Kout_max)
150 !-----
151 ! Rejection sampling method.
152 P=distribution(udm_Kin, Z, A, Kout) ! The
153 R=get_random_01()
154 ! If P > R, this Kout is accepted
155 if(P > R) then
156 ! Accepted!!!
157 ! Initialization required before fill
158 call initialize_udm_event_info
159 !-----
160 !-----
161 !-----
162 !-----
163 ! Set number of final state
164 set_final_state_number = 2
165 !-----
```

Second half

```
166 !-----
167 ! [Important Notice]
168 ! Here, you set the final state momenta to the following "set_*" array,
169 ! assuming the direction of the momentum of the incident particle to be
170 ! positive along the "Z-axis". The final state momenta are automatically
171 ! rotated based on the actual direction outside of this subroutine.
172 !-----
173 !-----
174 m_X=get_mass(kf_X)
175 E_X=Kout+m_X
176 p_X=sqrt(E_X**2-m_X**2)
177 theta_X=get_random(0.0d0,0.1d0)
178 !-----
179 ! Set 4-momentum of X
180 set_kf(1) = kf_X
181 set_Total_Energy_in_MeV(1) = E_X
182 set_Px_in_MeV(1) = p_X*sin(theta_X)
183 set_Py_in_MeV(1) = 0.0d0
184 set_Pz_in_MeV(1) = p_X*cos(theta_X)
185 !-----
186 m_e=get_mass(udm_kf_incident)
187 E_e=udm_Kin+m_e-E_X
188 p_e=sqrt(E_e**2-m_e**2)
189 theta_e=get_random(0.0d0,0.1d0)
190 !-----
191 ! Set 4-momentum of e+-
192 set_kf(2) = udm_kf_incident
193 set_Total_Energy_in_MeV(2) = E_e
194 set_Px_in_MeV(2) = p_e*sin(theta_e)
195 set_Py_in_MeV(2) = 0.0d0
196 set_Pz_in_MeV(2) = p_e*cos(theta_e)
197 !-----
198 !-----
199 ! [Additional Quantities]
200 ! set_isomer_level(1) = ?? ! (Default=0) 0: Not nuclear isom
201 ! set_excitation_energy_in_MeV(1) = ?? ! (Default=0.0) Excitation energy [
202 !-----
203 ! The final states are recorded.
204 call fill_final_state
205 !-----
206 !-----
207 !-----
208 return
```

Assuming that the direction of the incident particle in the starting state is the Z-axis, the momentum of the end state is calculated.

Assume the direction of the incident particle is the Z-axis, and assign the kf-code, energy, and momenta of the 1st final state.

In this example, we assign the information for the 2nd final state.

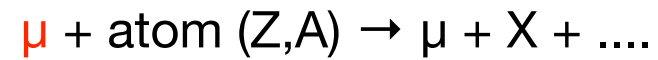
[Important]: Call `fill_final_state`, after the assignment.

[Important]: Initialize the array before setting the final state information.

[Important]: Specify the number of final states

(2) `udm_int_sample2.f90`

Code similar to `udm_int_sample1.f90`. However, the incident particles are **muons**.



```
16  character(len=99), parameter :: Name = "my_interaction_2"
17  double precision, allocatable, save :: Parameters(:) ! Par
18  integer, parameter :: num_initial = 2 ! The number of inci
19  integer, save :: kf_initial(num_initial) = (/ 13, -13 /) !
```

[Changes]

The incident particles are
 μ^- (13) and μ^+ (-13).

(3) udm_part_sample1.f90

Define the particle X generated by "udm_int_sample1.f90".

```
1  ! Particle Template Version = 1.0
2
3  !=====
4  module udm_part_sample_1
5  !=====
6
7  use udm_Parameter
8  use udm_Utility
9  implicit none
10 private ! Functions and variables are set to private by default
11 public :: caller ! The 'caller' subroutine should be public
12
13
14 Default variables
15
16 character(len=99), parameter :: Name = "my_particle_1"
```

Define an arbitrary **module name**.
This is used in udm_Manager.f90
described below.

Define a **Name** for this particle. This is
used in the input file.

```
145 !=====
146 end module udm_part_sample_1
147 !=====
```

(3) `udm_part_sample1.f90`

```
46  !=====
47  double precision function mass() ! Unit: MeV
48  !=====
49  mass=50.0 ! 50 MeV
50  return
51  end
52
53  !=====
54  double precision function lifetime() ! Unit: Second
55  ! The value should be greater than 0.
56  !=====
57  lifetime=0.1e-9 ! 0.1 nano second
58  return
59  end
60
61  !=====
62  subroutine decay
63  !=====
64  ! Kinetic Energy of the incident particle [MeV]
65  !-----
66  !-----
67  if(0.5 > get_random_0to1()) then
68      call two_body_decay_uniform(12,12) ! X -> 2 neutrinos (50%)
69  else
70      call three_body_decay_uniform(12,12,12) ! X -> 3 neutrinos (50%)
71  endif
72  end subroutine decay
```

Set mass to 50 MeV

Set lifetime to 0.1×10^{-9} second

Branching Ratio 50%

Get a random number in the range of 0 to 1

`decay` subroutine
defines decay pattern

`two_body_decay_uniform (kf1, kf2)`

Subroutine for 2-body decay, where kf-code of the final state is kf1 and kf2.

`three_body_decay_uniform (kf1, kf2, kf3)`

Subroutine for 3-body decay, where kf-code of the final state is kf1, kf2, and kf3.

In this case, 50% decays into two electron neutrinos (kf=12) and 50% into three electron neutrinos (unphysical, but for simplicity)

(4) udm_part_sample2.f90

Code similar to `udm_part_sample1.f90`.

The difference is that the mass and lifetime use the values entered in the [user defined particle] section.

```
4  module udm_part_sample_2
      :
16  character(len=99), parameter :: Name = "my_particle_2"
      :
46  !=====
47  double precision function mass() ! Unit: MeV
48  !=====
49  mass=Parameters(1) ! MeV
50  return
51  end
52
53  !=====
54  double precision function lifetime() ! Unit: Second
55  ! The value should be greater than 0.
56  !=====
57  lifetime=Parameters(2) ! second
58  return
59  end
```

(Example of input file)

```
[ user defined particle ]
n_part = 1

$ Name          kfcode    Parameters
my_particle_2   900000    100  1.0e-9
```

Parameters(1)

Parameters(2)

(5) udm_Manager.f90

Write the module name of the user code you want to use in `udm_Manager.f90`.

Line up **all modules**
you want to use.

```
1  module udm_Manager
2  use udm_Parameter
3
4  !=====
5  ! [udm_int]
6  use udm_int_sample_1, caller_udm_int_sample_1 => caller
7  use udm_int_sample_2, caller_udm_int_sample_2 => caller
8  ! [udm_part]
9  use udm_part_sample_1, caller_udm_part_sample_1 => caller
10 use udm_part_sample_2, caller_udm_part_sample_2 => caller
11 !=====
12
```

use <module name>, caller_<module name> => caller

Line up **all interactions**
you want to use.

```
15
16
17
18
19 subroutine user_defined_interaction(action,index)
20 integer action,index
21 !=====
22 call caller_udm_int_sample_1(action,index)
23 call caller_udm_int_sample_2(action,index)
24 !=====
25 end subroutine user_defined_interaction
```

call caller_<module name>(action,index)

Line up **all particles**
you want to use.

```
28
29
30 subroutine user_defined_particle(action)
31 integer action,index
32 do index=1,udm_part_nMax
33 !=====
34 call caller_udm_part_sample_1(action,index)
35 call caller_udm_part_sample_2(action,index)
36 !=====
37 enddo
```

call caller_<module name>(action,index)

```
39
40 end module udm_Manager
```