



Beginners' FLUKA Course

Overview

General concepts:

Analog vs. biased Monte Carlo calculation

Biasing options

(only the most important / common options available in FLUKA)

Importance biasing Weight Windows Leading particle biasing Multiplicity tuning Non-analogue neutron absorption Biasing mean-free paths - decay lengths biasing - hadronic inelastic interaction lengths

User-written biasing

Analog vs. Biased - 1

Analog Monte Carlo

- samples from actual phase space distributions
- predicts average quantities and all statistical moments of any order
- preserves correlations and reproduces fluctuations (provided the physics is correct...)
- is (almost) safe and can (sometimes) be used as "black box"

BUT

- is inefficient and converges very slowly
- fails to predict important contributions due to rare events

Analog vs. Biased - 2

Biased Monte Carlo

- samples from artificial distributions and applies a weight to the particles to correct for the bias
- predicts average quantities, but not the higher moments (on the contrary, its goal is to minimize the second moment)
- same mean with smaller variance, *i.e.*, faster convergence

BUT

- cannot reproduce correlations and fluctuations
- requires physical judgment, experience and a good understanding of the problem (it is not a "black box"!)
- in general, a user does not get the definitive result after the first run, but needs to do a series of test runs in order to optimize the biasing parameters

balance between user's time and CPU time

Reduce variance or CPU time ?

A Figure of Merit

Computer cost of an estimator = $\sigma^2 x t$

(σ^2 = Variance, *t* = CPU time per primary particle)

- some biasing techniques are aiming at reducing σ , others at reducing t
- often reducing σ increases *t*, and *viceversa*
- therefore, minimizing $\sigma^2 x t$ means to reduce σ at a faster rate than t increases or viceversa
- the choice depends on the problem, and sometimes a combination of several techniques is most effective
- bad judgment, or excessive "forcing" on one of the two variables can have catastrophic consequences on the other one, making computer cost explode

 it is the simplest, most "safe" and easiest to use of all biasing techniques

• importance biasing combines two techniques:

Surface Splitting (reduces σ but increases t) Russian Roulette (does the opposite)

 the user assigns a relative importance to each geometry region (the actual absolute value doesn't matter), based on

expected fluence attenuation with respect to other regions
 probability of contribution to score by particles entering the region

Surface Splitting

A particle crosses a region boundary, coming from a region of importance I_1 and entering a region of *higher* importance $I_2 > I_1$:

- the particle is replaced on average by $n=I_2/I_1$ identical particles with the same characteristics
- the weight of each "daughter" is multiplied by I_1/I_2

If I_2/I_1 is too large, excessive splitting may occur with codes which do not provide an appropriate protection.

An internal limit in FLUKA prevents excessive splitting if I_2/I_1 is too large (> 5), a problem found in many biased codes.

Russian Roulette

A particle crosses a region boundary, coming from a region of importance I_1 and entering a region of *lower* importance $I_2 < I_1$:

- the particle is submitted to a random survival test: with a chance I_2/I_1 the particle survives with its weight increased by a factor I_1/I_2
- with a chance $(1 I_2/I_1)$ the particle is killed

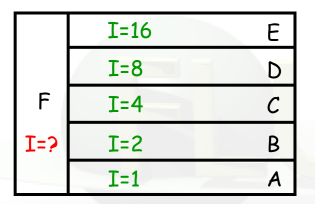
Importance biasing is commonly used to maintain a uniform particle population, compensating for attenuation due to absorption or distance. In FLUKA it can be tuned per particle type. Note:

In FLUKA, for technical reasons, importances are internally stored as integers. Therefore, importances can only take values between 0.0001 and 100000. An input values 0.00015 is read as 0.0001, 0.00234 is read as 0.0023, *etc.*

There is also a user routine USIMBS which allows to assign importances not only at boundaries, but at each step, according to any logic desired by the user (as a function of position, direction, energy,...). Very powerful, but *time-consuming* (it is called at each step!). The user must balance the time gained by biasing with that wasted by calls.

Problems

Although importance biasing is relatively easy and safe to use, there are a few cases where caution is recommended, *e.g.*:



Which importance shall we assign to region F? Whatever value we choose, we will get inefficient splitting/RR at the boundaries.

Another case is that of splitting in vacuum (or air). Splitting daughters are strongly correlated: it must be made sure that their further histories are differentiated enough to "forget" their correlation. (Difficult to do in ducts and mazes)

This applies in part also to muons: the differentiation provided by multiple scattering and by dE/dx fluctuations is not always sufficient.

Input card: **BIASING**

BIASING	0.0	0.0	4.64	8.	18.	2.PRINT	
The meaning of WHA	•••	WHAT(6) and SDUM	is differe	nt dependin	g on the	
f WHAT(1)							
WHAT (2	l) specifi	es the p	articles	to be bia	ased		
	= 1. = 2.	0 : hadr 0 : elec	particles ons and m trons, po energy ne	sitrons a	and photor	ıs	
WHAT (2	2) = (see	multipli	city tuni	ng)			
WHAT (3	_	_	ance (Def				

Input card: **BIASING**

Ψ						
BIASING	0.0	0.0	4.64	8.	18.	2.PRINT
If WHAT(1) >	= 0.0 :					
WHAT (4)	= lower bour	nd of th	e region i	.ndices/	names (Def	^f ault = 2.0)
WHAT (5)	= upper bou	nd of th	e region i	.ndices/	names (Def	fault = WHAT(4))
WHAT (6)	= step leng	th in as	signing ir	dices ([Default = 1.0))
SDUM	<pre>= PRINT : = NOPRINT: = USER: = NOUSER: = RRPRONLY: = (blank):</pre>	counters importan reset to	•	ted (cance cording to cels any pr	els any previ o the user d revious USE	

(Default: NOPRINT, NOUSER, multiplicity biasing for all generations)

Input card: **BIASING**

₽						
BIASING	2.0	0.0	10.0	7.0	11.0	2.0
BIASING	2.0	0.0	15.0	8.0	9.0	0.0
BIASING			3.0	4.0	0.0	
If WHAT(1) <	0.0 :					
WHAT(1)	_		g that all particle-d	-	_	es shall be
WHAT (2)	>= 0.0 : < 0.0 :	-	ng factor set to the		value 1.	0
WHAT (3)	= lower b	ound of	the parti	cle indi	ces/names	(Default: = 1.0)
WHAT (4)	= upper b (Defaul		-			articles otherwise)
WHAT (5)	= step le	ngth in	assigning	partic.	le indice	s (Default: 1.0)
WHAT(6)	= not use	d				
SDUM	NOPRIMA	Ry : imp	ortance bias ortance bias		•	
Defau	t = PRIMAR	У				

Input cards: WW-FACTO WW-THRES WW-PROFI

The weight window technique is a combination of splitting and Russian Roulette, but it is based on the absolute value of the weight of each individual particle, rather than on relative region importance.

The user sets an upper and a lower weight limit, generally as a function of region, energy and particle. Particles having a weight larger than the upper limit are split, those with weights smaller than the lower limit are submitted to Russian Roulette (killed or put back "inside the window").

Weight windows are a more powerful biasing tool than importance biasing, but they require also more experience and patience to set it up correctly. "It is more an art than a science" (Quote from the MCNP manual)

The use of weight windows is essential whenever other biasing techniques generate large weight fluctuations in a given phase space region.

Weight Windows - 2

Input cards: WW-FACTO WW-THRES WW-PROFI

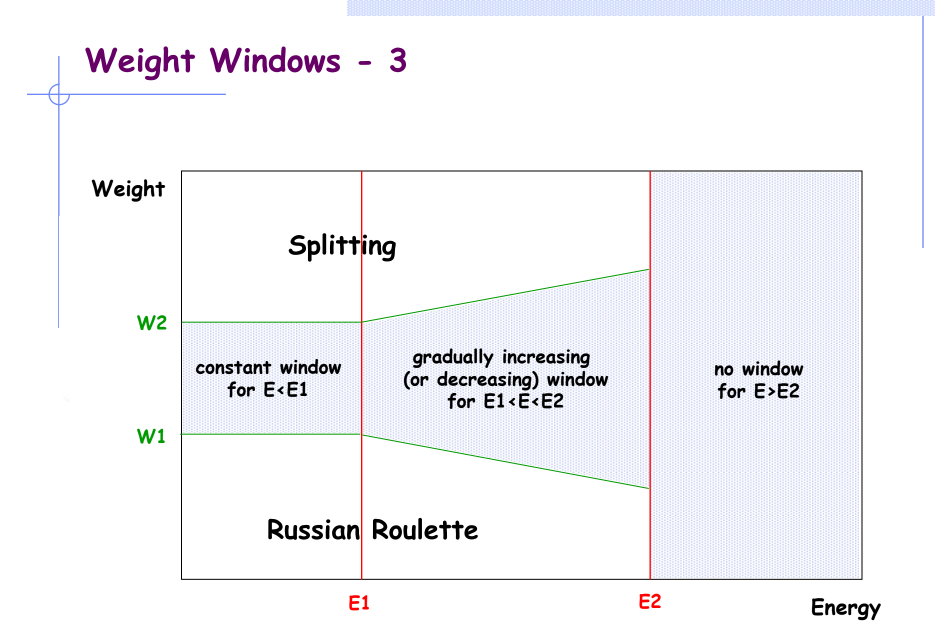
Killing a particle with a very low weight (with respect to the average for a given phase space region) decreases t but has very little effect on the score (and therefore on σ)

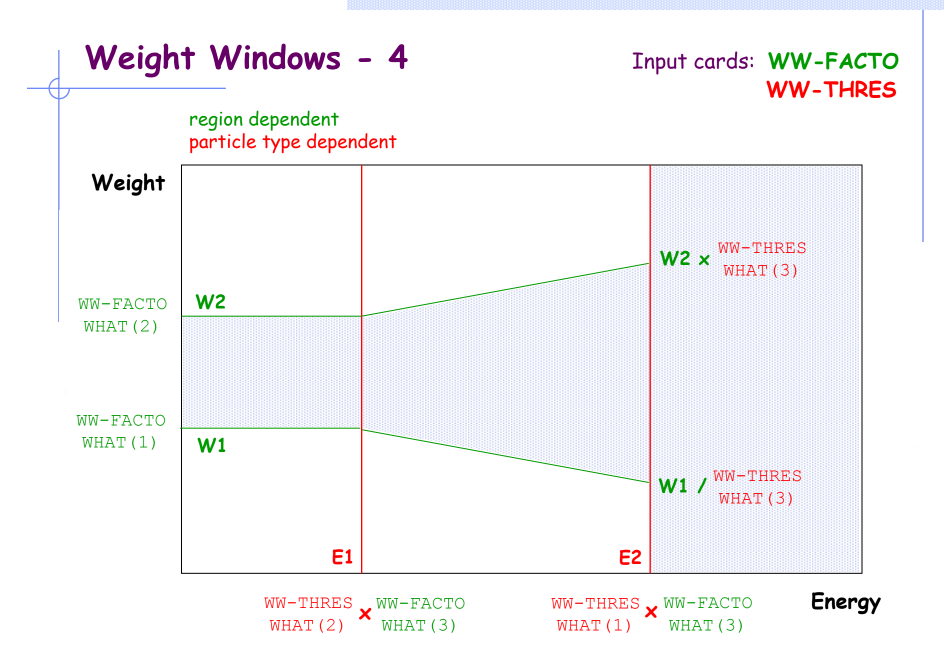
Splitting a particle with a large weight

- increases t (in proportion to the number of additional particles to be followed)
- but at the same time *reduces* σ by avoiding large fluctuations in the contributions to scoring.

The global effect is to reduce $\sigma^2 t$

A too wide window is of course ineffective, but also too narrow windows should be avoided. Otherwise, too much CPU time would be spent in repeated splitting / Russian Roulette. A typical ratio between the upper and the lower edge of the window is about 10. It is also possible to do Russian Roulette without splitting (setting the upper window edge to infinity) or splitting without Russian Roulette (setting the lower edge to zero)





```
Weight Windows - 5
                                                     Input card: WW-FACTO
               13.0
                       120.0
                                  1.5
                                                    31.0
                                                              2.0
WW-FACTO
                                           27.0
Defines Weight Windows in selected regions
     WHAT(1) >= 0.0 : Window "bottom" weight
              < 0.0 : resets to -1.0 (no Russian Roulette, Default)
               Weight below which Russian Roulette is played at the lower energy threshold
               (set by WW-THRES).
     WHAT(2) > 1.7 \times WHAT(1) : Window "top" weight
              = 0.0 : ignored
              =< 1.7 * WHAT(1) : resets to infinity (no Splitting, Default)
              Weight above which Splitting is applied at the lower energy threshold (set by
              WW-THRES).
             > 0.0 : Multiplicative factor (Default: 1.0)
     WHAT(3)
              = 0.0 : ignored
              < 0.0 : resets to 1.0
              Factor to be applied to the two energy thresholds for
              Russian Roulette / Splitting (set by WW-THRES)
```

Vindo	ws - 6			Inpu	t card:	WW-FACTO
13.0	120.0	1.5	27.0	31.0	2.0	
= lower	bound of	the regi	on indic	es/names	(Default	= 2.0)
= upper	bound of	the regi	on indic	es/names	(Default	= WHAT(4))
= step	length in	assignin	g indice	s/names (l)efault =	1.0)
<mark>weigh</mark> (see V	t-window pr	ofile to be (Default =	applied in 1.0)	the regions		· · ·
	13.0 = lower = upper = step : a numb weigh (see V	13.0 120.0 = lower bound of = upper bound of = step length in : a number from 1.0 weight-window pr (see WW-PROFI).	 lower bound of the regit upper bound of the regit step length in assigning a number from 1.0 to 5.0 in an weight-window profile to be (see WW-PROFI). (Default = 	 13.0 120.0 1.5 27.0 = lower bound of the region indic = upper bound of the region indic = step length in assigning indice : a number from 1.0 to 5.0 in any position, weight-window profile to be applied in (see WW-PROFI). (Default = 1.0) 	 13.0 120.0 1.5 27.0 31.0 = lower bound of the region indices/names = upper bound of the region indices/names = step length in assigning indices/names (I : a number from 1.0 to 5.0 in any position, indicating tweight-window profile to be applied in the regions (see WW-PROFI). (Default = 1.0) 	 13.0 120.0 1.5 27.0 31.0 2.0 = lower bound of the region indices/names (Default = upper bound of the region indices/names (Default = step length in assigning indices/names (Default = : a number from 1.0 to 5.0 in any position, indicating the low-e weight-window profile to be applied in the regions selected (see WW-PROFI). (Default = 1.0)

Attention: Option WW-FACTO alone is not sufficient to define a weight window. One or more WW-THRES cards are also necessary in order to activate the window.

Weight	Windo	ows - 7	,		Input	t card: W	N-THRE
WW-THRES	2.0	0.05	2.4	3.0	7.0	0.0	
Defines the	energy]	Limits and	particle	e-depender	nt modifi	cation fa	ctors
WHAT(1)	= 0.0:	: upper ki: Low-energ : ignored : any prev	y neutrons:	lower grou	p number (included)	led
WHAT (2)	>= 0.0	and < WHA	T(1): log y neutrons	ver kineti upper grou	ic energy p number (threshol included)	
WHAT (3)	> 0.0:	amplifica at the highe				ght window (by WHAT(1	
		The weight multiplying the same fo	window at by WHAT(actor the lo	the higher e 3) the upper ower weight	energy thro r weight lir limit. (Def	eshold is obt nit and by d ault = 10.0)	tained by ividing by
	< 0.0:	WHAT(3) weight limi (Default =	ts for the			the lower a WHAT(4-6)	• •

Weight V	Vindo	ws - 8			Inpu	t card: W	W-TH
W-THRES	2.0	0.05	2.4	3.0	7.0	0.0	
WHAT (4)	Note t	hat particle	e index 40 i	ndicates lo	w-energy ne	s (Default : eutrons (for 'gy > 20 MeV	this pu
WHAT(5)		bound of t = WHAT(
WHAT(6)	= step	length in	assignin	g indices	s (Default	= 1.0)	
SDUM	= PRIMA = NOPRI		ight window	v applies als	so to primar	y particles	(default

Selecting Weight Windows - 1

BIASING	0.0	0.0	4.64	8.	18.	2. PRINT	
FLUKA out	out file:						
Hadron importa	nce RR/Splitt:	ing counters					
-	RR <wt> in -00 0.00E+00</wt>	-		<wt> in <wt> k: 0.00E+00 0.00E+0</wt></wt>	-	N. of RR <wt> in 1.15E+05 9.31E-01</wt>	
				<wt> in <wt> or 0.00E+00 0.00E+0</wt></wt>		N. of Sp <wt> in 0.00E+00 0.00E+00</wt>	
-		-		<wt> in <wt> k: 3.22E-01 1.06E-0</wt></wt>	-	N. of RR <wt> in 6.03E+03 2.16E-01</wt>	
						N. of Sp <wt> in 8.24E+04 4.65E-01</wt>	
"N. of RR" " <wt> in" "<wt> kil"</wt></wt>	(i. Rus > Ave > Ave	e., particle sian Roulett rage weight	es undergo e nor spi of these	oing Russian H litting) particles	Roulette	and which are no as well as neit ng submitted to	her
"N. of Sp" " <wt> in" "<wt> out"</wt></wt>	> Ave	rage weight	of these	particles	-	n and which are itted to splitti	-

Selecting Weight Windows - 2

where

- A = "N. of RR" + "N. of Sp" = total number of particles entering the region
- B = ("<Wt> in"_RR * "N. of RR") + ("<Wt> in"_Sp * "N. of Sp")
 = total weight of the particles entering the region
- B/A = average weight of the particles entering the region

- Note -1: RR and splitting arising from Weight-Window biasing (options WW-FACTOR, WW-THRESh, WW-PROFI) or from multiplicity biasing (WHAT(2) in option BIASING) are not accounted for in the counters.
- Note 2: Separate counters are printed for hadrons/muons, electrons/photons and low-energy neutrons (referring to importance biasing requested by BIASING, respectively, with WHAT(1) = 1.0, 2.0 and 3.0, or = 0.0 for all).

Selecting Weight Windows - 3

Strategy:

1. run without any biasing and print counter, e.g.,

BIASING	0.0	1.0	1.0	1.	9.	PRINT
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2. analyze counter and adjust region importance biasing, *e.g.*, according to the inverse of the attenuation in shielding, add other biasing, *e.g.*, leading particle biasing, run and print counter again

BIASING	0.0	1.0	1.0	1.	9.	PRINT
BIASING	0.0	1.0	1.47	4.		
BIASING	0.0	1.0	2.15	5.		
BIASING	0.0	1.0	3.16	6.		
BIASING	0.0	1.0	4.64	7.		
BIASING	0.0	1.0	4.64	8.		

3. analyze counter, select Weight Windows (WW-THRES, WW-FACTO) around average weights and perform final (high-statistics) run

Leading particle biasing - 1

- Leading particle biasing is available only for e⁺, e⁻ and photons.
- It is generally used to avoid the geometrical increase with energy of the number of particles in an electromagnetic shower
- It is characteristic of EM interactions that two particles are present in the final state (at least in the approximation made by most MC codes).
- Only one of the two is randomly retained and its weight is adjusted so as to conserve weight x probability.
- The most energetic of the two particles is kept with higher probability (as it is the one which is more efficient in propagating the shower).
- Leading particle biasing is very effective at reducing t but increases σ by introducing weight fluctuations. If its application is not limited below a suitable energy threshold, it should be backed up by weight windows.

Leading particle biasing - 2 Input card: EMF-BIAS 16. 1022. 0. 5.E-4 20. 2.LPBEMF EMF-BIAS For SDUM = LPBEMF (default): WHAT(1) > 0.0: leading particle biasing (LPB) is activated WHAT(1) = $2^{0} \times b0 + 2^{1} \times b1 + 2^{2} \times b2 + 2^{3} \times b3 + 2^{4} \times b4 + 2^{4} \times b4$ $2^5 \times b5 + 2^6 \times b6 + 2^7 \times b7$ (b0 = 1 : LPB for bremsstrahlung and pair production) b1 = 1 : LPB for bremsstrahlung b2 = 1 : LPB for pair production b3 = 1 : LPB for positron annihilation at rest b4 = 1 : LPB for Compton scattering b5 = 1 : LPB for Bhabha & Moller scattering b6 = 1 : LPB for photoelectric effect b7 = 1 : LPB for positron annihilation in flight *Note:* WHAT(1) = 1022 activates LPB for all physical effects (values larger than 1022 are converted to 1022) (FLAIR uses 254)

```
< 0.0: leading particle biasing is switched off
= 0.0: ignored
```

Leading particle biasing - 3

Input card: **EMF-BIAS**

EMF-BIAS	1022.	0.	5.E-4	16.	20.	2.LPBEMF
WHAT (2)	> 0.0:		hreshold h s and posi		h LPB is	played for
			s: kinetic ene s: total ener <u>c</u>	rgy 3y plus rest r	nass energ	у
	< 0.0:			usly define Il energies, D		nold to infinity
	= 0.0:	ignored	- F,		,	
WHAT(3)	> 0.0:	energy t	hreshold h	below which	h LPB is	played for photons
	< 0.0:			usly define Ill energies, [nold to infinity
	= 0.0:	ignored				
WHAT (4)	= lowe:	r bound o	f the regi	lon indices	s/names ((Default = 2.0)
WHAT (5)	= uppe:	r bound o	f the regi	lon indice:	s/names	(Default = WHAT(4))
WHAT(6)	= step	length i	n assignir	ng indices,	/names (D)efault = 1.0)

Multiplicity tuning - 1

Input card: **BIASING**

- Multiplicity tuning is meant to be for hadrons what Leading Particle Biasing is for electrons and photons.
- A hadronic nuclear interaction at LHC energies can end in hundreds of secondaries. Thus, to simulate a whole hadronic cascade in bulk matter may take a lot of CPU time.
- Except for the leading particle, many secondaries are of the same type and have similar energies and other characteristics.
- Therefore, it is possible to discard a predetermined average fraction of them, provided the weight of those which are kept and transported be adjusted so that the total weight is conserved *(but the leading particle is never discarded)*.
- The user can tune the average multiplicity in different regions of space by setting a region-dependent reduction factor *(in fact, it can even be > 1 ! But this possibility is seldom used).*

Multiplic	ity tun	ning -	2			Input card	BIASIN
BIASING	1.0	0.7	1.0	8.	18.	1.	
WHAT(1)	specifies	the par	ticles to	o be bias	ed with	WHAT(3) (S	see before
	= 1.0		icles s and muons ns, positror		ons		
WHAT(2)		ries pro	-	a collis		erage num ld be redu	
WHAT(3)	= (see impo	ortance bi	asing)				
WHAT(4)	= lower b	ound of	the regio	on indice	s/names	(Default = 2	.0)
WHAT(5)	= upper b	ound of	the regio	on indice	es/names	(Default = V	/HAT(4))
WHAT(6)	= step le	ngth in	assignin	g indices	s/names (Default = 1.0)
SDUM	= RRPRONL = (blank)	•	•		ary particle	es only	

Non-analog neutron absorption - 1 Input card: LOW-BIAS

- Also called survival biasing
- It is implemented in most low-energy neutron transport codes, were the user must choose between two options:
 - at each neutron collision either analog scattering or absorption according to the actual physical probability σ_s / σ_T and (1- σ_s / σ_T), respectively
 - systematic survival with a weight reduced by a factor σ_s/σ_T
- In FLUKA there is also a different alternative: the user can force the neutron absorption probability to take an arbitrary value, pre-assigned on a region-by-region basis and as a function of energy.

When and how to use it?

- A small survival probability is often assigned to thermal neutrons to limit the number of scatterings in non-absorbing media.
- It is also very useful in material with unusual scattering properties (*e.g.*, iron)
- Survival probabilities too small with respect to the physical one σ_s / σ_T may introduce large weight fluctuations due to the very different number of collisions suffered by individual neutrons: in these cases a Weight Window should be applied. (Never set the absorption probability larger than a few times the physical one !!)

Non-analog neutron absorption - 3 Input card: LOW-BIAS

LOW-BIAS	60.0	47.0	0.95	5.0	19.0	0.0	
WHAT(1) > 0.0	Neutr	ons in e	nergy gro	ups with	number >=		ad
WHAT(2) > 0.0	(Defa Neut non-	ult: dep rons in analogue	ends on E energy gr absorpti	DEFAULTS s coups >= W con. If Wi	setting) WHAT(2) ur	MGP (=260)	
WHAT(3) > 0.0			survival ends on D			$_{s}/\sigma_{tot}$)	
WHAT $(4) = low$	er bound	of the	region in	dices/nar	mes (Defau	11t = 2.0)	
WHAT(5) = upp	er bound	of the	region in	dices/nar	mes (Defau	ult = WHAT(4))	
WHAT(6) = ste	p length	in assi	gning ind	lices/name	es (Defaul	lt = 1.)	

Biasing mean free paths - 1

Decay lengths:

- The mean life/average decay length of unstable particles can be artificially shortened.
- It is also possible to increase the generation rate of decay products without the parent particle actually disappearing.
- Typically used to enhance statistics of muon or neutrino production.
- The kinematics of the decay can also be biased (decay angle).

Biasing mean free paths - 2					Input card: LAM-BIAS		
LAM-BIAS	-3.E+3	1.	1.	13.	16.	0.GDECAY	
for SDUM = WHAT(1)	: mean de					the laboratory	
	(otherwi	se it is left	unchange	d).		al decay length . random choice)	
		after cro any case	eation of t and their	he decay pr weight adju	roducts. The	ill survive or not e latter are created in into account the ratio pability.	
	> 0.0	decay po weight V	pint the p o	article alwo b). Its daug	ays survives	ed probability: at the with a reduced viven a weight W=Pu/P	

Biasing mean free paths - 3

-0.5 1. 1. 13. 16. 0. LAM-BIAS for SDUM = blank: WHAT(1) : the mean life of the particle in its rest frame is reduced by a factor |WHAT(1)| (must be <= 1.0) < 0.0 : (as for SDUM=GDECAY) Russian Roulette > 0.0 : (as for SDUM=GDECAY) the particle always survives with a reduced weight for SDUM = blank or GDECAYWHAT (2) and WHAT (3) : (see interaction length biasing) WHAT(4) = lower bound of the particle index (Default = 1.0) WHAT(5) = upper bound of the particle index (Default = WHAT(4) if WHAT(4) > 0, 46 otherwise) WHAT(6) = step length in assigning indices (Default = 1.0)

Interaction lengths:

- In a similar way, the hadron or photon mean free path for nuclear interactions can be artificially decreased by a predefined particle-or material-dependent factor.
- This option is useful for instance to increase the probability for beam interaction in a very thin target or in a material of very low density.
- It is also necessary to simulate photonuclear reactions with acceptable statistics, the photonuclear cross section being much smaller than that for EM processes.

Biasing	mean f	ree po	ths - !	5	Inpu	ıt card: LAM-I	BIA
LAM-BIAS	0.0	0.02	11.	7.	0.	0.	
WHAT (1) :	(see deca	ay length	biasing)				
WHAT (2) :	biasing	factor fo	or hadroni	ic inelas	tic inter	actions	
			tic interacti (must be <=	-	f the partio	cle is reduced by	a
	< 0. :	decides v creation o any case o	vhether the of the secor	e particle d ndaries prod eight adjust	actually wil ducts. The l ted taking i	i.e. random choi survive or not c latter are create nto account the r pility.	after d in
	> 0. :	reduced their weig	weight. The	secondarie d taking int	es are creat o account t	s survives with a ted in any case ar he ratio between	d

Biasing mean free paths - 6

LAM-BIAS	0.0	0.02	11.	7.	0.	0.	
WHAT (3)	< (= (0.0 <	bias 0.0 : rese 0.0 : igne	ing factor l ets to th ored if a v erial, other =< 2.0 :	has to be ap le default value has be wise all mat	oplied. - a prev. as en previous terials	the inelastic signed value ly assigned to a	1 specific
WHAT (4)	= lower	bound of	the part	icle indi	lces/names	s (Default = 1.0))
WHAT(5)		bound of t = WHAT(-				
WHAT(6)	= step 3	length in	assignin	g indices	s (Default :	= 1.0)	

User-written biasing

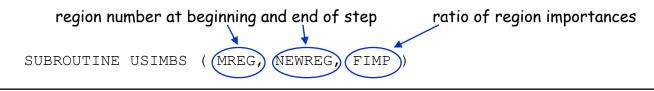
ubsset.f : User BiaSing SETting

- · called after reading in the input file and before first event
- allows to alter almost any biasing weight on a region-dependent basis

For example:	region number	region importances
SUBROUTINE UBS	SET (↓ (MPHAD, IMPLOW, IMPEMF,)
æ		NONA, PNONAN, IGDWSC, FDOWSC,
&	JWSHPP, WV	NLOW, WWHIG, WWMUL, EXPTR ,
&	ELECUT, GAN	MCUT, LPEMF, ELPEMF, PLPEMF)

usimbs.f: User defined IMportance BiaSing

- called at *every* particle step (!)
- allows to implement any importance biasing scheme based on region number and/or phase space coordinates
- enabled with BIASING/SDUM=USER



udcdrl.f: User defined DeCay DiRection biasing and Lambda

• only for neutrinos emitted in decays: bias on direction of emitted neutrino

Summary of main input cards

BIASING

1) region importance biasing (surface splitting or Russian Roulette)

2) multiplicity tuning at hadronic interactions

WW-FACTO / WW-THRES

definition of weight windows

EMF-BIAS

leading particle biasing for e+, e- and photon interactions

LOW-BIAS

non-analog neutron absorption (survival biasing)

LAM-BIAS

mean free path biasing (decay length biasing, hadronic interaction length biasing)