

| Amplitudes_2Pi | code/init/lowPhoton/twoPi_production/Amplitudes.f90 | | |
|----------------------------|---|---------|--|
| inMedium_delta_width | logical | .false. | turn of the in-medium-width of the delta |
| inMedium_delta_potential | logical | .false. | turn of the in-medium-potential of the delta |
| inMedium_nucleon_potential | logical | .false. | turn on the in-medium-potential of the nucleon |
| inMedium_pion_potential | logical | .false. | turn on the in-medium-potential of the pion |
| buuPotential | logical | .true. | use buu potentials, else constants |

| AnalyzeSpectra | code/analysis/analyzeSpectra.f90 | | |
|-----------------------|-----------------------------------|---------|---|
| realID | logical, dimen- sion(1:122) | .false. | Switch on/off the output for specific particle IDs from the real particles vector |
| pertID | logical, dimen- sion(1:122) | .false. | Switch on/off the output for specific particle IDs from the pert particles vector |

| angular_distribution | code/collisions/phaseSpace/winkelVerteilung.f90 | | |
|-------------------------------|---|--------|--|
| deltaPWave | logical | .true. | Switch for P-Wave decay of delta in pion nucleon Only relevant for deltas which are produced in pion-nucleon collisions. → see also master_2body Values: <ul style="list-style-type: none"> • .false.= isotropic in CM-Frame • .true. = $1+3*\cos(\theta)**2$ in CM Frame (θ is angle of producing pion to outgoing pion) |
| pionNucleon_backward | logical | .true. | Switch for backward peaked pion nucleon cross section: <ul style="list-style-type: none"> • .true.= use backward peaked distribution • .false.= isotropic |
| pionNucleon_backward_exponent | real | 26.5 | Exponent for backward peaked pion nucleon cross section. Distribution=(coeff-cos(theta)**exponent*(pole-sqrt(s)/pole) Only used if pionNucleon_backward=.true. . |
| pionNucleon_backward_coeff | real | 1.9 | Exponent for backward peaked pion nucleon cross section. Distribution=(coeff-cos(theta)**exponent*(pole-sqrt(s)/pole) Only used if pionNucleon_backward=.true. . |

| | | | |
|-----------------------|---------|---------|---|
| rho_pipi_nonIsotropic | logical | .true. | Switch for non-isotropic rho \rightarrow pi pi decay: <ul style="list-style-type: none"> • .false.= isotropic in CM-Frame • .true. = non-isotropic |
| NNisotropic | logical | .false. | if .true.: set isotropic nucleon-nucleon elastic cross section |
| iParam_gammaNVN | integer | 3 | for gamma N \rightarrow V N events, this parameter is given to the routine vecmesa and selects there, how dsigma/dt is calculated. Only if iParam_gammaNVN ≥ 0 the default value of that routine is overwritten. Possible values: <ul style="list-style-type: none"> • 0: 'old' parametrisation for gammaN\rightarrowVN (cf. Effenberger PhD): dsigma/dt \sim exp(Bt). Slope parameter B according ABBHHM collab, PR 175, 1669 (1968). • 1: Pythia parametrisation: Slope parameter $B=2*b_p+2*b_V+4*s**eps-4.2$ • 2: 'Donnachie, Landshoff' Select t according dsig/dt as given by VecMesWinkel/dsigdt, not by a given slope parameter • 3: as 1, but for rho and $W < \sim 6\text{GeV}$ slope parameter adjusted according CLAS experimental data [Morrow et al, EPJ A39, 5 (2009)] • 4: Muehlich PhD, Appendix E • 5: Rho0 Toy Init • 6: Rho0 Toy Init: Fit to PYTHIA-VMD • 7: Flat (not exp.) cf. VecMesWinkel/vecmesa for a detailed description. |
| NN_NR_noniso | logical | .false. | If .true., use non-isotropic angular distr. for NN \rightarrow NR, according to dsigma/dt = b/t**a. |

annihilation code/collisions/twoBodyReactions/annihilation/Annihilation.f90

| | | | |
|-------|---------|---|---|
| model | integer | 2 | Switch between the models of annihilation: <ul style="list-style-type: none"> • 1 – string based model, • 2 – statistical model |
|-------|---------|---|---|

| | | | |
|---------------|---------|---|---|
| position_flag | integer | 1 | Switch between the choices of position of outgoing mesons: |
| | | | <ul style="list-style-type: none"> • 1 – at the c.m. of the baryon and antibaryon, • 2 – at the antibaryon position |

| | | | |
|-------------------------|--|--------|---|
| barAntiBar_input | code/collisions/twoBodyReactions/baryonBaryon/barAntiBar.f90 | | |
| fact_LambdaBar | real | 1. | Enhancement factor of pbar p \rightarrow Lambda LambdaBar cross section (for larger statistics) |
| fact_JPsi | real | 1. | Enhancement factor of pbar p \rightarrow J/Psi cross section (for larger statistics) |
| fact_JPsi_width | real | 1. | Enhancement factor of the J/Psi total width (for larger statistics) |
| useAnni | logical | .true. | Flag whether to perform Baryon-Antibaryon annihilation or not at all |

| | | | |
|----------------------|---|--------------|---|
| barBar_barBar | code/collisions/twoBodyReactions/baryonBaryon/barBar_barBar.f90 | | |
| mat_NR | real, dimension(Delta:F37.1950) | ... | Squared matrix elements $M^{*2}/16\pi$ for $N N \rightarrow N R$. See http://arxiv.org/abs/1203.3557 . |
| mat_DR | real, dimension(Delta:F37.1950) | (/210., ...) | Squared matrix elements $M^{*2}/16\pi$ for $N N \rightarrow \Delta R$. See http://arxiv.org/abs/1203.3557 . |
| icugnon | integer | 1 | Switch for nucleon nucleon \rightarrow nucleon nucleon cross sections: <ul style="list-style-type: none"> • 0=old parametrization • 1=new parametrization (Alexej Larionov, Cugnon) |
| use_ND_ND_model | logical | .false. | Switch for delta nucleon \rightarrow delta nucleon cross sections: <ul style="list-style-type: none"> • false=old parametrization • true =one pion exchange model (Effenberger, Buss) |
| new_NR_NR | logical | .true. | <ul style="list-style-type: none"> • .false.= Switch off the NR\rightarrow NR improvement (improvement= better NN\leftrightarrowNN fit is being used) • only for debugging or comparing |
| NR_NR_massSHIFT | logical | .false. | <ul style="list-style-type: none"> • .true.= Shift the srts in NR\rightarrow NR elastic collisions. |
| oldOset_treatment | logical | .false. | <ul style="list-style-type: none"> • .true.= Use the old treatment for the Oset Delta width: Put everything into 3-body. • only for debugging or comparing |

| | | | |
|--------------------------|---------|---------|---|
| etafac | real | 6.5 | Parameter for enhancement of $p n \rightarrow N^*(1535) N$, relative to $p p \rightarrow N^*(1535) N$, in order to enhance eta production in pn collisions. See Calen et al., PRC 58 (1998) 2667. |
| rhofac | real | 1. | Parameter for enhancement of $p n \rightarrow N^*(1520) N$, relative to $p p \rightarrow N^*(1520) N$, in order to enhance rho production in p n collisions. |
| neufac | real | 1. | Parameter for enhancement of $p n \rightarrow N R$, relative to $p p \rightarrow N R$, affecting all resonances. |
| neufac_ropen | real | 2. | Parameter for enhancement of $p n \rightarrow N N^*(1440)$, relative to $p p \rightarrow N N^*(1440)$. See http://arxiv.org/abs/1203.3557 . |
| deltaN_densityDependence | logical | .false. | Switch to turn on the density dependence of the $NN \leftrightarrow N \Delta$ cross section. We use the density dependence from: Song/Ko, arXiv:1403.7363. The strength of the in-medium modification is controlled by the parameter alpha. |
| alpha | real | 1.2 | Parameter which controls the density dependence of the $NN \leftrightarrow N \Delta$ cross section. We use the density dependence from: Song/Ko, arXiv:1403.7363. See also deltaN_densityDependence. |

barBar_barBarMes `code/collisions/twoBodyReactions/baryonBaryon/barBar_barBarMes.f90`

| | | | |
|--------------|---------|--------|---|
| NNpi_BG | integer | 2 | Switch for the $N N \rightarrow N N \pi$ background: <ul style="list-style-type: none"> • 0 = no BG • 1 = BG according to Teis • 2 = BG according to Buss (improves threshold behavior, default) • 3 = BG according to Weil |
| NNV_BG | logical | .true. | Include a $N N \rightarrow N N V$ background term, where $V=\omega,\phi$ (in addition to possible resonance contributions). |
| isofac_omega | real | 1. | Isospin enhancement factor for $p n \rightarrow p n \omega$, relative to $p p \rightarrow p p \omega$. Data indicate that this is around 2, while theory predicts even larger values (up to 5). Reference: Barsov et al., EPJ A21 (2004) 521-527. |
| isofac_phi | real | 1. | Isospin enhancement factor for $p n \rightarrow p n \phi$, relative to $p p \rightarrow p p \phi$. Theory predicts values of 3-4, cf.: Kaptari, Kaempfer, Eur.Phys.J. A23 (2005) 291-304. |

| BarBar_to_barBar_model | code/collisions/twoBodyReactions/baryonBaryon/barbar_to_barbar_model.f90 | | |
|-------------------------------|--|-----|---|
| couplings_switch | integer | 2 | Possible values: <ul style="list-style-type: none"> • 1 = use couplings according to Dmitriev • 2 = use couplings according to Pascalutsa (default) |
| lambda_cutoff | real | 0.6 | Cutoff parameter in the form factor for ND→ND Possible values: <ul style="list-style-type: none"> • 0.6 (Dmitriev, default) • 1.2 (Doenges) |

baryonPotentialcode/potential/baryonPotential.f90

EQS_Type

integer

5

Switch for equation of state for nucleon resonances with spin 1/2.
Parameters for nucleon potentials:

- 0 = nucleon potential is set to zero
- 1 = soft, momentum dependent, $\lambda = 2.130$ (Teis PhD, $K = 215$ MeV)
- 2 = hard, momentum dependent, $\lambda = 2.126$ (Teis PhD, $K = 380$ MeV)
- 3 = soft, momentum independent (Teis PhD, $K = 215$ MeV)
- 4 = hard, momentum independent (Teis PhD, $K = 380$ MeV)
- 5 = medium, momentum dependent, $\lambda = 2.130$ (Teis PhD, $K = 290$ MeV)
- 6 = LDA potential (Birger Steinmueller)
- 7 = Deuterium potential Argonne V18 (not usable for eventtypes 'heavyIon' and 'hadron')
- 8 = LDA Potential Welke
- 9 = Buss PhD, Set#1 ($K = 220$ MeV, momentum dependent)
- 10 = Buss PhD, Set#2 ($K = 220$ MeV, momentum dependent)
- 11 = Buss PhD, Set#3 ($K = 220$ MeV, momentum dependent)
- 12 = Shanghai meeting 2014 (soft, momentum independent, $K = 240$ MeV)
- 13 = slightly modified Cooper potential, central depth = - 67.5 MeV at $p=0$ (see #14)
- 14 = Potential fitted by Cooper et al, Fig. 6 in PRC 47 (1993) 297
- 98 = use pre-stored values
- 99 = variable Skyrme : E_bind, p_0, U_0, rho_0 must be defined!

NOTES

References:

- for 1-5, see the PhD thesis of S. Teis, chapter 3.3.2 / table 3.1
 - for 9-11, see the PhD thesis of O. Buss, chapter 7.2.3 / table 7.1
-

| | | | |
|-------------------------|---------|---------|---|
| DeltaPot | integer | 1 | <p>Switch for potential of spin=3/2 resonances:</p> <ul style="list-style-type: none"> • 0 = no potential • 1 = nucleon (spin=1/2) potential times 2/3 [according to Ericson/Weise book] • 2 = 100 MeV * rho/rhoNull • 3 = nucleon (spin=1/2) potential |
| HypPot | integer | 1 | <p>Switch for potential of hyperons:</p> <ul style="list-style-type: none"> • 0 = no potential • 1 = nucleon (spin=1/2) potential times (3+S)/3 (i.e. according to the share of the light quarks) • 2 = nucleon (spin=1/2) potential |
| symmetryPotFlag | integer | 0 | <p>Switch for the asymmetry term in the nucleon potential. NOTES Possible values:</p> <ul style="list-style-type: none"> • 0 = none (default) • 1 = linear (strength given by 'dsymm') • 2 = stiffer, $E_{sym}=E_{sym_rho_0} * U^{\gamma}=31. * U^{\gamma}$, $\gamma=2$ • 3 = stiff, linear increasing $E_{sym}=E_{sym_rho_0} * U=31. * U$ • 4 = soft, $U_c=3$, can give negative $E_{sym}=E_{sym_rho_0} * U * (U_c - U) / (U_c - 1)$ |
| symmetryPotFlag_Delta | logical | .false. | <p>Switch for the asymmetry term in the Delta potential. NOTES If .true., a symmetry potential will be used also for the Delta (but only if symmetryPotFlag>0). It is closely related to the symmetry potential of the nucleon.</p> |
| noPerturbativePotential | logical | .false. | <p>Switch for potential of perturbative particles. If .true. then perturbative baryons feel no potential.</p> |
| rho_0 | real | 0.16 | <p>Nuclear matter density for EQS_Type=99 NOTES</p> <ul style="list-style-type: none"> • Units : fm⁻³ |

| | | | |
|-----------------|---------|---------|--|
| p_0 | real | 0.8 | momentum for which $U(p_0, \rho = \rho_0) = 0$ for EQS_Type=99 NOTES • Units : GeV |
| U_0 | real | 0.075 | $U(p=0, \rho = \rho_0)$ for EQS_Type=99 NOTES • Units : GeV |
| bindingEnergy | real | 0.016 | Nuclear matter binding energy for EQS_Type=99 NOTES • Units : GeV |
| compressibility | real | 0.290 | Nuclear matter compressibility for EQS_Type=99 NOTES • Units : GeV |
| SurfacePotFlag | logical | .false. | Switch for the surface term in the nucleon potential. NOTES • Do not use it together with yukawa! |
| nLoopReAdjust | integer | 10 | number of iterations, if density is readjusted (cf. <code>type(nucleus)%ReAdjustForConstBinding</code>) NOTES It is necessary to reiterate (at least for momentum dependent potentials), since we calculate the potential for a given pF and then calculate for the readjusting a new pF. |
| dsymm | real | 0.03 | Parameter for symmetry potential in GeV. NOTES Value is only used for <code>symmetryPotFlag = 1</code> |

BaryonWidth

| | code/width/baryonWidth.f90 | | |
|------------|----------------------------|---------|---|
| readTable | logical | .true. | There is a tabulation of the widths saved in <code>buinput</code> which is used to initialize (' <code>baryonWidthVacuum.dat.bz2</code> '). If you don't want to use this pre-tabulated input, then you can set " <code>readTable=.false</code> ". This is useful for runs on a cluster where you want to minimize input/output. Also it is necessary if the decay channels have been modified (cf. <code>DecayChannels.dat</code>). |
| writeTable | logical | .false. | This flag determines whether we write out a new tabulation of the widths (' <code>baryonWidthVacuum.dat.bz2</code> '). It will only have an effects if <code>readTable == .false.</code> or reading of the tabulation file fails for some reason. |

| BaryonWidthMedium_tables code/width/baryonWidthMedium_tables.f90 | | | |
|---|---------|---------|--|
| minRes | integer | -1000 | Read the data table starting at this minimal resonance ID. ONLY FOR TESTING!!! |
| maxRes | integer | 1000 | Read the data table up to a maximum resonance ID. ONLY FOR TESTING!!! |
| inMediumParameterset | integer | 2 | chooses the parameters for the inMediumWidth (1 electron, 2 neutrino) |
| onlyNucleon | logical | .false. | Only for debugging: only nucleon width is read in. |
| deltaOset | logical | .false. | Use delta width according to Oset et al. NPA 468 (1987) |
| extrapolateAbsP | logical | .false. | if(true) then set absP to maxAbsP if absP is larger |

| BaryonWidthVacuum code/width/baryonWidthVacuum.f90 | | | |
|---|---------|--------|---|
| use_cutoff | logical | .true. | <ul style="list-style-type: none"> • Switch on and off the use of cut off parameters. • These cut-offs are necessary when working with dispersion relations to deduce the real part. |
| deltaRho_cutoff | real | 0.85 | <ul style="list-style-type: none"> • Cut off parameter for the decay of a resonance into delta rho. • Units of GeV |
| baryon_cutoff | real | 2.0 | <ul style="list-style-type: none"> • Cut off parameter for the decay of a resonance into an unstable baryon and a meson. • Units of GeV |
| meson_cutoff | real | 1.6 | <ul style="list-style-type: none"> • Cut off parameter for the decay of a resonance into a baryon and an unstable meson. • Units of GeV |
| Delta_width | integer | 1 | <p>Select a parametrization for the Delta width:</p> <ul style="list-style-type: none"> • 1 = Manley (GiBUU default, cf. Manley/Saleski, Phys. Rev. D 45, 1992) • 2 = Dmitriev (Dmitriev/Sushkov/Gaarde, Nucl. Phys. A 459, 1986) • 3 = Moniz (Koch/Moniz/Ohtsuka, Ann. of Phys. 154, 1984) • 4 = Verwest (Phys. Lett. B 83, 1979) • 5 = UrQMD (Bass et al., Prog. Part. Nucl. Phys. 41, 1998) |

| BB_BYK code/collisions/twoBodyReactions/baryonBaryon/barBar_BarHypKaon.f90 | | | |
|---|------------------------|--------|---|
| enable | logical | .true. | Enable the production of BB \rightarrow B Hyperon Kaon channels. B=Nucleon ^{0,1} ,Delta ^{-,0,+} ,++; Hyperon=Lambda ⁰ ,Sigma ^{0,-,+} ; Kaon=K ^{+,0} |
| parameter_set | integer | 2 | Select a particular parameter set for BB \rightarrow BYK collisions. Possible values: <ul style="list-style-type: none"> • 1 = original Tsushima model: Tsushima et al., PRC59 (1999) 369 • 2 = extended/adjusted model, fitted to HADES data: Agakishiev et al., arXiv:1404.7011 • 3 = custom parameters based on Tsushima values (as given by the array 'a' in the jobcard; those values not given in the jobcard are adopted from Tsushima, i.e. parameter set 1) • 4 = custom parameters based on HADES values (as given by the array 'a' in the jobcard; those values not given in the jobcard are adopted from HADES, i.e. parameter set 2) |
| a | real, dimension(1:Nch) | -1. | This array contains the "a" parameters (in microbarn) for the 30 primary channels, see: <ul style="list-style-type: none"> • Tsushima et al., PRC59 (1999) 369, table III • Agakishiev et al., arXiv:1404.7011, chapter IV <p>Note: The values given in the jobcard are only used for parameter_set = 3 and 4.</p> |

| Box code/init/initBox.f90 | | | |
|----------------------------------|------------------------|---------|---|
| thermalInit | logical | .false. | flag how to initialize |
| nDens | real | 1.0 | particle density [fm ⁻³] |
| ChargeSelection | integer | 0 | define the type of the charge selection: <ul style="list-style-type: none"> • 0: only pi0 • 1: 50% pi+, 50% pi- • 2: 33% for +,0,- |
| pInit | real | 0.5 | initial momentum of particles [GeV/c] |
| BoostZ | real | 0.0 | additional boost for all particles in z-direction |
| Temp | real, dimension(1:122) | 0.0 | for thermal init: temperature of every meson species in GeV, if larger than 0. otherwise this species is not initialized |

| | | | |
|------------------|------------------------|-----|--|
| Fugacity | real, dimension(1:122) | 1.0 | for thermal init: fugacity of every hadron species. |
| correctMovingBox | integer | 1 | switch to indicate, whether a correction of the momenta after initialization should be done to enforce vanishing 3-momenta. possibilities are: <ul style="list-style-type: none"> • 0 : no correction • 1 : global correction • 2 : per ensemble correction |

| | | | |
|--------------------|-------------------------------|---------|---|
| BoxAnalysis | code/analysis/BoxAnalysis.f90 | | |
| do_Tmunu | logical | .false. | Switch for Tmunu output. default: Only one file for all ensemble! you may change this with the flag perEnsemble_Tmunu |
| perEnsemble_Tmunu | logical | .false. | Switch for Tmunu output. One file per ensemble! NOTES this may slow down the execution dramatically, since huge output to the hard drive is induced. You may observe this, if e.g the cpu load drops permanently to 30%. Thus: switch it on, only if you want it! |
| do_P | logical | .false. | Switch for dN/p ² dp output |
| do_velrel | logical | .false. | Switch for calculating velrel |
| do_Cumulants | logical | .false. | Switch for calculating cumulants |
| useSet | | | |

| | | | |
|-------------------------|---------------------|---------|--|
| Checks | code/run/checks.f90 | | |
| Do_CheckDensity | logical | .false. | Flag to indicate whether the density check routine should be called. |
| Do_CheckCoulomb | logical | .false. | Flag to indicate whether the Coulomb check routine should be called. |
| Do_CheckFermiSurface | logical | .false. | Flag to indicate whether the Fermi-surface check routine should be called. |
| Do_CheckRadius | logical | .false. | Flag to indicate whether the radius check routine should be called. |
| Do_CheckMomentumDensity | logical | .false. | Flag to indicate whether the momentum-density check routine should be called. |
| Do_CheckEnergyLDA | logical | .false. | Flag to indicate whether the local density approximation check routine should be called. |
| Do_OccupiedReal | logical | .false. | Flag to indicate whether the occupation check routine for the real particle vector should be called. |

| | | | |
|----------------------|---------|---------|---|
| Do_OccupiedPert | logical | .false. | Flag to indicate whether the occupation check routine for the perturbative particle vector should be called. |
| Do_CheckEnergy | logical | .false. | Flag to indicate whether the energy check routine should be called. |
| Do_TachyonsReal | logical | .false. | Flag to indicate whether the tachyon check routine for the real particle vector should be called. |
| Do_TachyonsPert | logical | .false. | Flag to indicate whether the tachyon check routine for the perturbative particle vector should be called. |
| TachyonIsBlocking | logical | .false. | Select whether the occurrence of a 'tachyon' in the check routines will stop the code or not (error messages will hopefully occur later in the code). |
| Do_CheckPertFlag | logical | .true. | Flag to indicate whether the flag '%perturbative' is set correctly in the particle vectors |
| Do_CheckConservation | logical | .false. | Flag to indicate whether conservation of energy/momentum, baryon number and strangeness between time steps for the real particles should be checked |

coll_BaB code/collisions/twoBodyReactions/HiEnergy/DoColl_BaB.f90

| | | | |
|------|---------|---|--|
| iset | integer | 1 | Switch to choose an initialization of jets: <ul style="list-style-type: none"> • 1: phase space distribution, also the charge is conserved (new prescription) • 2: first jet along inPart(1) momentum, 3-d jet opposite, others orthogonal, charge is not conserved (old prescription) |
|------|---------|---|--|

coll_Manni code/collisions/twoBodyReactions/HiEnergy/DoColl_Manni.f90

| | | | |
|-----------------|---------|----|---|
| angDistribution | integer | 2 | Switch to select the angular distribution: <ul style="list-style-type: none"> • 1: isotropic • 2: diquark/quark aligned like baryon/meson |
| itry_max | integer | 10 | maximum number of tries |

collCriteria code/collisions/twoBodyReactions/collisionCriteria.f90

| | | | |
|------------------|---------|---------|---|
| kodama_evalFrame | logical | .false. | Set to .true., this logical will cause the kodama criterion to be evaluated in the laboratory/evaluation frame, not CM frame. |
|------------------|---------|---------|---|

| collHistory code/storage/CollHistory.f90 | | | |
|---|---------|---------|--|
| DoCollHistory | logical | .false. | Flag to switch on/off the whole Collision History machinery. You may set this variable via your jobcard, namelist "collHistory" |

| collisionterm code/collisions/collisionTerm.f90 | | | |
|--|---------|---------|--|
| oneBodyProcesses | logical | .true. | Switch on/off one-body-induced processes. |
| twoBodyProcesses | logical | .true. | Switch on/off two-body-induced processes. |
| threeBodyProcesses | logical | .true. | Switch on/off three-body-induced processes. |
| threeMesonProcesses | logical | .false. | Switch on/off three-meson-induced processes. These are the backreactions for e.g. $\omega \rightarrow \pi \pi \pi$ etc. |
| threeBarMesProcesses | logical | .false. | Switch on/off baryon-meson-meson induced processes. These are the backreactions for e.g. $N \pi \rightarrow N \pi \pi$ etc. |
| twoPlusOneBodyProcesses | logical | .false. | Switch on/off 2+1 body processes (two really colliding particles plus one nearby). |
| twoBodyProcessesRealReal | logical | .true. | Switch on/off two-body-induced processes between two real particles. |
| twoBodyProcessesRealPert | logical | .true. | Switch on/off two-body-induced processes between a real and a perturbative particle. |
| oneBodyAdditional | logical | .true. | Switch on/off additional Pythia one-body-induced processes. |
| doForceDecay | logical | .true. | switch on/off the forced decays at the end of the simulation |
| NOTES | | | |
| <ul style="list-style-type: none"> • Do not touch, unless you know what you are doing! • You may set this to .false., if you are e.g. running box calculations with excited states. Decaying all these particles would need a much larger particle vector... | | | |
| energyCheck | real | 0.01 | Precision of energy check for each collision in GeV. |
| maxOut | integer | 100 | Maximal number of produced particles in one process. |
| collisionProtocol | logical | .false. | Write a protocol of all real-real collisions to the file 'fort.990'. Includes the time, IDs, charges, invariant masses and 3-momenta of both collision partners and an indicator for Pauli blocking. |
| printPositions | logical | .false. | Switch on/off output of positions in real-pert collisions. Produces statistical output. |
| useStatistics | logical | .false. | Generate statistical information using the module statistics. |

| | | | |
|-----------------------|---------|---------|---|
| noNucNuc | logical | .false. | Switch on/off perturbative NN reactions. |
| storeRho0Info | logical | .false. | Flag whether in a rho0 decay the particle numbers of the resulting charged pions are stored or not. |
| storeRho0InfoOnlyDifr | logical | .false. | Flag, whether the flag storeRho0Info is valid for all decays or only for rho0, which are marked to be diffractive. |
| DoJustAbsorptive | logical | .false. | If this flag is true, then: for perturbative simulations all final state particles in a collision are set to zero; for real simulations %event index of incoming hadron is changed in the case of collision, but actual collision is not simulated. This is a way to mimick Glauber like calculations. NOTES The "absorption" is done with sigmaTot, not just by sigmaInEl. |
| annihilate | logical | .false. | If this flag is true, then an annihilation of the antibaryons with the closest baryons will be simulated (by hand) starting from annihilationTime. |
| annihilationTime | real | 1000. | Time moment (in fm/c) when the annihilation will be started. NOTES This flag has an influence only when annihilate = .true. Before annihilationTime all the collision processes are not activated. They start to act (if the corresponding switches oneBodyProcesses,twoBodyProcesses etc. are .true.) only after annihilationTime. |
| justDeleteDelta | logical | .false. | Deletes final-state products in Delta N N → NNN and Delta N → N N. NOTE: Only for testing and comparing with the old Effenberger code! DO NOT USE OTHERWISE: Violates energy conservation! This switch is meant to simulate the treatment of the Delta in the old code. Only implemented for perturbative runs. |
| noRecollisions | logical | .false. | Outgoing particles of collisions are inserted somewhere in the particle vector. Due to implementation issues, these outgoing particles may interact during the same timestep. Setting this flag to true, the parameter '%lastCollisionTime' is checked against the actual time variable and collisions of these particles are excluded. |

| | | | |
|---------------------|---------------------------------------|---------|---|
| collReporter | code/collisions/collisionReporter.f90 | | |
| UseCollReporter | logical | .false. | Enable or disable the collision reporter. |
| cR_sizeT | integer | 200 | Number of timestep bins. |
| cR_sizeE | integer | 100 | Number of sqrt(s) bins. |
| cR_DeltaT | real | 0.1 | Size of timestep bins. |

| | | | |
|-----------|------|-----|-----------------------|
| cR_DeltaE | real | 0.1 | Size of sqrt(s) bins. |
|-----------|------|-----|-----------------------|

| | | | |
|---------------------|--|---------|---|
| ColStat | code/collisions/twoBodyReactions/twoBodyStatistics.f90 | | |
| flag_sqrts | logical | .false. | If .true., then the calculation and output of the sqrts distributions from subroutine sqrts_distribution will be done |
| flag_rate | logical | .false. | If .true., then the calculation and output of the collision rates from subroutine rate will be done |
| flag_varirate | logical | .false. | If .true., then the calculation and output of the collision rates from subroutine varirate will be done |
| sqrts_mode | integer | 1 | This flag determines the way how sqrt(s) is calculated (if flag_sqrts = .true.). 1 = use vacuum sqrt(s) 2 = use in-medium, i.e. full sqrts |
| varirate_chargeZero | logical | .true. | If .true., then all charge states are combined together |
| varirate_size | integer | 100 | size of array to hold all rates |
| varirate_filterPhi | logical | .false. | If .true., then only channels involving a phi meson are reported |

| | | | |
|-----------------|------------------------------------|---------|--|
| coulomb | code/potential/coulomb/coulomb.f90 | | |
| coulombFlag | logical | .false. | Switch to turn on/off the Coulomb potential. If turned on, also 'symmetryPot-Flag' (in namelist 'baryonPotential') needs to be turned on. |
| magnetFieldFlag | logical | .false. | Switch to turn on/off elm. vector potential. NOTES The vector potential is not yet fully implemented! Please do not use this option. |

| | | | |
|--|---------|---------|--|
| cutMomentumPotential | real | 0.025 | <p>If larger than 0, the coulomb potential is set to zero for all particles with momentum larger than $\text{minmass}^2/(2*\text{cutMomentumPotential})$ The cut-off is smeared out, if $\text{cutMomentumWidth}>0$</p> <p>This cut is needed in non-RMF mode since</p> <p>$\text{pre}_i m_{\text{eff}}^2 = (\text{sqrt}(m^2+p^2)+U_C)^2-p^2$ can become smaller than zero for</p> <p>$\text{pre}_i p > U_C/2 - m^2/(2*U_C)$. In this case we have a 'tachyon'.</p> <p>NOTES</p> <ul style="list-style-type: none"> • for RMF mode you do not need this cut • This value should correspond to the maximum vale of the Coulomb potential. Therefore you should readjust this for every nucleus. • For the pion, we take the mass (0.138) instead of minmass, since here minmass is zero! |
| cutMomentumWidth | real | 0.100 | <p>If $\text{cutMomentumPotential}>0$ and $\text{cutMomentumWidth}>0$, then the cut-off is smeared by some linear interpolation:</p> <p>$\text{pre}_i (A_t-\text{Width}) = 1.0 \dots (A_t+\text{Width}) = 0.0$ with $A_t = \text{minmass}^2/(2*\text{cutMomentumPotential})$ and the width given here in GeV.</p> <p>This width is introduced in order to destroy numerical problems due to sharp edges.</p> |
| <hr/> | | | |
| DecayChannels code/database/decayChannels.f90 | | | |
| rhoDelta_is_sigmaDelta | logical | .false. | <p>If true, the rho-Delta decay channel will be replaced by sigma-Delta. For discussion, see e.g. Effenberger PhD, chapter 6.3.2.</p> |
| <hr/> | | | |
| deltawidth code/width/deltaWidth.f90 | | | |
| deltaSwitch | integer | 3 | <p>Switch for different prescriptions for the delta width.</p> <p>NOTES</p> <ul style="list-style-type: none"> • 1 = use Oset self energies+BUU input • 2 = Spreading potential • 3 = use Oset self energies • 4 = density dependent with BUU input |

| detailed_diff | | | |
|-----------------------|------------------------------------|---------|--|
| | code/analysis/neutrinoAnalysis.f90 | | |
| EkinMin | real | 0. | if detailed_diff_output is TRUE: Maximal kinetic energy for dsigma/dEkin for hadrons |
| EkinMax | real | 2. | if detailed_diff_output is TRUE: Maximal kinetic energy for dsigma/dEkin for hadrons |
| dEkin | real | 0.01 | if detailed_diff_output is TRUE: Delta(eKin) for dsigma/dEkin for hadrons |
| forPion | logical | .true. | If .true. then also the detailed output of differential cross sections is produced |
| forEta | logical | .false. | If .true. then also the detailed output of differential cross sections is produced |
| forKaon | logical | .false. | If .true. then also the detailed output of differential cross sections is produced |
| forKaonBar | logical | .false. | If .true. then also the detailed output of differential cross sections is produced |
| forDmeson | logical | .false. | If .true. then also the detailed output of differential cross sections is produced |
| forDbar | logical | .false. | If .true. then also the detailed output of differential cross sections is produced |
| forDs_plus | logical | .false. | If .true. then also the detailed output of differential cross sections is produced |
| forDs_minus | logical | .false. | If .true. then also the detailed output of differential cross sections is produced |
| forNucleon | logical | .true. | If .true. then also the detailed output of differential cross sections is produced |
| forLambda | logical | .false. | If .true. then also the detailed output of differential cross sections is produced |
| forSigmaResonance | logical | .false. | If .true. then also the detailed output of differential cross sections is produced |
| forXi | logical | .false. | If .true. then also the detailed output of differential cross sections is produced |
| forOmegaResonance | logical | .false. | If .true. then also the detailed output of differential cross sections is produced |
| deuteriumFermi | | | |
| | code/init/deuterium.f90 | | |
| waveFunction_switch | integer | 1 | Possible values are: <ul style="list-style-type: none"> • 0 – No Wave functions! Pointlike Deuterium • 1 – Wave functions according to Bonn potential • 2 – Wave functions according to Argonne V18 |

| | | | |
|---------------|---------|-----|--|
| iParam | integer | 1 | Choose parameterization of momentum distribution when using the Bonn potential. Possible values: <ul style="list-style-type: none"> • 1 – Full Bonn (MaH87) • 2 – OBEPQ (MaH87) • 3 – OBEPQ-A (Mac89) • 4 – OBEPQ-B (Mac89) • 5 – OBEPQ-! (Mac89) • 6 – OBEPR (MaH87) self-made • 7 – Paris References: MaH87: R. Machleidt et al. Phys. Rep. 149, 1 (1987) Mac89: R. Machleidt, Advances in Nucl. Phys. Vol 19 |
| pMax | real | 0.5 | Cut-off parameter for Fermi momentum |
| scaleMomentum | real | 1.0 | The selected momentum is multiplied by this factor afterwards, i.e. some rescaling is done |

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|--|---------|---------|--|
| DileptonAnalysis code/analysis/DileptonAnalysis.f90 | | | |
| Enable | logical | .false. | If .true. the dilepton analysis will be performed, otherwise not. |
| Extra | logical | .false. | If .true. an extended analysis will be performed, writing out many extra histograms (beyond the basic ones: mass, pT and rapidity). |
| DeltaDalitz | integer | 2 | Choose between different parametrizations of the Delta Dalitz decay width ($\Delta \rightarrow N e^+e^-$): <ul style="list-style-type: none"> • 1 = Wolf, http://inspirehep.net/record/306273 • 2 = Krivoruchenko (default), http://inspirehep.net/record/555421 • 3 = HadronTensor • 4 = Ernst, http://inspirehep.net/record/452782 |

| | | | |
|---------------|---------|-------|---|
| DeltaDalitzFF | integer | 1 | <p>Choose a parametrization of the electromagnetic N-Delta transition form factor for the Delta Dalitz decay (only used for DeltaDalitz = 2):</p> <ul style="list-style-type: none"> • 1 = constant (default) • 2 = Dipole • 3 = MAID 2005 • 4 = simple VMD • 5 = Wan/Iachello, Int. J. Mod. Phys. A 20 (2005) 1846, http://inspirehep.net/record/689265 • 6 = Ramalho/Pena, Phys.Rev. D85 (2012) 113014, http://inspirehep.net/record/1114321 |
| omegaDalitzFF | integer | 1 | <p>Choose between different parametrizations of the omega Dalitz decay ($\omega \rightarrow \pi^0 e^+e^-$) form factor:</p> <ul style="list-style-type: none"> • 0 = constant • 1 = Effenberger/Bratkovskaya (default) • 2 = standard VMD • 3 = Terschluesen/Leupold |
| b_pi | real | 5.5 | <p>This constant represents the b parameter in the form factor of the pi0 Dalitz decay (in GeV^{-2}), cf. Effenberger Diss. eq. (2.141). Originally taken from L.G. Landsberg, Phys. Rep. 128, 301 (1985).</p> |
| lambda_eta | real | 0.716 | <p>This constant represents the Lambda parameter in the form factor of the eta Dalitz decay in GeV. Values:</p> <ul style="list-style-type: none"> • L.G. Landsberg, Phys. Rep. 128, 301 (1985): Lambda = 720 MeV • HADES pp@2.2, B. Spruck, Diss. (2008): Lambda = 676 MeV • NA60, Arnaldi et al, PLB 677 (2009): Lambda = 716 MeV (default) • CB/TAPS, Berghuser et al, PLB 701 (2011): Lambda = 722 MeV |

| | | | |
|------------------|---------|---------|--|
| etaPrimeDalitzFF | integer | 0 | Choose between different parametrizations of the eta' Dalitz decay ($\eta' \rightarrow e^+e^- \gamma$) form factor: <ul style="list-style-type: none"> • 0 = constant (default) • 1 = eta FF (cf. lambda_eta) • 2 = generic VMD • 3 = Genesis / Lepton-G • 4 = standard VMD (Terschluesen) |
| angDist | integer | 1 | This switch determines the angular distribution of the pseudoscalar Dalitz decays $P \rightarrow e^+ e^- \gamma$ (with $P=\pi^0, \eta, \eta'$): <ul style="list-style-type: none"> • 0 = isotropic decay • 1 = anisotropic decay according to $1 + B \cos^2(\theta)$ with $B=1$ • 2 = the Dalitz decays of π^0 and η will be done via Pythia. |
| brems | integer | 1 | This switch determines how the bremsstrahlung contribution is obtained: <ul style="list-style-type: none"> • 0 = none • 1 = soft-photon approximation (SPA) • 2 = according to the one-boson-exchange (OBE) model by R. Shyam (for NN bremsstrahlung only, no em. form factors) • 3 = as 2, but with pion em. form factor (for pn) |
| nEvent | integer | 10 | Number of events to generate for each dilepton decay (to enhance statistics). |
| nEvent_BH | integer | 1000 | Number of events for Bethe-Heitler simulation. BH typically needs a lot more statistics than the other dilepton channels. Therefore nEvent_BH should be much bigger than nEvent. |
| kp_cut | logical | .false. | Perform a cut on (k^*p) in the dilepton analysis, where k is the photon 4-momentum, and p is the electron or positron 4-momentum. This is useful for suppressing the BH contribution. Cf. "kp_min". |
| kp_min | real | 0.01 | If $kp_cut=.true.$ a cut on (k^*p) is performed. kp_min determines the position of this cut. Only events with $(k^*p) > kp_min$ are taken into account. |
| binsz | real | 0.01 | This determines the bin size of the dilepton mass spectrum in GeV. Default is 10 MeV. |

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|-----------------|---------------------|----|--|
| filter | integer | 0 | <p>If filter is nonzero, a filtering algorithm will be applied to the dilepton pairs, otherwise they will be written to the histograms unfiltered. For details on the filtering parameters see routine 'CS'. Choices:</p> <ul style="list-style-type: none"> • 0 = no filter • 1 = DLS • 2 = HADES (simple cuts on polar angle, absolute momentum and opening angle) • 3 = HADES (full acceptance filter, using pair acceptance) • 4 = HADES (full acceptance filter, using single-particle acceptance) • 5 = g7/CLAS @ JLab • 6 = KEK E325 (cuts on rapidity, transverse momentum and opening angle) • 7 = JPARC E16 <p>NOTES For filtering modes 3 and 4, the file containing the acceptance matrices must be specified (cf. <code>hadesFilterFile</code>).</p> |
| hadesFilterFile | character(len=1000) | | This character string determines the location of the file containing the HADES acceptance matrices (filename with absolute or relative path). It has to be set for filtering modes 3 and 4. |
| WriteEvents | integer | 0 | <p>This switch decides whether we write out the simulated events. Possible values:</p> <ul style="list-style-type: none"> • 0: Don't write events (default). • 1: We write out only the lepton pair information (including charge, four-momentum, perturbative weight, source channel and filter result). All of this will be written to a file called 'Dilepton_Events.dat'. • 2: As 1, but only writing exclusive events ($NN \rightarrow NNe+e-$). • 3: We write out all produced particles in the event (including the lepton pair) to a file called 'Dilepton_FullEvents.dat'. • 4: As 3, but only writing exclusive events ($NN \rightarrow NNe+e-$). • 5: As 4, but only writing out $R \rightarrow Ne+e-$ events (with $R=N^*, \Delta^*$). |
| p_lep_min | real | 0. | This switch sets a lower bound on the lepton momentum. Only leptons with momenta larger than this threshold will pass the filter. This switch is only used for filtering mode 5 (JLab). |

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|-----------------|----------------------|----------------------------------|--|
| beta_gamma_cut | real | 1.25 | This is an upper bound on the beta*gamma value of the lepton pair. Since beta*gamma = p/m, it cuts on slow pairs. |
| massBinning | real, dimension(1:4) | (/ 0.150, 0.550, 9.999, 9.999 /) | We produce several histograms (e.g. p,pT,mT,y,theta_cm) not only for the full mass range, but also for (up to 5) different mass bins. The borders of these bins are given by this array. |
| particle_source | logical | .true. | This switch determines whether the mass spectrum will contain separate channels for different sources of particles, such as decays ($R \rightarrow \rho N$) or collisions ($\pi \pi \rightarrow \rho$, $K K \rightarrow \phi$). Currently this is only done for the rho and phi mesons. Note: If using this switch, the "sum" channel in the mass histogram should not be used, since the rho and phi contributions will enter twice. |

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|-------------------|------------------------------|---|--|
| elementary | code/init/initElementary.f90 | | |
| impactParameter | real | -1. | <ul style="list-style-type: none"> • ≥ 0: this is the actual impact parameter • < 0 : Impact parameter integration up to an automatically determined b_max. The actual impact parameter is randomly sampled in the interval $[0, b_max]$ with a proper geometrical weight. |
| particleId | integer, dimension(2) | (/1,1/) | Id of particles |
| particleAnti | logical, dimension(2) | (/.false.,.false./)if .true. then particles are antiparticles | |
| particleCharge | integer, dimension(2) | (/0,0/) | Charge of the particles |
| srtsRaiseFlag | logical | .false. | <ul style="list-style-type: none"> • if .true. then the srts stepping is done • if .false. then the ekin_lab stepping is done |
| ekin_lab | real | 1. | kin. energy of first particle in the rest frame of second particle (starting value for the energy scan: the number of different energies is set by parameter num_Energies in the namelist "input") |
| delta_ekin_lab | real | 0.03 | kin. energy step for energy scan |
| srts | real | 3. | invariant energy (starting value for the energy scan) |
| delta_srts | real | 1. | srts step for srts scan |

| Elementary_Analysis code/analysis/ElementaryAnalysis.f90 | | | |
|---|---------|---------|--|
| DoOutChannels | logical | .false. | switch on/off: reporting of all final state channels |
| DoH2d | logical | .false. | if .true. than make output of 2-dimensional histograms (they could produce files of size 240 mb) |
| Do45ForAllEvents | logical | .false. | flag to decide, whether DoElementaryAnalysis4(5).dat is written for all events or just for events, where the output channel consist of pions |
| DodNNbar | logical | .false. | |
| DoPanda | logical | .false. | if .true., elementary analysis for channels with S = -2 and -3 (Xi, Omega) |
| Dodsigdt | logical | .false. | |
| Do2Part | logical | .false. | |

| eN_Event code/init/ElectronGenerator/eN_event.f90 | | | |
|--|---------|--------|---|
| selectFrame | integer | 2 | select frame, in which the calculaton of W_free is done: <ul style="list-style-type: none"> • 0 = doNOT — do NOT remove • 1 = CM • 2 = CALC • 3 = THRE prescription from correct threshold behaviour, used in heavy ion collisions • 4 = NucleonRest : boost nucleon in the rest frame, set free mN, recalculate boson momentum • 5 = THRE2 threshold with m^2: $s_{free}=s+m^2-m^{*2}$ |
| restingNucleon | logical | .true. | if this flag is .false., we use the momentum of the target nucleon in the calculation of the flux |

| EventOutput code/analysis/EventOutputAnalysis.f90 | | | |
|--|---------|---------|---|
| WritePerturbativeParticles | logical | .false. | Flag to write out the perturbative particle vector to an output file. The switch 'EventFormat' determines which format is used. |
| WriteRealParticles | logical | .false. | Flag to write out the real particle vector to an output file. The switch 'EventFormat' determines which format is used. |

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| EventFormat | integer | 1 | <p>This switch determines the format of the event output files. Possible values:</p> <ul style="list-style-type: none"> • 1 = Les Houches format (default) • 2 = OSCAR 2013 format • 3 = Shanghai 2014 format • 4 = ROOT <p>NOTES</p> <ul style="list-style-type: none"> • For Les Houches, the output will be written to files called EventOutput.Pert.lhe and EventOutput.Real.lhe. • For OSCAR, the output files are called EventOutput.Pert.oscar and EventOutput.Real.oscar. • For Shanghai, the output files are called EventOutput.Pert.dat and EventOutput.Real.dat. • For ROOT, the output files are called EventOutput.Pert.root and EventOutput.Real.root. |
| Interval | integer | 0 | <p>Interval for event output, i.e. number of timesteps after which output is written. If zero, only final output at the end of the time evolution is produced.</p> |

externalSystem

code/init/initExternal.f90

| | | | |
|-----------------|----------------|----------------|---|
| inputFile | character*1000 | './source.inp' | <p>the absolute name of the input file with hadrons to be propagated. possible values:</p> <ul style="list-style-type: none"> • if not set, default is './source.inp' • if given, but does not contain '/': default is './[inputFile]' • otherwise: filename is absolute, including path <p>NOTE if you want to use the file 'XXX.inp' in the actual directory, give it as './XXX.inp'</p> |
| DoPerturbative | logical | .false. | <p>if true, the particles will be inserted into the perturbative particle vector, the real particles have to be initialized via some nucleus definition</p> |
| NumberingScheme | integer | 1 | <p>The way, how particles%event will be numbered:</p> <ul style="list-style-type: none"> • 1: event = iPart, i.e. the particle number in the ensemble (historical, but does not work for fullensemble) • 2: event = -999 (should work for perturbative init) |

| ff_QE | code/init/lepton/formfactors_QE_nucleon/FF_QE_nucleonScattering.f90 | | |
|------------------|---|---------|---|
| parametrization | integer | 3 | <ul style="list-style-type: none"> • 0 = dipole approximation • 1 = BBA03 parametrization • 2 = BBBA05 parametrization • 3 = BBBA07 parametrization |
| MV2 | real | 0.71 | vector mass squared in the dipole parametrization of the vector form factors |
| MA_in | real | 1.0 | axial mass (only if useNonStandardMA=.true.) |
| useNonStandardMA | logical | .false. | if one wants to use a specific axial mass, set this to true and choose value for MA_in |
| deltas | real | -0.15 | strange contribution to the axial ff. |
| axialMonopole | logical | .false. | use axial ff. of Gari, Kaulfuss PLB 138 (1984) |

| FinalState_Full | code/collisions/phaseSpace/finalState_Full.f90 | | |
|------------------------|--|---------|---|
| maxbwd_scalingFactor | real | 1. | • Rescales maxBWD |
| silentMode | logical | .true. | • Switches error messages off in massAss. Errors can still be seen looking at massAssStatus.dat |
| NYK_isotropic | logical | .false. | If .true., the angular distribution in Nucleon-Hyperon-Kaon production is assumed to be isotropic. If .false., a non-isotropic distribution is used, as described in Larionov/Mosel, Phys.Rev. C 72 (2005) 014901. See also momenta_in_3Body_BYK. |

| formfactors_pion | code/init/lepton/formfactors_pionProduction/formfactors_A_input.f90 | | |
|-------------------------|---|---|--|
| which_MaidVersion | integer | 2 | choice of MAID version: 1=2003, 2=2007 |

| Freezeout | code/analysis/FreezeoutAnalysis.f90 | | |
|------------------------|-------------------------------------|---------|--|
| FreezeoutAnalysis_Pert | logical | .false. | Flag to do freeze out analysis for perturbative particles |
| FreezeoutAnalysis_Real | logical | .false. | Flag to do freeze out analysis for real particles |
| potThreshold | real | 0.005 | threshold value in GeV. If the absolute value of the potential is below this value, the particle is considered to be 'free', e.g. it 'escaped' |

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|----------------------------|---|--------|--|
| gamma_2Pi_Xsections | code/init/lowPhoton/twoPi_production/gamma2Pi_Xsections.f90 | | |
| experimentalXsections | logical | .true. | <ul style="list-style-type: none"> • If .true. then the Xsections are taken from the experiment • If .false. then the theoretical values are given |

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|-----------------|--------------------------|---------|--|
| hadron | code/init/initHadron.f90 | | |
| impactParameter | real | 0. | smaller 0: Impact parameter will be chosen randomly in the interval $[0; \text{abs}(\text{impactParameter})]$ (see subroutine setGeometry). It is recommended to take very large negative value of impactParameter in order to have good automatic random choice, e.g. <code>impactParameter=-100</code> . |
| bRaiseFlag | logical | .false. | if .true.: actual impact parameter will be raised by <code>deltaB</code> after <code>nRunsPerB</code> subsequent runs. Starting value is given by the <code>impactParameter</code> variable. |
| deltaB | real | 0. | impact parameter step (relevant if <code>bRaiseFlag=.true.</code>) |
| nRunsPerB | integer | 1 | number of subsequent runs per impact parameter (relevant if <code>bRaiseFlag=.true.</code>) |
| perturbative | logical | .false. | if .true. then the hadron is a perturbative particle |
| numberParticles | integer | 200 | Number of projectile testparticles per ensemble in the case of a perturbative treatment |
| particleId | integer | 1 | Identity of the projectile hadron. |
| antiParticle | logical | .false. | if .true. then the hadron is an antiparticle |
| particleCharge | integer | 0 | Charge of the hadron |
| ekin_lab | real | 1. | kinetic energy of the hadron in the rest frame of the target nucleus (GeV) NOTES If <code>ekin_lab < 0</code> . — initialization according to the binding energy |
| E_bind | real | 0. | binding energy of initialized hadron (GeV) NOTES Active for <code>iniType= 0,2</code> if <code>ekin_lab < 0</code> . is set. |
| iniType | integer | 0 | <ul style="list-style-type: none"> • 0: usual initialization for the hadron-nucleus collision • 1: position and momentum of the hadron is chosen according to the Gaussians centered, resp., at the centre of the nucleus and at zero momentum (<code>impactParameter</code>, <code>distance</code> and <code>ekin_lab</code> have no effect in this case) • 2: gaussian in coordinate space, but usual sharp momentum choice (<code>impactParameter</code>, <code>distance</code> and <code>ekin_lab</code> work as usual) |

| | | | |
|--------------------------------------|---------|-----|--|
| zChoice | integer | 1 | <ul style="list-style-type: none"> • 1: hadron is initialised at fixed distance delta from nuclear surface • 2: hadron is initialised at fixed z |
| Relevant for iniType=0 or iniType=2. | | | |
| delta | real | 0.5 | <ul style="list-style-type: none"> • for zChoice=1: distance from nuclear surface, at which the hadron is initialised [fm] • for zChoice=2: maximum distance from the edge of nucleus in transverse direction which restricts the choice of actual impact parameter for impactParameter < 0 (for impactParameter > 0 no restriction) |
| Relevant for iniType=0 or iniType=2. | | | |
| deltaZ | real | 5. | z = -deltaZ - R_nucleus, where z is z-coordinate of the hadron Relevant for iniType=0,2 and zChoice=2. |
| width | real | 1. | Width of a gaussian density profile [fm]. Only relevant for iniType= 1 and 2. |

HadronAnalysis

code/analysis/hadronAnalysis.f90

| | | | |
|--------------|---------|---------|---|
| flagAnalysis | logical | .false. | <p>If true, perform the output of a hadron at the latest time step before the hadron disappeared (file DoHadronAnalysisTime.dat). The output hadron has the same baryon/meson type and antiparticle-flag as the beam particle. In case if the hadron did not disappear, the output is done at the end of the time evolution. The output for the hadron is also done in three another files if its momentum becomes for the first time less than the cut values pCut1 and pCut2 (files DoHadronAnalysisTime1.dat and DoHadronAnalysisTime2.dat) and if it becomes bound (DoHadronAnalysisTime3.dat)</p> <p>NOTES</p> <p>Presently feasible only for real particle simulations.</p> |
|--------------|---------|---------|---|

hadronformation

code/collisions/twoBodyReactions/hadronFormation.f90

| | | | |
|-------------|------|-----|---|
| tauProda | real | 0.5 | in formation time concept 2) and 3) for "error particles": production time of non-leading in rest frame of hadron (in fm) |
| tauForma | real | 0.8 | in formation time concept 1) and in concept 2),3) for "error particles": formation time in rest frame of hadron (in fm) |
| tauFormaFak | real | 1.0 | in formation time concept 1): scale factor for constituent quark model, rescales #(lead quarks)/#quarks |

| | | | |
|----------------|---------|---------|---|
| useJetSetVec | logical | .true. | Flag to select fragmentation time estimates: <ul style="list-style-type: none"> • false → old concept 1) • true → new concepts 2) and 3) NOTES select false in case of calculations on a nucleon (speed!). |
| powerCS | real | 1.0 | in formation time concept 2): power of 't' (constant, linear, quadratic) |
| useTimeFrom | integer | 1 | in formation time concept 2): encode time XS starts to evolve: 1: tP_min, 2: tP_max, 3: tF |
| useTimeTo | integer | 3 | in formation time concept 2): encode time XS stops to evolve: 1: tP_min, 2: tP_max, 3: tF |
| GuessDiffTimes | logical | .true. | if true, then the times for diffractive particles are treated like them of all other particles, otherwise particles from "diffractive" events hadronize immediately. |
| useJetSetVec_Q | logical | .true. | if useJetSetVec, then also use Q2 as measure for XS-pedestal, i.e. select concept 3) instead of concept 2) |
| useJetSetVec_R | logical | .true. | if not useJetSetVec_Q, then use rLead as measure for XS-pedestal |
| pedestalCS | real | 0.0 | in formation time concept 2): encode time XS stops to evolve: 1: tP_min, 2: tP_max, 3: tF |
| useQDM | logical | .false. | If true, then use the quantum diffusion model of G.R. Farrar et al., PRL 61, 686 (1988). It means that the cross section grows as $(t-t_{int})^{**powerCS}$ for $t_{int} < t < t_{form}$, where t_{int} is the interaction time (=0 for electron-nucleus case) and $t_{form} = t_{int} + 2*p/dM2$. So hadrons with equal momenta have equal formation times (lengths). Also allows to control the space-time scale of hadronization. Attention: setting useQDM = .true. overrides other switches of this module. |
| dM2 | real | 0.7 | Mass denominator in the coherence length. Relevant only for quantum diffusion model (when useQDM =.true.) |
| use_pCut | logical | .false. | If true, then only particles with momentum $p < pCut$ will interact. |
| pCut | real | 1. | Momentum cutoff. Relevant only when use_pCut =.true. |

| | | |
|-----------------------------|---|--------|
| HadronTensor_ResProd | code/init/lepton/hadronTensor_ResProd.f90 | |
| speedup | logical | .true. |

| heavyIon | | code/init/initHeavyIon.f90 | |
|---------------------|---------|----------------------------|---|
| impact_parameter | real | 0. | Impact parameter b [fm]. There are three options: <ul style="list-style-type: none"> • $b \geq 0$: The impact parameter is fixed to the given value. • $-100 < b < 0$: The impact parameter will be chosen randomly in each run between 0 and $\text{abs}(b)$. • $b \leq -100$: "Minimum bias". The impact parameter will be chosen randomly in each run (maximum = sum of radii plus twice the sum of surfaces). |
| impact_profile | integer | 0 | This switch provides impact-parameter distributions for trigger-biased setups. Only used for <code>impact_parameter < 0</code> . Possible values: <ul style="list-style-type: none"> • 0: minimum bias (default) • 1: HADES C+C at 1.00 AGeV • 2: HADES C+C at 2.00 AGeV • 3: HADES Ar+KCl at 1.76 AGeV • 4: HADES Au+Au at 1.23 AGeV (all) • 5: HADES Au+Au at 1.23 AGeV (0-10% central) • 6: HADES Au+Au at 1.23 AGeV (10-20% central) • 7: HADES Au+Au at 1.23 AGeV (20-30% central) • 8: HADES Au+Au at 1.23 AGeV (30-40% central) |
| distance | real | 0. | Distance between centers of nuclei along z (i.e. beam)-direction [fm]. This will be readjusted automatically in case it is too small. |
| coulomb | logical | .false. | If <code>.true.</code> , then a Coulomb propagation from <code>coulombDistance = 10000 fm</code> to distance is performed. |
| ekin_lab_Target | real | 0. | Kinetic energy per nucleon of target nucleus in lab frame [GeV]. |
| ekin_lab_Projectile | real | 0. | Kinetic energy per nucleon of projectile nucleus in lab frame [GeV]. |
| adjustGridFlag | logical | .false. | If <code>.true.</code> , the grid spacing in z-direction will be readjusted. |
| cmsFlag | logical | .true. | If <code>.true.</code> , the collision takes place in the CM frame of the two nuclei (default option). If <code>.false.</code> , the collision takes place in the LAB frame (target at rest). |

HICanalysis_Input code/analysis/HeavyIonAnalysis.f90

| | | | |
|-----------------|---------|---------|---|
| flag_outputReal | logical | .false. | If <code>.true.</code> , then the output of the real particle vector will be written to the file 'DoHIA.dat'. |
|-----------------|---------|---------|---|

| | | | |
|---------------------|----------------------|---|---|
| flag_outputPert | logical | .false. | If .false., then the output of the perturbative particle vector will be written to the file 'DoHIA_pert.dat'. |
| flag_outputDetailed | logical | .false. | Print out more detailed information at each time step from subroutine HeavyIon_evol: <ul style="list-style-type: none"> • rhorad_*.dat • rhoz_*.dat • rhozx_*.dat • Fields_*.dat • pauli_*.dat • dens_max.dat |
| pionAnalysis | logical | .false. | This flag generates various pion spectra (p-T, m-T, y, etc). The analysis operates under the assumption of a fixed target, and expects the collision to be performed in the CMS system (cf. cmsFlag in namelist /heavyIon/). The analysis matches the one applied to the HADES data in Agakishiev et al., Eur.Phys.J. A40 (2009) 45-49. |
| rapBinning | real, dimension(0:7) | (/ -0.75, -0.45, -0.15, 0.15, 0.45, 0.75, 1.05, 1.35 /) | Rapidity binning for the pion analysis (only used if pionAnalysis = .true.). The numbers represent the binning borders in y0. For each of the seven y0 bins, a separate mT spectrum will be generated. |
| KaonAnalysis | logical | .false. | This flag generates various Kaon spectra and Kaon-related analyses. |
| DensityPlot | logical | .false. | This flag select printing the density for several time steps |
| NucleonMassPlot | logical | .false. | This flag select printing the (invariant) mass of the nucleons for several time steps |
| do_Tmunu | logical | .false. | Switch for Tmunu output. |
| rotateZ_Tmunu | logical | .false. | select, whether the particles are first rotated to be aligned to the z-axis |
| correctPot_Tmunu | integer | 0 | select, whether the energy is corrected for the potential or not: <ul style="list-style-type: none"> • 0: no correction • 1: full potential added to p0 • 2: only $U_b/2+U_r$ added to p0 |

| HiGammaNucleus | code/init/ElectronGenerator/eventGenerator_eN_HiEnergy.f90 | | |
|-----------------------|--|---------|--|
| DoLowEv | logical | .true. | If this flag is set true, then for $W_{\text{free}} < \text{HighEnergyThreshold}$ we will call the low energy model routines. |
| DoTransEv | logical | .false. | flag: use transitionEvent in order to replace PYTHIA events by events where we give the cross section explicitly and do the remaining stuff by FRITIOF NOTES this replaces the flag "FRITIOF" in the namelists "HiLeptonNucleus" and "HiPhotonNucleus" |
| useHermesPythiaPars | logical | .false. | flag: Use "PYTHIA tuning done by HERMES collab" |
| DoDiffr | logical | .true. | flag: Generate diffractive events |
| PYTHIAthresh | real | 2.0 | Below this value for W , PYTHIA is not used to generate (G)VMD events NOTES This value is transferred to PyVP.f. you can access this value by the function "GetPYTHIAthresh()". |
| useVMD_VM | logical, dimension(4) | .true. | These flags can be used to switch on/off some VM in the VMD description of the events generated by "transitionevent" NOTES <ul style="list-style-type: none"> • The VMD events of PYTHIA are not affected. (We could change this!) |
| useRes | logical, dimension(2:nres+1) | .true. | Switch for including/excluding specific resonances |
| allowRes | logical | .true. | Switch for including/excluding resonance contribution. If this is set to .true., 1pion events will just be generated as for the background, but according the full MAID cross section (if at all) |
| allow1pi | logical | .true. | Switch for including/excluding 1pion contribution. Depending on the switch allowRes, 1 pion events will be done according the full cross section or just as a background. |
| allow2piBack | logical | .true. | Switch for including/excluding additional 2pion background. |
| allowDIS | logical | .true. | Switch for including/excluding DIS contribution |
| allowVMDrho | logical | .true. | Switch for including/excluding the VMD gamma $N \rightarrow \rho_0 N$ contribution |
| DoToyModel_pi | logical | .false. | flag: Use a Toy model instead of realistic event generation. Only a single pion is generated |

| | | | |
|--------------------------|---------|---------|---|
| DoToyModel_rho | logical | .false. | flag: Use a Toy model instead of realistic event generation Only rho0 N events are generated. Additional assumptions: (c.f.UseFormTime_ToyModel_rho) * tau_F = 0 * tau_F = m with t_F = E (boost according E/m, not E_string/M_string) In the latter case we suffer also the following simplifications: <ul style="list-style-type: none"> • no Q2 dependance • XS starts with n.L/n = 0.5 (should be 0.66 for the nucleon) |
| UseFormTime_ToyModel_rho | logical | .false. | flag: if .true., we set the formation times of the particles produced in the Toy-Model_rho equals to the energy of the particle (t_f/fm=E/GeV) representing the assumption tau_f/fm = m/GeV plus a boost according E/m. (Otherwise the formation time is set to zero.) |
| DoExclPiModel | logical | .false. | flag: Use a model for exclusive pion production. Only those events are generated |
| ExclPiCharge | integer | 1 | variable to specify the charge of the pion produced, if DoExclPiModel is selected |
| flagTwoJets | logical | .false. | If .true. - the events without two jets with large transverse momentum are marked with XS_tot=100000 mub. |

| HiLepton_Analysis | | code/analysis/HiLeptonAnalysis.f90 | |
|--------------------------|---------|------------------------------------|---|
| DoTimes | logical | .false. | switch on/off: reporting of times |
| DoOutChannels | logical | .false. | switch on/off: reporting of all final state channels |
| DoInvMasses | logical | .false. | switch on/off: reporting of pairwise-invariant-masses |
| DoFindRho0 | logical | .false. | switch on/off: reconstructing rho0 from final pions |
| DoClasie | logical | .false. | switch on/off: Do pion analysis as Clasie et al., arXiv:0701.1481 |
| DoMorrow | logical | .false. | switch on/off: Do pion analysis as Morrow et al., Morrow:2008ek |
| DoBrooks | logical | .false. | switch on/off: Do pi+ pT2 spectra for Brooks delta pT2 |
| DoMandelT | logical | .false. | switch on/off: Do pion analysis with Mandelstam t. |
| DoClassifyFirst | logical | .false. | Classifying 'FirstEvent' into some classes Needs DoEventAdd. |
| DoFSIsqrts | logical | .false. | switch on/off: Estimate potential/future final state interactions Plot the sqrt(s) distribution of potential final state interactions of perturbative particles with the nucleus (real) particles). (The interactions do not happen, this is calculated before every propagation.) In order to select the particle class for which one wants to report the FSI, please change directly the code. |

| | | | |
|--------------------|---------|---------|--|
| DoCentralN | logical | .false. | switch on/off: Do centrality analysis with slow nucleons |
| DoLeptonKinematics | logical | .false. | switch on/off lepton kinematics output |
| DoHadronKinematics | logical | .false. | switch on/off hadron kinematics output |
| flagDoIt | logical | .true. | switch on/off using DoHiLeptonAnalysis |

HiLeptonNucleuscode/init/initHiLepton.f90

iExperiment

integer

0

choice of experiment, detector and energy
possible values are:

- 0: no experiment/fixed kinematics
- 1: Hermes, 27GeV, D,N,Kr
- 2: Hermes, 27GeV, Ne
- 3: Hermes, 27GeV, H
- 4: JLAB, 12GeV
- 5: JLAB, 5GeV
- 6: EMC, 100GeV
- 7: EMC, 120GeV
- 8: EMC, 200GeV
- 9: EMC, 280GeV
- 10: Hermes, 12GeV
- 11: Hermes, 27GeV, arXiv:0704.3270
- 12: Mainz, Yoon: Ebeam=1.5GeV
- 13: Hermes, 27GeV, arXiv:0704.3712 (pT-broadening)
- 14: JLAB, 5GeV, rho0 experiment
- 15: JLAB, 4GeV, rho0 experiment
- 16: EIC, E_e and E_A given explicit (3+30,11+30,4+100)
- 17: no detector, total cross section, Ebeam
- 18: E665, 470GeV
- 19: CLAS/JLAB, 12GeV RunGroupA optimized 10.6 GeV
- 20: CLAS/JLAB, 12GeV RunGroupA theoretical

please note: The entry "iExperiment == 0" replaces the old HiPhoton event
type.

shadow

logical

.true.

flag: Consider shadowing or not

minimumMomentum

real

0.1

minimal momentum considered. (in GeV)

| | | | |
|-------------------|---------|---------|--|
| ModusCalcFluxNorm | logical | .false. | if this flag is true, than we do not really generate events. We only select nu and Q2 according an equal distribution and plot the flux (and the flux multiplied with AccWeight). Normally we choose nu and Q2 according flux*Accweight via von-Neumann- rejection method (where we loose access to the absolute normalisation). |
| iDetector | integer | -1 | This sets the treatment of the detector: <ul style="list-style-type: none"> • -1 : not valid/not initialized/use default • 0 : no detector, as AccFlag=.false. • 1 : HERMES, full efficiency • 2 : EMC, full efficiency • 3 : CLAS, only cuts (th_e=12..50, th_hadron=6..143) • 4 : CLAS, full efficiency + cuts as for 5GeV • 5 : CLAS, electron: cuts (th_e=12..50), hadrons: efficiency+cuts as for 5GeV • 90 : full acceptance |
| EIC_Ee | real | -99.9 | the electron beam energy, if iExperiment=EIC |
| EIC_EA | real | -99.9 | the hadron beam energy, if iExperiment=EIC |
| realRun | logical | .false. | Flag to indicate, whether we produce real or perturbative particles. NOTES run with real particles untested !!! |
| DoStatistics | logical | .false. | switch on/off statistical output of init routines |
| user_numin | real | -99.9 | user given value for numin, overrides default value if reasonable |
| user_numax | real | -99.9 | user given value for numax, overrides default value if reasonable |
| user_costmin | real | -99.9 | user given value for costmin, overrides default value if reasonable |
| user_costmax | real | 99.9 | user given value for costmax, overrides default value if reasonable |
| user_ymax | real | -99.9 | user given value for ymax, overrides default value if reasonable |
| user_smin | real | -99.9 | user given value for smin, overrides default value if reasonable |
| user_xBmin | real | -99.9 | user given value for xBmin, overrides default value if reasonable |
| user_qsqmin | real | -99.9 | user given value for qsqmin, overrides default value if reasonable |
| user_qsqmax | real | -99.9 | user given value for qsqmax, overrides default value if reasonable |
| user_maxw | real | -99.9 | user given value for maxw, overrides default value if reasonable |

| | | | |
|------------|---------|---------|---|
| earlyPauli | logical | .false. | <p>Flag to indicate, whether we should check Pauli blocking already during generation or only at the end.</p> <p>if .false. (default), events will be generated in a first stage without Pauli blocking. This is then tested afterwards. If the generated event is blocked, it will be redone! Thus Pauli blocking does <i>*not*</i> change the total cross section, only the relative strength will be reshuffled.</p> <p>if .true., then blocked events will be excluded from the Monte Carlo decision and the total cross section will be reduced.</p> <p>NOTES</p> <p>The behaviour, if no event at all is possible, is at the moment a little bit unpredictable ;)</p> |
|------------|---------|---------|---|

| HiPhotonKinematics | code/init/initHiLepton.f90 | | |
|---------------------------|----------------------------|-------|--|
| nu | real | -99.9 | Photon energy [GeV] |
| Q2 | real | -99.9 | transfer four momentum squared [GeV ²] |
| eps | real | -99.9 | Photon polarisation [1] |
| srts | real | -99.9 | sqrt(s) of electron nucleon system [GeV] |
| W | real | -99.9 | sqrt(s) of photon nucleon system [GeV] |
| xBj | real | -99.9 | Bjorken x [1] |
| Ebeam | real | | electron beam energy [GeV] |

| HiPion_Analysis | code/analysis/HiPionAnalysis.f90 | | |
|------------------------|----------------------------------|---------|--|
| Enable | logical | .true. | If .true. the HiPion analysis will be performed, otherwise not. |
| EnablePerTime | logical | .false. | If .true. the HiPion analysis per timestep will be performed, otherwise not. |
| DoSimpleKin | logical | .false. | switch on/off: Analysis for simple kinematics: pZ-, pT-spectra etc. |
| DoHarp | logical | .false. | switch on/off: Analysis for the HARP experiment |
| DoBlobel | logical | .false. | switch on/off: Analysis according Blobel et al. |
| DoInvMasses | logical | .false. | switch on/off: reporting of pairwise-invariant-masses |
| DoOutChannels | logical | .false. | switch on/off: reporting of all final state channels |

| HiPionNucleus | code/init/initHiPion.f90 | | |
|----------------------|--------------------------|---------|--|
| distance | real | 15. | Distance in z-direction from the nucleus center in fm, where the projectiles are initialized. If negative, the distance will be chosen automatically. |
| impact_parameter | real | 0. | Impact parameter of the projectiles in fm. If positive (or zero), this fixed value is used for all projectiles. If negative, the impact parameter is chosen by Monte Carlo, so that the projectiles are distributed over a certain disk. Cf. 'setPosition'. |
| ProjectileCharge | integer | 0 | Charge of projectile particles. |
| ProjectileID | integer | pion | ID of projectile particles. |
| ProjectileAnti | logical | .false. | Antiparticle flag of projectile particles. |
| nTestparticles | integer | 200 | Number of projectile testparticles per ensemble. |
| ekin_lab | real | -99.9 | Kinetic energy of projectile particles in lab frame [GeV]. |
| p_lab | real | -99.9 | Momentum of projectile particles in lab frame [GeV/c]. |
| DoPerturbativeInit | logical | .false. | <p>If this flag is set to .true., the first collision of the projectile particles with a nucleon in the target nucleus will be done in this init routine (at timestep 0). This enables you to treat the first (hard) collision different from those in the FSI.</p> <p>If this flag is set to .false., the projectile particles have to be propagated onto the nucleus as in the default transport treatment.</p> <p>See documentation of 'initHiPionInducedCollide' and 'initHiPionInducedCollideFull' for further information.</p> |
| DoOnlyOne | logical | .true. | <p>If the first interaction of beam and target particles is treated already here in the init (cf. DoPerturbativeInit), you may select whether a beam particle may interact only once (flag set to .true.) or with all other target nucleons (flag set to .false.).</p> <p>See documentation of 'initHiPionInducedCollide' and 'initHiPionInducedCollideFull' for further information.</p> |
| minimumMomentum | real | 1.0 | Minimal momentum of particles (in GeV) produced in the init routines. Only particles with an absolute momentum larger than this will be further propagated. |
| useHermesPythiaPars | logical | .false. | flag: Use "PYTHIA tuning done by HERMES collab" |
| NucCharge | integer | -1 | Select charge state of nucleons to scatter on. If this value is ≥ 0 , then we only scatter on nucleons with the respective charge, i.e. only on neutrons if NucCharge==0 and only on protons if NucCharge==1. Useful e.g. for selecting only pn events in a pd collision. |

| History | | code/collisions/history.f90 | |
|-----------------------|---------|-----------------------------|---|
| IncGeneration_Decay | logical | .true. | This flag determines whether we will increase the stored 'generation' of the daughter particles in a resonance decay. |
| IncGeneration_Elastic | logical | .true. | This flag determines whether we will increase the stored 'generation' of particles in an elastic collision. Setting it to .false. will also prevent elastic collisions from showing up as parents in the history. |

| InABoxAnalysis | | code/analysis/InABoxAnalysis.f90 | |
|-----------------------|---------|----------------------------------|--|
| Enable | logical | .true. | Flag to enable or disable the box analysis altogether. |
| Interval | integer | 20 | Interval for output, i.e. number of timesteps after which output is written. |

| initDatabase | | code/database/particleProperties.f90 | |
|------------------------|---------|--------------------------------------|---|
| propagationSwitch | integer | 3 | <ul style="list-style-type: none"> • 0 = propagate resonances with more than 1 star in their rating (irrespar=0 in old code) • 1 = propagate just the Delta (irrespar=2 in old code) • 2 = propagate no resonance (irrespar=3 in old code) • 3 = propagate all resonances (default) |
| usageForXsectionSwitch | integer | 2 | <ul style="list-style-type: none"> • 0 = use resonances with more than 1 star rating for cross sections • 1 = use all resonances for cross sections • 2 = use all resonances besides the 1* star I=1/2 resonances • 3 = use only the Delta |
| rho_dilep | logical | .false. | If .false. (default), the rho meson width will be exclusively given by the 2pi decay and its minmass will be 2m_pi. If .true., the dilepton width will be included in the width and spectral function of the rho, and the minmass will be 2m_e. This is important for dilepton spectra, in order to get contributions from the rho below the 2pi threshold. |

| | | |
|-----------------------|------------------|---|
| FileNameDecayChannels | character(1000)" | <p>The absolute filename of the file containing decay channel infos. possible values:</p> <ul style="list-style-type: none"> • if not set, default is '[path_To_Input]/DecayChannels.dat' • if given, but does not contain '/': default is '[path_To_Input]/[FileNameDecayChannels]' • otherwise: filename is absolute, including path <p>NOTE if you want to use the file 'XXX.dat' in the actual directory, give it as './XXX.dat'</p> |
|-----------------------|------------------|---|

| | | | |
|----------------------|--------------------------|-----------------|---|
| initDensity | | | |
| | code/density/density.f90 | | |
| densitySwitch | integer | 1 | <p>This switch decides whether the density is static or dynamic during the run. ("Static" makes sense only for fixed target scenarios!) One can use a static density if the nucleus stays roughly in its ground state during the collision. possible values:</p> <ul style="list-style-type: none"> • 0: Density is set to 0. • 1: Dynamic density according to test-particle distribution. • 2: Static density (not for heavy-ion collisions). • 3: Resting matter: Density is given by the two input parameters "densityInput_neutron" and "densityInput_proton". |
| linearInterpolation | logical | .true. | If this switch is 'true', then the dynamic-density mode uses linear interpolation to determine the density in between the gridpoints. |
| densityInput_proton | real | 0.084 | Assumed proton density if densitySwitch=3 |
| densityInput_neutron | real | 0.084 | Assumed neutron density if densitySwitch=3 |
| gridSize | real, dimension(1:3) | (/12.,12.,12./) | Size of density grid in fm. |
| gridPoints | integer, dimension(1:3) | (/30,30,30/) | Number of gridpoints in each space direction. |
| setnewsmearing | logical | .false. | Readjust the smearing to a different width if .true. |
| newsmearing | real | 1. | Use a smearing width as in a grid with newsmearing times the gridspaceing |
| numberLargePoints | integer | 2 | Number of points which are considered to the left and right to smear density on |

| initInABox | | code/init/initInABox.f90 | |
|----------------------|---------|--------------------------|---|
| proton_Density | real | 0.084 | • proton Density [fm ⁻³] |
| neutron_Density | real | 0.084 | • neutron Density [fm ⁻³] |
| fermiMotion | logical | .true. | • true = switch on Fermi motion • false = switch off Fermi motion |
| temp | real | 0. | If fermiMotion is true, this switch determines the temperature (in GeV) used in the Fermi distribution. |
| energy_density | real | 0. | Energy density in GeV/fm ³ . If a finite positive number is given, the box will be boosted to a frame with the given energy density. |
| standing_wave_number | integer | 0 | If this number is larger than zero, the initial density distribution will not be uniform, but is modulated with a standing wave in z direction. The given number determines the number of oscillations throughout the box. The amplitude of the oscillations is currently fixed to be 20% of the (average) nucleon density. |

| initNbarN_to_NbarDelta | | code/collisions/twoBodyReactions/baryonBaryon/NbarN_to_NbarDelta.f90 | |
|-------------------------------|---------|--|---|
| delta_mass | real | 0.01 | • grid step on a delta mass (GeV) |
| maxPoints_mass | integer | 150 | • number of the grid points on the delta mass |
| delta_srts | real | 0.01 | • grid step on an invariant energy (GeV) |
| maxPoints_srts | integer | 100 | • number of the grid points on the invariant energy |

| InitNucleus_in_PS | | code/init/initNucPhaseSpace.f90 | |
|--------------------------|---------|---------------------------------|---|
| improvedMC | logical | .false. | • If this flag is set to .true. then we use the information of the already initialized nucleons to decide on the position of a nucleon which has to be initialized. • This prescription does only work properly if the smearing with is really small. Therefore it is switched off by default. |
| improvedMC_speedup | integer | 500 | • If improvedMC is set to .true. then this variable defines the speedup of the algorithm. • The number defines how often the density field is updated. • A large value of this parameter yields a less accurate test-particle distribution and a faster initialization. |
| HiTail | logical | .false. | If HiTail is set to .true., then a simple parametrization of n(p) is used to initialize the nucleon momenta (cf. function chooseAbsMomentum for details). |

| | | | |
|--|---------|---------|---|
| determine_Fermi_momentum_by_binding_energy | logical | .false. | If set to .true., the Fermi momentum will be determined by $E_B = p_f^2 / (2m) + U(\rho, p_F)$, where E_B is the binding energy per nucleon. |
| determine_Fermi_new_NucDLDA | logical | .false. | If set to .true., the Fermi momentum will be set to a value such that there are no unbound nucleons at the initialisation. |
| useEnergySF | logical | .false. | If set to .true., then a spectral function is used to choose the energy. |
| compressedFlag | logical | .false. | If set to .true., then a spherically deformed nucleus is initialized (isotropic compression/expansion; protons and neutrons in phase). This type of deformation corresponds to a giant-monopole resonance mode. |
| ScaleFactor | real | 1. | If compressedFlag=.true., then rescale coordinates by ScaleFactor. |
| useCdA | logical | .false. | Instead of the usual momentum distribution according to a Fermi gas, use the momentum parametrizations as given in: <ul style="list-style-type: none"> • C. Ciofi degli Ati, S. Simula, PRC 53, 1689 (1996) These exist only for 2H,3He,4He,12C,16O,40Ca,56Fe,208Pb |
| zeroNucleusMomentum | logical | .true. | Indicate whether a procedure should be called to try to find a momentum initialization where the sum of all nucleon momenta (per ensemble) is zero (or at least close to zero). At the moment, only a hill climbing algorithm is available, which changes the directions of the momenta randomly. The resulting averaged nucleus momentum is in the order of 10 MeV. Without that, the average nucleus momentum goes $\sim 0.17 \text{ GeV} \cdot \sqrt{A}$. (Applies only for $A > 2$.) |

| initPauli | | | |
|-------------------|--------------------------------|---------|--|
| | code/density/pauliBlocking.f90 | | |
| pauliSwitch | integer | 1 | <ul style="list-style-type: none"> • 0 : No Pauli blocking • 1 : dynamic Pauli blocking (use actual phase space densities) • 2 : analytic Pauli blocking (use ground state assumption) (not possible for Heavy Ions!) |
| densDepMomCutFlag | logical | .false. | if .true. - the radius in momentum space for selecting nucleons around given nucleon will depend on local Fermi momentum NOTES Used only for dynamic pauli blocking. |
| Gauss | real | 1.0 | Smearing for dynamic pauli blocking |
| cutGauss | real | 2.2 | Cutoff for gauss Smearing |

| | | | |
|--------------|---------|---------|---|
| cutMom | real | 0.08 | <ul style="list-style-type: none"> • for densDepMomCutFlag=.false. — radius of phase space box in momentum space • for densDepMomCutFlag=.true. — minimum radius of phase space box in momentum space |
| cutPos | real | 1.86 | Radius of phase space box in position space |
| nGridPos | integer | 30 | number of points in position space to save weights on |
| ensembleJump | integer | 5 | Parameter for speedup. Only every "ensemblejump"th ensemble is considered to evaluate the probability for pauli blocking. |
| DoHistogram | logical | .false. | if .true., a histogram is filled representing the blocking probability as function of the fermi momentum. You have to call 'WriteBlockMom' explicitly for writing the histogram |

initRandom code/numerics/random.f90

| | | | |
|-------------|---------|---------|---|
| Seed | integer | 0 | Random Seed (used to initialize the random number generator), accessible through the namelist 'initRandom'. If Seed is zero (default), then it is set via "SYSTEM_CLOCK()". |
| resetRandom | logical | .false. | Reread random generator, used by setRandom, useful for debugging. |

initThermoDynamics code/density/thermoDyn.f90

| | | | |
|---------------------|---------|--------|---|
| temperatureSwitch | integer | 1 | <ul style="list-style-type: none"> • 1=groundstate calculations (T=0,mu=E_F) • 2=the full procedure |
| linearExtrapolation | logical | .true. | <ul style="list-style-type: none"> • .true.= Use linear extrapolation for temperature between gridPoints • .false.= Do not use it |

input code/inputOutput/input.f90

| | | | |
|---------------|-----------------|---------------|---|
| path_To_Input | character(1000) | '../buuinput' | Path to input files. This switch needs to be set to the local path of the 'buuinput' directory, which contains various input files for GiBUU. |
| numEnsembles | integer | 300 | Number of parallel ensembles |
| eventtype | integer | 3 | Switch for the type of event possible values: see module eventtypes |

| | | | |
|---------------------|---------|---------|--|
| fullEnsemble | logical | .false. | Switch for the type of simulation: <ul style="list-style-type: none"> • .false.=parallel ensembles • .true.=full ensemble See also "localEnsemble". |
| localEnsemble | logical | .false. | Switch for the type of simulation: <ul style="list-style-type: none"> • .false. = parallel or full ensembles (depending on the value of the fullEnsemble switch). • .true. = fullEnsemble with "local collisionCriteria", see Lang/Babovsky et al., J. Comput. Phys. 106 (1993) 391-396. Setting localEnsemble = .true. will implicitly set fullEnsemble = .true. (disregarding its value in the jobcard). |
| delta_T | real | 0.2 | time difference for time stepping |
| numTimeSteps | integer | 100 | Number of time steps |
| variableTimeStep | logical | .false. | Switch for using of variable time step: <ul style="list-style-type: none"> • .false.= use constant time step delta_T (see above). • .true.= use time step computed from the frequency of collisions. In this case the input delta_T is used as the maximum allowed time step. |
| time_max | real | 30. | Maximum time until which the time evolution will be computed in the case of variableTimeStep = .true. |
| num_energies | integer | 1 | Number of different energies for energy scans |
| num_runs_sameEnergy | integer | 1 | Number of runs with the same energy in the initialization. |
| checkGridSize_Flag | logical | .false. | Switch for checking if particles escape out of grid. possible values: <ul style="list-style-type: none"> • .false.= no check. • .true. = check is performed, and a warning flag is printed out, in case that particles are outside of the grid. • check valid only for real particles. |

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| continousBoundaries | logical | .false. | <ul style="list-style-type: none"> • Switch to turn on continous boundary conditions. • Implications for density and propagation. • This means that particles are propagated according to continous boundaries. A particle leaving the grid will move back in from the opposite side. The densities are carefully constructed such that places at the opposite side contribute to places on the near side. • What is still missing is the full implementation in collision criteria, this is not done yet for the two body collisions! Be careful therefore with the 2-Body-collisions at the edges. A particle at one edge does not see its scattering partner at the opposite edge. |
| FinalCoulombCorrection | logical | .false. | Switch for Coulomb correction at the end of each run of the outgoing particles |
| length_perturbative | integer | -1 | Length of perturbative particle vector (per ensemble). If negative, it will be determined automatically by event type. |
| length_real | integer | -1 | Length of real particle vector (per ensemble). If negative, it will be determined automatically by event type. |
| freezeRealParticles | logical | .false. | Switch for not propagating real particles |
| printParticleVectors | logical | .false. | Switch to turn on the printing of the particle vector at the start and end of a run. |
| printParticleVectorTime | logical | .false. | <ul style="list-style-type: none"> • Switch to turn on the printing of the particle vector as function of time. • Useful for event classes using real particles (HeavyIon,Hadron). • See also 'timeForOutput' and 'timeSequence'. |
| printParticleVectorsFormat | integer | 1 | Select the format for printing the particle vectors. Possible values are: <ul style="list-style-type: none"> • 1: ASCII • 2: binary |
| timeForOutput | real | 50. | <ul style="list-style-type: none"> • Time (fm/c) after which the particle vector is printed during run (see also variable "timeSequence"). • valid only if printParticleVectorTime = .true. |
| timeSequence | real | 10. | <ul style="list-style-type: none"> • Time sequence (fm/c) of time dependent printing of the particle vector • valid only if printParticleVectorTime = .true. |
| DoPrLevel | | | |
| povray_switch | logical | .false. | Switch for generating Povray-Output |

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| LRF_equals_CALC_frame | logical | .false. | <ul style="list-style-type: none"> • Switch to turn on the assumption that calculation frame and LRF frame coincide • Only useful for reactions close to ground state !!! |
| DoFragmentNucleons | logical | .false. | <ul style="list-style-type: none"> • Switch to turn on/off adding of nucleons stemming from fragmentation of bound clusters. |
| PrintCollisionList | | | |

input_FF_Delta code/init/lepton/formfactors_Delta/FF_Delta_production.f90

| | | | |
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| FF_Delta | integer | 1 | This switch decides whether the Paschos form factors (FF_Delta=1) or the Maid form factors (FF_Delta=0) are used. Default is FF_Delta=1. |
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input_FF_ResProd code/init/lepton/formfactors_ResProd/formFactor_ResProd.f90

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| FF_ResProd | integer | 0 | <p>With FF_ResProd (namelist "input_FF_ResProd" in the Jobcard) one can choose how the form factors are calculated:</p> <ul style="list-style-type: none"> • 0: MAID's helicity amplitudes (Luis' helicity expressions - CM frame) • 1: fit of Lalakulich (PRD 74, 014009 (2006)) • 2: MAID's helicity amplitudes (Lalakulich's helicity expressions - LAB frame) |
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| aDelta | real | -0.25 | fit parameter for C_5^A (Adler) |
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| bDelta | real | 0.04 | fit parameter for C_5^A (Adler) |
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| cDelta | real | 3. | fit parameter for C_5^A (Paschos) |
|--------|------|----|-------------------------------------|

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| DeltaAxFF | integer | 1 | <p>choose between different axial form factors for the Delta:</p> <ul style="list-style-type: none"> • 1: Adler • 2: Paschos • 3: dipol |
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|-------------|---------|---------|--|
| HNV_axialFF | logical | .false. | <p>With .true. or .false. HNV_axialFF (namelist "input_FF_ResProd" in the Jobcard) one can choose which axial form factors to use for Delta-resonance:</p> <ul style="list-style-type: none"> • .true. is Hernandez-Nieves-Valverde fit with $C_5^A=0.867, MA=0.985$ (PRD 76) • .false. is as it was used by Lalakulich et al in PRD 74 |
|-------------|---------|---------|--|

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|-----------------------|---------|---------|---|
| nenner_C5A_Lalakulich | real | 3.0 | Factor wich appear in the Lalakulich parameterization of the axial C_5^A form factor 3.0 was fitted to BNL and used in Lalakulich PRD71 and PRD 74 fit of ANL gave 0.5 |
| refit_barnu_axialFF | logical | .false. | With .true. refit_barnu_axialFF (namelist "input_FF_ResProd" in the Jobcard) means that the axial form factors are refitted to explain the low value of antineutrino cross section (exper data Bolognese PLB81,393 (1979)) |
| W_cutOff_lambda | real | 1.071 | Value for lambda in the W-dependent cut-off function. |
| W_cutOff_switch | logical | .false. | Switch to include a W-dependent cut-off function for the vector form factor of the Delta: <ul style="list-style-type: none"> • false = excluded • true = included |
| vector_FF_switch | logical | .true. | Switch to turn off the vector form factors: <ul style="list-style-type: none"> • false = off • true = on |
| axial_FF_switch | logical | .true. | Switch to turn off the axial form factors: <ul style="list-style-type: none"> • false = off • true = on |
| W_cutOff_switchAll | logical | .false. | Switch to include a W-dependent cut-off function for the vector and the axial form factor of all resonances: <ul style="list-style-type: none"> • false = excluded • true = included <p>NOTES we assume the same dependence as for the Delta vector form factor</p> |
| DeltaCouplrelErr | real | 0. | error in percent for $C_5^A(0)$ for the Delta |
| MA | real | 0.95 | delta resonance axial mass parameter. |

insertion

code/collisions/insertion.f90

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|---------------|------|-------|--|
| minimumEnergy | real | 0.005 | Minimal kinetic energy in GeV for produced perturbative nucleons. If their energy is below this threshold, then they are not propagated, i.e. they are not inserted in the particle vector. NOTES This value was formerly given in the namelist "collisionterm". |
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| propagateNoPhoton | logical | .true. | If .true. then we eliminate all photons, such that they are not propagated and do not show up in the particle vector. If .false. then photons are explicitly propagated. |
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Lepton2p2h

code/init/lowElectron/lepton2p2h.f90

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|---------------|----------------------|--------------|--|
| ME_Version | integer | 4 | indicate the type of matrix element parametrisation NOTES possible values: <ul style="list-style-type: none"> • 1: const ME_Norm_XX ! const for CC fitted to MiniBooNE is 1.8e-6 • 2: constant transverse and decreasing with Enu • 3: "Dipole transverse" transverse, fall with Q2 as 4-th power • 4: MEC from E. Christy (8/2015), with parametrization for longitudinal • 5: MEC from Bosted arXiv:1203.2262, with parametrization for longitudinal • 6: MEC additional parametrization, with parametrization for longitudinal • not yet implemented |
| ME_Norm_QE | real, dimension(1:3) | (/1.0, 1.0/) | 1.0, Overall strength of 2p2h matrix element with 2N out for (EM,CC,NC) NOTES The value == 1 gives the coded strength |
| ME_Norm_Delta | real, dimension(1:3) | (/1.0, 1.0/) | 1.0, Overall strength of 2p2h matrix element with NDelta out for (EM,CC,NC) NOTES The value == 1 is a dummy value |
| ME_Mass_QE | real, dimension(1:3) | (/1.0, 1.0/) | 1.0, Cutoff-mass in some parametrizations of 2p2h matrix element for NN out for (EM,CC,NC) NOTES The value == 1 is a dummy value |
| ME_Mass_Delta | real, dimension(1:3) | (/1.0, 1.0/) | 1.0, Cutoff-mass in some parametrizations of matrix element for NDelta out for (EM,CC,NC) NOTES The value == 1 is a dummy value |

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|-----------------|----------------------|-----------------|------|--|
| ME_Transversity | real, dimension(1:3) | (/1.0, 1.0/) | 1.0, | Parametrisation of structure functions for (EM,CC,NC) NOTES The value = 1 chooses structure function W2 so that 2p2h is pure transverse |
| ME_LONG | real, dimension(1:3) | (/0.0, 0.0/) | 0.0, | Parametrization of structure functions for (EM,CC,NC) NOTES The value = 0 turns any additional longitudinal contribution to structure funct. W2 off |
| ME_W3 | real, dimension(1:3) | (/0.,1.0, 1.0/) | | Overall strength factor for structure function W3 only for (CC,NC) NOTES overall strength parameter for structure function W3 |
| ME_ODW | integer | 1 | | switch for choosing the connection between structure functions W1(electron) and W1(neutrino) and W3(neutrino): <ul style="list-style-type: none"> • 1: for expressions from Martini et al • 2: for expressions from O'Connell et al • 3: for expression with relativistic version of O'Connell only for (CC,NC) NOTES <ul style="list-style-type: none"> • O'Connell et al: PR C6 (1972) 719 • Martini et al: PR C80 (2009) 065501 |
| inmedW | integer | 1 | | Controls which inv mass W is used in parametrization of 2p2h W1 NOTES <ul style="list-style-type: none"> • 1: W = static inv. mass in 2p2h parametrization of W1 • 2: W = inv mass for Fermi moving nucleons in potential • 3: W = inv mass for Fermi moving nucleons without potential |
| T | real | 99 | | target isospin, affects only neutrino 2p2h structure function NOTES <ul style="list-style-type: none"> • T = 0, 1 , ... • T = 99 gives T = (N-Z)/2 |

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|------|---------|---|--|
| Adep | integer | 2 | Switch for A-dependence of 2p2h structure function NOTES |
| | | | <ul style="list-style-type: none"> • 1: A-dependence for zero-range force (Mosel, Gallmeister, 2016) • 2: linear A-dependence, normalized to C12 |

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|-------------------|-------------------------------------|------|--|
| lepton_bin | code/init/neutrino/initNeutrino.f90 | | |
| cost_min | real | -1.0 | if detailed_diff_output is TRUE: Minimal cos(theta) of outgoing leptons, used in 2D dsigma/dEdcos(theta) This cut affects *only* the outgoing lepton |
| cost_max | real | +1.0 | if detailed_diff_output is TRUE: Maximal cos(theta) of outgoing leptons, used in 2D dsigma/dEdcos(theta) This cut affects *only* the outgoing lepton |
| delta_cost | real | 0.1 | stepsize of cos(theta) of outgoing leptons, used in 2D dsigma/dEdcos(theta) |
| Elept_min | real | 0.0 | stepsize of min. energy of outgoing leptons, used in 2D dsigma/dEdcos(theta) |
| Elept_max | real | 2.0 | stepsize of maximal energy of outgoing leptons, used in 2D dsigma/dEdcos(theta) |
| delta_Elept | real | 0.01 | stepsize of energy of outgoing leptons, used in 2D dsigma/dEdcos(theta) |

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| low_photo_induced | code/init/lowPhoton/initLowPhoton.f90 | | |
| energy_gamma | real | 0. | Energy of incoming photon in nucleus rest frame (in GeV). |
| delta_energy | real | 0. | Increase of energy for energy scans. |
| energy_weighting | integer | 0 | Determines the relative weight of different photon energies in energy scans Possible values: <ul style="list-style-type: none"> • 0 = flat distribution (all energies are weighted equal) • 1 = exponential distr. (energies are weighted $\sim 1/E$) |

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|--------------------|---------------------------------------|---|--|
| LowElectron | code/init/lowElectron/lowElectron.f90 | | |
| runType | integer | 1 | <ul style="list-style-type: none"> • If runType=1, then we make runs at some fixed angle defined by lowElectron/theta_lf. • If runType=2, then we make runs at some fixed QSquared defined by lowElectron/QSquared |

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| inputType | integer | 1 | Decides which set of variables is used to determine the final electron energy <code>energy_lf</code> and the step size <code>delta_energy_lf</code> : <ul style="list-style-type: none"> • If <code>inputType=1</code>, then we use directly <code>energy_lf</code> and <code>delta_energy_lf</code> as input • If <code>inputType=2</code>, then we use <code>W_min</code> and <code>W_max</code> as input. For this we assume the nucleon to be at rest to calculate <code>energy_lf</code> out of <code>W</code>. • If <code>inputType=3</code>, then we use <code>energy_lf_min</code> and <code>energy_lf_max</code> as input. |
| theta_lf | real | 10. | Theta scattering angle of outgoing electron with respect to the incoming one. Only relevant of <code>runType=1</code> . |
| phi_lf | real | -10. | Phi scattering angle of outgoing electron with respect to the incoming one. If less than 0, then we do a Monte-Carlo-Integration over phi! |
| energy_li | real | 1.2 | Energy of incoming electron in GeV. |
| energy_lf | real | 0.8 | Energy of final state electron in GeV. * Only used if <code>inputType=1</code> |
| energy_lf_min | real | 0.1 | Minimal <code>energy_lf</code> * Only used if <code>inputType=3</code> |
| energy_lf_max | real | 0.1 | Maximal <code>energy_lf</code> * Only used if <code>inputType=3</code> |
| delta_energy_lf | real | 0.8 | <code>delta(Energy)</code> of final state electron in GeV for energy scans. * Only used if <code>inputType=1</code> |
| W_min | real | 0.9 | Minimal <code>W</code> at the hadronic vertex assuming a resting nucleon * Only used if <code>inputType=2</code> |
| W_max | real | 1.9 | Maximal <code>W</code> at the hadronic vertex assuming a resting nucleon * Only used if <code>inputType=2</code> |
| QSquared | real | 0.5 | QSquared of virtual photon. Only relevant of <code>runType=2</code> . |
| Do_QE | logical | .true. | Switch for including or excluding Quasi-Elastic (QE) processes |
| Do_1Pi | logical | .true. | Switch for including or excluding direct Single pion production processes. If the resonances are included (<code>Do_Res=.true.</code>) then only the background part is included. |
| Do_Res | logical | .true. | Switch for including or excluding resonance production processes |
| Do_2Pi | logical | .true. | Switch for including or excluding direct Double pion production processes. If the resonances are included (<code>Do_Res=.true.</code>) then only the background part is included. |
| Do_DIS | logical | .true. | Switch for including or excluding deeply inelastic scattering (DIS) events. Only relevant for <code>W>1.4-1.5 GeV</code> . |
| Do_2p2hQE | logical | .false. | Switch for including or excluding event according <code>gamma* N1 N2 → N1' N2'</code> |

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| Do_2p2hDelta | logical | .false. | Switch for including or excluding event according gamma* N1 N2 → N' Delta |
| minMass_QE | real | 0.3 | Minimal mass of a nucleon in QE event. Prevents super-luminous nucleons when embedded in a Skyrme potential. |
| minEnergy_1pi | real | 0.16 | Minimal q_0 such that pion production processes are considered. |
| onlyDelta | logical | .false. | Switch for including only delta resonance |
| nuclearTarget_corr | logical | .true. | <ul style="list-style-type: none"> • If the input is a nuclear target, then the target nucleus is at rest and we calculate the cross section for nuclear target: use flux with respect to the nucleus. • Use .false. only for debugging. |

lowElePhoto_Analysis code/analysis/lowElectronAnalysis.f90

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| dOmega_switch | logical | .false. | If .true. then also dSigma/dOmega is produced, if false not.. |
| dE_switch | logical | .false. | If .true. then also dSigma/dE is produced, if false not.. |

lowPhotonAnalysis code/analysis/LowPhotonAnalysis.f90

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| outputEvents | logical | .false. | If .true. then all events are printed to file. |
| outputEvents_onlyFree | logical | .false. | If outputEvents=.true. then only particles which may leave the nucleus, i.e. may become "free", are printed to file. |
| KruscheOutput | logical | .false. | If .true. then we perform an analysis as in EPJA22 347-351 (2004) |
| KruscheAnalyse_cut | real | 0. | Value of the cut for the deltaE cut in EPJA22 347-351 (2004). |
| FissumOutput | logical | .false. | If .true. then we perform an analysis as in PRC 53,#3 pages 1278 ff. (1996) Produces dsigma/dOmega/dT_pi for pi^+ |
| photonAnalyse | logical | .false. | Special analysis for final state photons |
| TwoPiOutput | logical | .false. | If .true. then we perform an analysis for 2pi production, including statistics for the mass of the pi-pi pair. |
| pi0gamma_analysis | logical | .false. | Do analysis of pi0 gamma pairs (dsigma/dm), to reconstruct invariant mass spectrum of omega mesons. |
| pi0gamma_momcut | real | 0.5 | Cut on the absolute omega three momentum in GeV, being applied to the pi0 gamma spectrum. |

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| pi0gamma_masscut | real, dimension(1:2) | (/0.,2./) | Cuts on the pi0-gamma invariant mass in GeV, being applied to all pi0-gamma spectra (except the mass spectrum). First component is lower limit, second component is upper limit. |
| pi0gamma_mombin | real | 0.050 | Bin size for the pi0 gamma momentum spectrum in GeV. |
| pi0gamma_massres_sigma | real | 0.025 | Sigma parameter for the exp. resolution smearing (width of the Gauss or Novosibirsk function in GeV). See also pi0gamma_massres_tau. |
| pi0gamma_massres_tau | real | -0.090 | Skewness parameter tau of the Novosibirsk function (for exp. resolution smearing). See also pi0gamma_massres_sigma. |
| Ekin_pi0_cut | real | 0. | Cut on the kinetic energy of neutral pions in the pi0gamma_analysis. Only pions with kinetic energies larger than this cutoff are used for the analysis. |

MassAssInfo code/typeDefinitions/MassAssInfoDefinition.f90

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| UseMassAssInfo | logical | .true. | This switch indicates, whether we want to use the whole MassAssInfo machinery or stick to the old prescription of mass assignment. You may set this switch via the jobcard. Anyhow, if your selection of switches for baryon and medium switches leads to cases which are not yet implemented, this flag is set to false automatically. |
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master_1body code/collisions/oneBodyReactions/master_1Body.f90

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| correctEnergy | logical | .true. | Scale final state momenta to fulfill energy and momentum conservation. If .false. energy conservation is violated |
| StableInFormation | logical | .true. | Particles during its formation time are considered to be stable or not. |
| omegaDecayMediumInfo | logical | .false. | Write out information about all decaying omega mesons to a file called "omegaMediumInfo.dat" (decay point, momentum, density, etc). |

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| omegaDecay_restriction | integer | 0 | <p>This switch, like omegaDecayMediumInfo, helps to analyze omega \rightarrow pi0 gamma decays. It will only have an effect for omegaDecayMediumInfo = .true. Possible values:</p> <ul style="list-style-type: none"> • 0 = none (default) • 1 = vacuum (rho < 0.1 rho0) • 2 = medium (rho > 0.1 rho0) <p>With the default value (0), all omega decays are carried out as usual. For the value 1, the decay products are only kept, if the decay happens in the vacuum (i.e. at rho < 0.1 * rho0). For the value 2, the decay products are only kept, if the decay happens in the medium (i.e. at rho > 0.1 * rho0). If the density does not meet these conditions, the decay products are simply removed and will not be put in the particle vector (and thus they will not appear in the analysis).</p> |
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| master_2body | | code/collisions/twoBodyReactions/master_2Body.f90 | |
| correctEnergy | logical | .true. | Scale final state momenta to fulfill energy and momentum conservation. If .false. energy conservation is violated |
| baryonBaryonScattering | logical | .true. | Switch to turn off baryon-baryon-Scattering |
| baryonMesonScattering | logical | .true. | Switch to turn off baryon-Meson-Scattering |
| mesonMesonScattering | logical | .true. | Switch to turn off meson-Meson-Scattering |
| usePythia | integer | 1 | <p>This flag decides whether to use Fritiof or Pythia for high-energy collisions:</p> <ul style="list-style-type: none"> • 0: use Fritiof • 1: use Pythia <p>NOTES</p> <ul style="list-style-type: none"> • This flag is not used in the baryon-antibaryon channel |
| usePythia_BaB | integer | 0 | <p>This flag decides whether to use Fritiof or Pythia for high-energy baryon-antibaryon collisions:</p> <ul style="list-style-type: none"> • 0: use Fritiof • 1: use Pythia |
| useHiEnergy | logical | .true. | <p>Switch to turn HiEnergy on/off. Formerly known as "useFritiof".</p> <p>NOTES</p> <p>Please be very sure what you are doing when setting this parameter to .false.!</p> |

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|----------------------------------|----------------------|--------------|--|
| HiEnergyThresholdBarMes | real | 2.2 | Sqrt(s) threshold for HiEnergy in Baryon-Meson Reactions |
| HiEnergyThresholdBarMesDelta | real | 0.2 | width for the Sqrt(s) threshold for HiEnergy in Baryon-Meson Reactions |
| HiEnergyThresholdBarBar | real | 3.4 | Sqrt(s) threshold for HiEnergy in Baryon-Baryon Reactions |
| HiEnergyThresholdBarBarDelta | real | 0.1 | width for the Sqrt(s) threshold for HiEnergy in Baryon-Baryon Reactions |
| HiEnergyThresholdBarAntibar | real | 2.38 | Sqrt(s) threshold for HiEnergy in Baryon-Antibaryon Reactions |
| HiEnergyThresholdBarAntibarDelta | real | 0.0 | width for the Sqrt(s) threshold for HiEnergy in Baryon-Antibaryon Reactions |
| useManni | logical | .true. | Flag, whether to use meson-baryon annihilation as proposed by Markus Wagner (Diploma, Giessen 2004), but with some enhanced treatment |
| ElastAngDist | integer | 3 | Choice of angular distribution in (high-energy) elastic collisions (cf. Do-Coll_Elast): <ul style="list-style-type: none"> • 1 = isotropic • 2 = J. Cugnon et al., NPA 352, 505 (1981) • 3 = Pythia (default) |
| flagElastBB | logical | .false. | If .true., use a constant elastic baryon-baryon cross section of 40 mb and no inelastic baryon-baryon scattering. |
| coarse | real, dimension(1:3) | (/3.,4.,4./) | coarse maximal impact parameter (in fm) |
| bmax_nucleonNucleon | real | 2.52 | Real maximal impact parameter for nucleon-nucleon-scattering. Maximal crosssection is <pre>!pre! bMax**2 * pi * 10 mb/fm**2 = (2.52**2*pi*10) mb = 199.5 mb</pre> |
| bmax_nucleonResonance | real | 1.60 | Real maximal impact parameter for nucleon-resonance scattering. Maximal crosssection is <pre>!pre! bMax**2 * pi * 10 mb/fm**2 = (1.60**2*pi*10) mb = 80.4 mb</pre> |
| bmax_hyperonNucleon | real | 2.52 | Real maximal impact parameter for hyperon-nucleon-scattering. Maximal crosssection is <pre>!pre! bMax**2 * pi * 10 mb/fm**2 = (2.52**2*pi*10) mb = 199.5 mb</pre> |
| bmax_baryonPion | real | 2.52 | real maximal impact parameter for baryon pion scattering |
| bmax_baryonMeson | real | 2.52 | real maximal impact parameter for baryon-Meson-scattering |
| bmax_mesonMeson | real | 2. | real maximal impact parameter for meson-meson-scattering |
| correctEnergy_message | logical | .true. | Switch off error message for energy correction failures. |

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| OverrideSigma_PiN | real | -99.9 | Parameter to replace the calculated cross section for pi+N collision by a fixed value (in mb). Only in use if ≥ 0 . The elastic cross section is assumed to be 1/10 of the given value. |
| OverrideSigma_RhoN | real | -99.9 | Parameter to replace the calculated cross section for rho+N collision by a fixed value (in mb). Only in use if ≥ 0 . The elastic cross section is assumed to be 1/10 of the given value. |
| OverrideSigma_PiPi | real | -99.9 | Parameter to replace the calculated cross section for pi+pi collision by a fixed value (in mb). Only in use if ≥ 0 . We set $\text{sigma_elast} = \text{sigma_tot}$ |
| Override_PiPi_ResIsElast | logical | .false. | Flag to replace the calculated cross section for pi+pi collision; The calculated resonant cross section will be transformed into the elastic cross section. Thus no resonances will be propagated explicitly, but they show up in the cross section We set $\text{sigma_elast} = \text{sigma_Res}$, $\text{sigma_Res} = 0$, $\text{sigma_tot} = \text{sigma_elast}$ please note: background processes as $\text{pi pi} \leftrightarrow \text{K K}^{\sim}$ are *not* affected by this switch. You have to disable those additionally by hand, see <code>mesMes_do2to2</code> |
| omega_K_factor | real | 2. | Modification factor for the inelastic omega-nucleon cross section. Necessary to describe transparency ratio data measured by CBELSA/TAPS, see: http://arxiv.org/abs/1210.3074 |
| coulombCorrect | logical | .false. | Since the new particles are initialized at new positions, also the total coulomb energy might change. If .true. than this is taken into account and some correction to $\text{sqrt}(s)$ is done. NOTES Should only be used if the new finalstate particles are initialized in the middle of the two initial state particles! According to OB, this parameter, which switches on/off the usage of the routine "CoulombDifference", has more or less some nostalgic reasons. Better not to use it nowadays anymore! |
| mesMes_do2to2 | logical | .true. | flag whether to do $m m' \leftrightarrow \text{K K}^{\sim}$, $\text{K K}^{*\sim}$ etc. |
| mesMes_useWidth | logical | .false. | flag whether to use the width in $m m' \leftrightarrow \text{K K}^{\sim}$, $\text{K K}^{*\sim}$ etc. This is needed to enforce detailed balance. Otherwise only pole masses are used. |
| NN_inmed | logical | .true. | flag for using the Li-Machleidt (PR C 48 (1993) 1702, Eq 15 , PR C 49 (1994) Eq 1) attenuation of NN cross sections in medium. |

| | | | |
|---------------------|--|---------|--|
| master_3body | code/collisions/threeBodyReactions/masterThreeBody.f90 | | |
| correctEnergy | logical | .true. | Scale final state momenta to fulfill energy and momentum conservation. If .false., energy conservation is violated. |
| radiusNukSearch | real | 2.9 | Radius for the search of nucleons, i.e. the radius in which nucleons shall be searched for at rho_0. |
| deltaThreeBody | | | |
| pionThreeBody | logical | .true. | Switch for the NNpion \rightarrow NN processes (false=OFF). |
| positionNNpi | logical | .false. | This switch determines where the final state particles in NNpi \rightarrow NN are positioned: <ul style="list-style-type: none"> • true: pion position • false: center of NNpi (default) |

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|------------------------|--------------------------------------|---------|--|
| MatrixElementQE | code/init/lepton/matrixElementQE.f90 | | |
| useQEextraterm | logical | .true. | switch on/off an extra term appearing in the current due to different masses of in- and outgoing nucleons |
| useCorrelations | logical | .false. | switch on/off RPA correlations according to Nieves, Amaro, Valverde, PRC70, 055503 (2004) |
| nievesCorr_para | integer | 2 | if RPA correlations are switched on, this parameter decides which set of variables to use: <ul style="list-style-type: none"> • 1: modified Nieves et al., PRC70, 055503 (2004) • 2: original Nieves et al., PRC70, 055503 (2004) • 3: Tselyaev, Speth et al., PRC75, 014315 (2007) |
| gp | real | 0.63 | vary gp if RPA correlations are switched on |
| withScalarInt | logical | .true. | switch on/off scalar interactions |

| | | | |
|---------------------|-------------------------|-------|--|
| MediumModule | code/density/medium.f90 | | |
| mediumCutOff | real | 1.E-8 | If the density is lower than this value, then we treat the medium like vacuum. |

| mesonPotential | | code/potential/mesonPotential.f90 | |
|-------------------------|---------|-----------------------------------|--|
| pionPot_Switch | integer | 0 | Switch for pion potential: <ul style="list-style-type: none"> • 0 = no potential • 1 = Oset potential (NPA 554), which is valid up to 50 MeV kinetic energy • 2 = Kapusta suggestion for pion potential (rather unusual) • 3 = Delta-Hole potential, which is valid up to 130 MeV kinetic energy • 4 = Smooth spline transition between switch 1 and 3. NOTES Can be set in namelist mesonPotential. |
| noPerturbativePotential | logical | .false. | Switch for potential of perturbative particles. If .true. then perturbative mesons feel no potential. NOTES Can be set in namelist mesonPotential. |
| vectorMesonPot | integer | 0 | Switch for medium-modification of vector mesons: <ul style="list-style-type: none"> • 0 = no modification • 1 = Brown-Rho-Scaling • 2 = Brown-Rho-Scaling with momentum dependence according to Kondratyuk (see page 162 in Effenberger's thesis). Currently not available! NOTES Can be set in namelist mesonPotential. |
| brownRho | real | 0.16 | Brown-Rho scaling parameter alpha. |

| MesonWidthVacuum | | code/width/mesonWidthVacuum.f90 | |
|-------------------------|---------|---------------------------------|--|
| omega_width | integer | 1 | Select a parametrization for the omega vacuum width: <ul style="list-style-type: none"> • 1 = GiBUU default (a la Manley) • 2 = Muehlich |

| ModifyParticles | | |
|------------------------|---|--|
| | code/database/particleProperties.f90 | |
| mass | real, -1.0 dimension(1:pion+nMes-1) | Input array for modifications on the particle mass NOTES This array is intended to "input" values for the mass of the particles, which are different from the default. Therefore only entries, which are positive after reading the file are stored in the internal database. |
| width | real, -1.0 dimension(1:pion+nMes-1) | Input array for modifications on the particle width NOTES This array is intended to "input" values for the width of the particles, which are different from the default. Therefore only entries, which are positive after reading the file are stored in the internal database. |
| stabilityFlag | integer, -1 dimension(1:pion+nMes-1) | Input array for modifications on the particle stability NOTES This array is intended to "input" values for the stability of the particles, which are different from the default. Therefore only entries, which are >-1 after reading the file are stored in the internal database. The index of the array is the particle ID. The value encodes on a bitwise level, how the particle may decay (cf. also master.lBody): <ul style="list-style-type: none"> • 1: particle may decay during run, if $\Gamma > \text{gammaCutOff}$ • 2: particle may decay at the end of the run, if $\Gamma > 0$. • 4: particle may decay at the end via Jetset, if there the parameters allow for a decay. The default values are one of the following: <ul style="list-style-type: none"> • 0: particle may not decay at all (i.e. it is stable) • 3: particle may decay both during run and at the end (combination of 1 and 2) |

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| neutrino_induced | <code>code/init/neutrino/initNeutrino.f90</code> | | |
|-------------------------|--|--|--|

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|------------|---------|---|---|
| process_ID | integer | 2 | Determine the process (cf. module leptonicID): |
| | | | <ul style="list-style-type: none"> • 1 = EM • 2 = CC • 3 = NC • -1 = antiEM • -2 = antiCC • -3 = antiNC |

| | | | |
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| flavor_ID | integer | 2 | Determine the lepton flavor: |
| | | | <ul style="list-style-type: none"> • 1 = electron • 2 = muon • 3 = tau |

| | | | |
|----------------|---------|---|---|
| nuXsectionMode | integer | 0 | <p>To choose which kind of Xsection is calculated. All values set in module neutrino_IDTable.f90 possible values:</p> <ul style="list-style-type: none"> • 0 = integratedSigma: required input: enu • 1 = dSigmaCosThetaElepton: required input: enu, costheta, elepton • 2 = dSigmaQsdElepton: required input: enu, Qs, elepton • 3 = dSigmaQs: required input: enu, Qs • 4 = dSigmaCosTheta: required input: enu, costheta • 5 = dSigmaElepton: required input: enu, elepton • 6 = dSigmaMC: required input: enu • 7 = dSigmaW: required input: enu, W <p>calculation for specific experiments taking into account the flux (choose your favorite experiment with flag nuExp):</p> <ul style="list-style-type: none"> • 10 = EXP_dSigmaEnu • 11 = EXP_dSigmaCosThetaElepton • 12 = EXP_dSigmaQsdElepton • 13 = EXP_dSigmaQs • 14 = EXP_dSigmaCosTheta • 15 = EXP_dSigmaElepton • 16 = EXP_dSigmaMC • 17 = EXP_dSigmaW |
|----------------|---------|---|---|

| | | | |
|-----------|---------|--------|---|
| nuExp | integer | 0 | <ul style="list-style-type: none"> • 0 = no specific experiment • 1 = MiniBooNE neutrino flux (in neutrino mode = positive polarity) • 2 = ANL • 3 = K2K • 4 = BNL • 5 = MiniBooNE antineutrino flux (in antineutrino mode = negative polarity) • 6 = MINOS muon-neutrino in neutrino mode • 7 = MINOS muon-antineutrino in neutrino mode • 8 = NOVA neutrino (medium energy NuMI, 14 mrad off-axis), FD • 9 = T2K neutrino off-axis 2.5 degrees (at ND280 detector) • 10 = uniform distribution from Eflux_min to Eflux_max (see namelist nl_neutrino.energyFlux in the module expNeutrinoFluxes) • 11 = MINOS muon-neutrino in antineutrino mode • 12 = MINOS muon-antineutrino in antineutrino mode • 13 = MINERvA muon neutrino, old flux • 14 = MINERvA muon antineutrino, old flux • 15 = LBNF/DUNE in neutrino mode • 16 = LBNF/DUNE in antineutrino mode • 17 = LBNO neutrino in neutrino mode • 18 = NOMAD • 19 = BNB nue BNB= Booster Neutrino Beam • 20 = BNB nuebar • 21 = BNB numu • 22 = BNB numubar • 23 = NOvA ND • 24 = T2K on axis • 25 = MINERvA, 2016 flux • 99 = user provided input file |
| includeQE | logical | .true. | include QE scattering |

| | | | |
|-------------------|------------------|---------|---|
| includeDELTA | logical | .true. | include Delta excitation |
| includeRES | logical | .true. | include excitation of higher resonances |
| include1pi | logical | .false. | include one-pion cross section see neutrinoXsection.f90 for details: there one might choose between different models and also whether it is taken as background or as total cross section |
| include2pi | logical | .false. | include 2 pion background channel |
| includeDIS | logical | .false. | include DIS contribution |
| include2p2hQE | logical | .false. | include 2p2h QE contribution |
| include2p2hDelta | logical | .false. | include 2p2h Delta contribution |
| sigmacut | real | 10e-4 | events with a cross section smaller than this value are skipped. |
| realRun | logical | .false. | Do not initialize the final state particles as perturbative particles but as real ones. |
| printAbsorptionXS | logical | .false. | flag to produce output about inclusive (absorption) cross sections |
| FileNameFlux | character(1000)" | | The absolute filename of the file containing flux info, if user supplied possible values: <ul style="list-style-type: none"> • if given, but does not contain '/': default is '[path.To.Input]/[FileNameFlux]' • otherwise: filename is absolute, including path ('~' is okay) NOTE if you want to use the file 'XXX.dat' in the actual directory, give it as './XXX.dat' |
| storeNucleon | integer | 2 | indicate which kind of struck nucleon to save: <ul style="list-style-type: none"> • 1: free Nucleon (i.e. potential removed) • 2: bound nucleon NOTES real check of energy and momentum conservation only possible with '2' |

| | | | |
|----------------------------|---|-----|---|
| neutrino_MAIDlikeBG | code/init/neutrino/singlePionProductionMAIDlike.f90 | | |
| b_proton_pinull | real | 3. | strength of 1pi BG for CC, multiplies EM BG 3. is tuned to ANL, 6. is tuned to BNL |
| b_neutron_piplus | real | 1.5 | strength of 1pi BG for CC, multiplies EM BG 1.5 is tuned to ANL, 3. is tuned to BNL |

| | | | |
|-------------------------------|--------------------------------------|---|--|
| neutrino_matrixelement | code/init/neutrino/matrixelement.f90 | | |
| which_resonanceModel | integer | 0 | to change between different realizations of the matrix elements: |
| | | | <ul style="list-style-type: none"> • 0 = with Fortran calculated matrix elements containing all resonances (default) • 1 = with Mathematica calculated matrix elements (only Delta) • 2 = Rein and Sehgal's matrix elements |

| | | | |
|---|------------------------------------|---------|--|
| NeutrinoAnalysis | code/analysis/neutrinoAnalysis.f90 | | |
| detailed_diff_output | logical | .false. | If .true. then also the detailed output of differential cross sections is produced |
| include_W_dist | logical | .false. | If .true. then the invariant mass distributions for events with 1 pion and 1 nucleon in the final state are produced |
| kineticEnergyDetectionThreshold_lepton | real | 0.0 | kineticEnergyDetectionThreshold only lepton kinetic energies above this threshold can be detected This cut affects *all* events, not just the outgoing lepton! |
| AngleUpperDetectionThresholdDegrees_lepton | real | 180.0 | lepton angles up to this value can be detected This cut affects *all* events, not just the outgoing lepton! |
| kineticEnergyDetectionThreshold_nucleon | real | 0.0 | kineticEnergyDetectionThreshold lower detection threshold for nucleon kinetic energies |
| AngleUpperDetectionThresholdDegrees_nucleon | real | 180.0 | nucleon angles up to this value can be detected |
| kineticEnergyDetectionThreshold_chargedpion | real | 0.0 | kineticEnergyDetectionThreshold |
| AngleUpperDetectionThresholdDegrees_chargedpion | real | 180.0 | charged pion angles up to this value can be detected |
| kineticEnergyDetectionThreshold_neutralpion | real | 0.0 | kineticEnergyDetectionThreshold |
| AngleUpperDetectionThresholdDegrees_neutralpion | real | 180.0 | neutral pion angles up to this value can be detected |
| inclusiveAnalysis | logical | .false. | If .true. then we don't care whether particle has left the nucleus or not |
| Fissum_analysis | logical | .false. | do analysis with cuts as needed for Fig 25 in Fissum et al, PRC 70, 034606 (2004) |
| ZeroPion_analysis | logical | .false. | produce output of xsec for various final states with 0 pions and 2 pions see file see sigma_0pions.dat for the list of the final states see files neutrino_0pions.dat, neutrino_0pions_QE.dat, neutrino_0pions_Delta.dat, ... for output |
| calorimetric_analysis | logical | .false. | do calorimetric energy-transfer and neutrino-energy reconstruction (for each QE, Delta, ...) as in the MINOS experiment |

| | | | |
|-----------------------------|---------|---------|---|
| radialScale | real | 0.0 | If radial position of nucleon < radialScale*target radius, then the nucleon is assumed to be bound |
| reconstruct_neutrino_energy | logical | .false. | reconstruct neutrino energy for final state in "specificEvent_analysis" NOTES .true. must be combined with specificEvent_analysis=.true. and at least one specific event .true. |
| outputEvents | logical | .false. | If .true. then all events are printed to the file 'FinalEvents.dat'. |
| specificEvent_analysis | logical | .false. | do analysis for specific final states values can be changed in the namelist nl_specificEvent |
| Xsection_analysis | logical | .false. | If .true. then files "..._total_Xsection..." and "..._dSigmadEkin..." are printed. |

| nl_calorimetric_analysis code/analysis/neutrinoAnalysis.f90 | | | |
|--|------|------|---|
| numin | real | 0. | for calorimetric analysis: values for transferred energy; only work if calorimetric_analysis is .true. set the min, max and bins for nu distributions |
| numax | real | 10.0 | for calorimetric analysis: values for transferred energy; only work if calorimetric_analysis is .true. set the min, max and bins for nu distributions |
| nubin | real | 0.1 | for calorimetric analysis: values for transferred energy; only work if calorimetric_analysis is .true. set the min, max and bins for nu distributions |
| Enumin | real | 0. | for calorimetric analysis: values for neutrino energy; only work if calorimetric_analysis is .true. set the min, max and bins for nu distributions |
| Enumax | real | 10.0 | for calorimetric analysis: values for neutrino energy; only work if calorimetric_analysis is .true. set the min, max and bins for nu distributions |
| Enubin | real | 0.1 | for calorimetric analysis: values for neutrino energy; only work if calorimetric_analysis is .true. set the min, max and bins for nu distributions |

| nl_dSigmadcostheta code/init/neutrino/neutrinoXsection.f90 | | | |
|---|------|------|--|
| enu | real | -10. | neutrino energy, read in by namelist |
| costheta | real | -10. | cosine of the angle between the neutrino (z-direction) and the outgoing lepton |
| delta_costheta | real | -10. | value by which costheta is increased |

| nl_dSigmaDCosThetadElepton <small>code/init/nuetrino/nuetrinoXsection.f90</small> | | | |
|--|------|------|--|
| enu | real | -10. | neutrino energy, read in by namelist |
| costheta | real | -10. | cosine of the angle between the neutrino (z-direction) and the outgoing lepton |
| elepton | real | -10. | energy of the outgoing lepton |
| delta_elepton | real | -10. | value by which elepton is increased |

| nl_dSigmaDElepton <small>code/init/nuetrino/nuetrinoXsection.f90</small> | | | |
|---|------|------|--------------------------------------|
| enu | real | -10. | neutrino energy, read in by namelist |
| elepton | real | -10. | energy of the outgoing lepton |
| delta_elepton | real | -10. | value by which elepton is increased |

| nl_dSigmaDQs <small>code/init/nuetrino/nuetrinoXsection.f90</small> | | | |
|--|------|------|--------------------------------------|
| enu | real | -10. | neutrino energy, read in by namelist |
| Qs | real | -10. | momentum transfer squared |
| delta_Qs | real | -10. | value by which Qs is increased |

| nl_dSigmaDQsdElepton <small>code/init/nuetrino/nuetrinoXsection.f90</small> | | | |
|--|------|------|--------------------------------------|
| enu | real | -10. | neutrino energy, read in by namelist |
| Qs | real | -10. | momentum transfer squared |
| elepton | real | -10. | energy of the outgoing lepton |
| delta_elepton | real | -10. | value by which elepton is increased |

| nl_dSigmaDW <small>code/init/nuetrino/nuetrinoXsection.f90</small> | | | |
|---|------|------|--------------------------------------|
| enu | real | -10. | neutrino energy, read in by namelist |
| W | real | -10. | invariant mass defined as $(p+q)^2$ |
| delta_W | real | -10. | value by which W is increased |

| nl_fluxcuts <small>code/init/nuetrino/esample.f90</small> | | | |
|--|---------|---------|--|
| Enu_lower_cut | real | 0. | cut events with neutrino energy below Enu_lower_cut; for ANL experiment, for example, Enu_lower_cut=0.5 for some analysis of ppi+ |
| Enu_upper_cut | real | 200. | cut events with neutrino energy above Enu_upper_cut; for ANL experiment, for example, Enu_upper_cut=1.5 for ppi0 and npi+ final state, but 5.98 for ppi+ |
| energylimit_for_Qsrec | logical | .false. | switch for using the energylimits Enu_upper_cut and Enu_lower_cut for the Q ² reconstruction; values: .true. or .false. (default: .false.) |

| nl_integratedSigma <small>code/init/nuetrino/nuetrinoXsection.f90</small> | | | |
|--|------|------|---|
| enu | real | -10. | neutrino energy, read in by namelist |
| delta_enu | real | -10. | value by which the neutrino energy is increased |

| nl_neutrino_energyFlux <small>code/init/nuetrino/expNeutrinofluxes.f90</small> | | | |
|---|------|-------|---|
| Eb | real | 0.034 | contant binding energy used for energy and Q2 reconstruction based on QE scattering kinematics |
| Eflux_min | real | 0.2 | minimum energy for uniform flux distribution minimum and maximum energies for the uniform neutrino flux (nu-Exp=10 in the namelist neutrino_induced) can be changed in the namelist nl_neutrino_energyFlux |
| Eflux_max | real | 2.5 | maximum energy for uniform flux distribution minimum and maximum energies for the uniform neutrino flux (nu-Exp=10 in the namelist neutrino_induced) can be changed in the namelist nl_neutrino_energyFlux |

| nl_neutrinoxsection <small>code/init/nuetrino/nuetrinoXsection.f90</small> | | | |
|---|---------|-----|---|
| integralPrecision | integer | 3 | precision for the Gauss integration (reduce it for nuXsectionMode.eq.0 (sigma) to e.g. 2) |
| integralPrecisionQE | integer | 500 | precision for the Gauss integration over the QE peak (reduce it for nuXsection-Mode.eq.0 (sigma) to e.g. 300) |
| singlePiModel | integer | 1 | to change between different models for the pion nucleon cross section: <ul style="list-style-type: none"> • 0 = pi N according to Nieves et al (hep-ph/0701149) • 1 = MAID-like model |

| | | | |
|---------------------|---------|--------|--|
| invariantMassCut | real | 100. | cut events with invariant Mass above this value (in GeV); cut pion production from Delta and DIS on $W_{rec} = \sqrt{M^2 + 2*M*\nu - Q^2}$ |
| invariantMassCut_BG | real | 100. | cut MAID-like background events with invariantMass_BG above this value (in GeV); cut 1pi BG on $W_{rec} = \sqrt{M^2 + 2*M*\nu - Q^2}$ |
| DIScutW1 | real | 1.55 | lower W-cut for linear onset of DIS |
| DIScutW2 | real | 1.65 | upper W-cut for linear onset of DIS |
| REScutW1 | real | 1.90 | lower W-cut for linear turn-off of resonances |
| REScutW2 | real | 2.00 | upper W-cut for linear turn-off or resonances |
| DISformfakEM | integer | 2 | Introduce an additional form factor for the DIS cross section, when processed via a photon: <ul style="list-style-type: none"> • 0: no form factor • 1: $Q^2/(mcutDIS^2+Q^2)$ • 2: $Q^4/(mcutDIS^2+Q^2)^2$ In case of electron induced events, we need choose 2 in order to be compatible with Pythia's electron machinery. |
| DISformfakNCCC | integer | 1 | Introduce an additional form factor for the DIS cross section, when processed via W or Z boson: <ul style="list-style-type: none"> • 0: no form factor • 1: $Q^2/(mcutDIS^2+Q^2)$ • 2: $Q^4/(mcutDIS^2+Q^2)^2$ In case of electron induced events, we need choose 2 in order to be compatible with Pythia's electron machinery. |
| mcutDIS | real | 0.5 | parameter to control Q^2 dependence of DIS |
| Q2cut | real | 1.0 | parameter to control Q^2 dependence of DIS, DIS is cut out for all $Q^2 < Q2cut$ in the resonance region |
| dQ2 | real | 0.1 | parameter to control sharpness of DIS-cutoff Q2cut |
| DISrespectHad | logical | .true. | Flag to indicate, whether hadronization failures should be respected and affect the overall DIS cross section Pythia is run to generate the DIS cross section. But not every of the generated events may lead to a correct hadronic final state. |
| DISdoMSTP23 | logical | .true. | Flag to indicate, whether in Pythia for neutrino-DIS the value MSTP(23)=1 should be used or not |

| nl_NievesHadronTensor | | |
|------------------------------|--|--------|
| | <i>code/init/neutrino/NievesHadronTensor.f90</i> | |
| DeltaPole | logical | .true. |
| crossedDelta | logical | .true. |
| nucleonPole | logical | .true. |
| crossedNucleonPole | logical | .true. |
| contactTerm | logical | .true. |
| pionPole | logical | .true. |
| pionInFlight | logical | .true. |

| nl_SigmaMC | | | |
|-------------------|--|------|--------------------------------------|
| | <i>code/init/neutrino/neutrinoXsection.f90</i> | | |
| enu | real | -10. | neutrino energy, read in by namelist |
| MC_xmax | real | 2.0 | |

| | | | |
|---------------------------------------|--|---|--|
| nl_singlePionProductionNHVlike | code/init/neutrino/singlePionProductionNHVlike.f90 | | |
| integrate_over | integer | 2 | possible values: <ul style="list-style-type: none"> • 1 = costhetaPi • 2 = Epi • 3 = over cosThetaPi_star_qz in CM frame <p>which 3-pl differential cross sectio to use for integration:</p> <ul style="list-style-type: none"> • 1= dsigma/dcostheta/dElepton/dcosThetaPion was originally used and works for nuclei. disadvantage: for some cosThetaPion there are two solutions for Epi, this leads to fluctuations on the cross section • 2= dsigma/dcostheta/dElepton/dEPion has an advantage, that for a given pion energy there is only one solution for the angle between the resonance and pion momenta. so the integration is simpler and results should be smoother <p>NOTES</p> <ul style="list-style-type: none"> • for 1 : the only option checked for nucleus • for 2 : code works better and faster, gives significantly smoother results below Delta peak. disadvantage: now for the free nucleon only, TO DO : nuclei |

| | | | |
|-------------------------|------------------------------------|---------|--|
| nl_specificEvent | code/analysis/neutrinoAnalysis.f90 | | |
| no_pi | logical | .false. | do analysis for specific final states: specificEvent=1, no_pi (for example, for QE-like MiniBooNE) values can be changed in the namelist nl_specificEvent |
| p_Xn_no_pi | logical | .false. | do analysis for specific final states: specificEvent=2 values can be changed in the namelist nl_specificEvent |
| piplus | logical | .false. | do analysis for specific final states: specificEvent=3, 1 pi+ X nucleons mesons of other flavor values can be changed in the namelist nl_specificEvent |
| piplus_MULTI | logical | .false. | do analysis for specific final states: specificEvent=4 >=1 pi+ X other pions (incl pi+) X nucleons values can be changed in the namelist nl_specificEvent |

| | | | |
|---------------|---------|---------|---|
| pi0 | logical | .false. | do analysis for specific final states: specificEvent=5, 1 pi0 X nucleons, plus mesons of other flavor values can be changed in the namelist nl_specificEvent |
| pi0_MULTI | logical | .false. | do analysis for specific final states: specificEvent=6, >=1 pi0 X other pions X nucleons, (pi0 K2K) values can be changed in the namelist nl_specificEvent |
| piminus | logical | .false. | do analysis for specific final states: specificEvent=7 1 pi- X other pions X nucleons values can be changed in the namelist nl_specificEvent |
| piminus_MULTI | logical | .false. | do analysis for specific final states: specificEvent=8 >=1 pi- X other pions X nucleons values can be changed in the namelist nl_specificEvent |
| pp_no_pi | logical | .false. | do analysis for specific final states: specificEvent=9 2 protons, X neutrons, 0 pions values can be changed in the namelist nl_specificEvent |
| pn_no_pi | logical | .false. | do analysis for specific final states: specificEvent=10 1 neutron, 1 proton, 0 pions values can be changed in the namelist nl_specificEvent |
| nn_no_pi | logical | .false. | do analysis for specific final states: specificEvent=11 2 neutrons, X protons, 0 pions values can be changed in the namelist nl_specificEvent |
| pp_Xn_no_pi | logical | .false. | do analysis for specific final states: specificEvent=12 2 protons, X neutrons, 0 pions values can be changed in the namelist nl_specificEvent |
| nn_Xp_no_pi | logical | .false. | do analysis for specific final states: specificEvent=13 2 neutrons, X protons, 0 pions values can be changed in the namelist nl_specificEvent |
| ppp_Xn_no_pi | logical | .false. | do analysis for specific final states: specificEvent=14 3 protons, X neutrons, 0 pions values can be changed in the namelist nl_specificEvent |
| pppp_Xn_no_pi | logical | .false. | do analysis for specific final states: specificEvent=15 4 protons, X neutrons, 0 pions values can be changed in the namelist nl_specificEvent |
| p_no_pi | logical | .false. | do analysis for specific final states: specificEvent=16 1 proton, 0 neutron, 0 pion values can be changed in the namelist nl_specificEvent |

| | | | |
|-------------|---------|---------|---|
| n_no_pi | logical | .false. | do analysis for specific final states: specificEvent=17 1 neutron, 0 proton, 0 pion values can be changed in the namelist nl_specificEvent |
| Xn_no_pi | logical | .false. | do analysis for specific final states: specificEvent=18, 0 proton, X neutrons, 0 pions values can be changed in the namelist nl_specificEvent |
| binsizeQ2 | real | 0.01 | do analysis for specific final states: binning for reconstruction of Q2 and Enu values can be changed in the namelist nl_specificEvent |
| binsizeEnu | real | 0.02 | do analysis for specific final states: binning for reconstruction of Q2 and Enu values can be changed in the namelist nl_specificEvent |
| maxQ2 | real | 5.0 | do analysis for specific final states: max values for reconstruction of Q2 and Enu values can be changed in the namelist nl_specificEvent |
| maxEnu | real | 5.0 | do analysis for specific final states: max values for reconstruction of Q2 and Enu values can be changed in the namelist nl_specificEvent |
| excl_hadron | logical | .false. | do analysis for specific final states: specificEvent=19,20,21 exclusive 1 pion, no other pions or other mesons of different flavor There could be still other mesons which are heavier than the D, Such events (very rare at DUNE energies) could be counted as exclusive single-meson cross section. This could be cured by extending the list of stable mesons value can be changed in the namelist nl_specificEvent |
| QEp | logical | .false. | if .true, do analysis for specific analysis for QE-like event with 1 mu, 0 pi, X p values can be changed in the namelist nl_specificEvent |
| full_incl | logical | .true. | do analysis for specific final states: specificEvent=22 fully inclusive event, all hadrons in final state value can be changed in the namelist nl_specificEvent |

| OffShellPotential | | | |
|-----------------------------|----------------------------------|---------|---|
| | code/width/offShellPotential.f90 | | |
| useOffShellPotentialBaryons | logical | .false. | Switch on or off whether the offshellness should be used for baryons. NOTES <ul style="list-style-type: none"> • must be set to "TRUE" if mediumSwitch_coll (see module Baryon-WidthMedium) is .true. • if .true. then delta_T (see module inputGeneral) must be ≤ 0.05 AND delta_P (see module propagation) must be ≤ 0.002; AND delta_E (see module propagation) must be ≤ 0.002; slows down propagation by a factor of 10 |
| useOffShellPotentialMesons | logical | .false. | Switch on or off whether the offshellness should be used for mesons. |
| extrapolateBaryonWidth | logical | .true. | Whether to extrapolate the baryon width below minimal mass or not. |
| max_offshellparameter | real | 5. | The maximal value for the offshell parameter. Note: empirical value! This only applies to baryons. For mesons we have no restrictions on the offshell parameter. |
| relativistic | logical | .false. | <ul style="list-style-type: none"> • false: Use non-rel. off-shell parameter $x = \Delta m / \Gamma$, which obeys Stefan Leupold's non-rel. EOM. • true: Use rel. off-shell parameter $x = \Delta m^2 / \Gamma$, which obeys Cassing's rel. EOM. |

| paramEP | | | |
|----------------|-----------------------------------|---|--|
| | code/init/lowElectron/ParamEP.f90 | | |
| useParam | integer | 2 | select, which parametrization to use: <ul style="list-style-type: none"> • 1: Brasse • 2: Bosted |

| photonXS | | | |
|-----------------|--|--------|--|
| | code/collisions/twoBodyReactions/HiEnergy/photonXS.f90 | | |
| iParam | integer | 2 | Switch to select the kind of parametrization for $\gamma N \rightarrow V N$: <ul style="list-style-type: none"> • 1: "old parametrization", fit to experimental data, cf. Effenberger PhD, p.53 • 2: Pythia, cf. Friberg/Sjstrand hep-ph/0007314 • 3: Donnachie, Landshoff [citation needed] |
| omega_saphir | logical | .true. | If .true. an improved fit (to SAPHIR data) will be used for $\gamma N \rightarrow \omega N$. cf. "calcXS_omega_saphir" |

| pionAnalysis code/analysis/pionXsection.f90 | | | |
|--|---------|---------|--|
| CMFrame | logical | .false. | If .true. Xsection is evaluated in CM-Frame of the incoming pion and a resting nucleon, else in calculation frame. |
| dsigma_dOmegadE_switch | logical | .false. | If .true. then dsigma/dOmega and dSigma/dOmega/dE are evaluated. |
| twoPi_switch | logical | .false. | If .true. then 2Pi output is evaluated. |

| pionNucleus code/init/initPion.f90 | | | |
|---|---------|---------|---|
| UseCoulomb | logical | .false. | if .true. then a Coulomb propagation from CoulombDistance to distance is performed |
| CoulombDistance | real | 200. | distance from where the Coulomb propagation starts |
| distance | real | 15. | initialization distance |
| impact_parameter | real | 0. | impact parameter. If less than 0, than an impact parameter integration is performed |
| charge | integer | 0 | charge of pion |
| numberPions | integer | 200 | number of initialized pions per ensemble |
| ekin_lab | real | 0. | kinetic energies of pions in lab frame. |
| delta_ekin_lab | real | 0.01 | step size for kinetic energies in energy scans |

| pn_medium code/width/proton_neutron_width_medium.f90 | | | |
|---|---------|---------|---|
| density_dependent | logical | .false. | the density of the spectral function |
| pn_medium_switch | logical | .true. | If .true. medium_modifications will be used |
| form_factor | logical | .true. | If .true. the form factor for the width is used |

| projectile code/density/nucleus.f90 | | | |
|--|---------|--------|--|
| Projectile_A | integer | 0 | Mass A of projectile nucleus (= number of nucleons). If zero, a default isotope is chosen for the given projectile_Z. |
| Projectile_Z | integer | 20 | Charge Z of projectile nucleus (= number of protons). |
| fermiMotion | logical | .true. | Determines whether particles in target nucleus feel Fermi motion or not. |

| | | | |
|----------------------|---------|-------|---|
| densitySwitch_static | integer | 3 | <p>This switch is important, because it decides, which static density is used to set up the testparticles in the nuclei before the first time-step.</p> <p>Possible values:</p> <ul style="list-style-type: none"> • 0 : density=0.0 • 1 : Static density uses Woods-Saxon according to H. Lenske • 2 : Static density according to NPA 554 • 3 : Static density according to Horst Lenske, implements different radii for neutrons and protons • 4 : Static density according oscillator shell model • 5 : Density distribution is a sphere with density according to the input value of "fermiMomentum_input". • 6 : Static Density based on LDA, implemented by Birger Steinmueller • 7 : Static Density based on LDA + Welke potential • 8 : Static Density prescription according Relativistic Thomas-Fermi (Valid only in RMF-mode) <p>Possible nuclei for the different prescriptions:</p> <ul style="list-style-type: none"> • 1 : $A > 2$ (only $A > 16$ makes sense) • 2 : • 3 : 6→C(12), 8→O(16),O(18), 13→Al(27), 20→Ca(40),Ca(44), 79→Au(197) 82→Pb(208) • 4: 2→He(4), 4→Be(9), 5→B(11), 6→C(12), 8→O(16) |
| fermiMomentum_input | real | 0.225 | Input value of the fermi momentum for densitySwitch_static=5 (in GeV). |

| Propagation | | | |
|--------------------|----------------------------------|--------|---|
| | code/propagation/propagation.f90 | | |
| delta_P | real | 0.01 | Delta Momentum in derivatives |
| delta_E | real | 0.01 | Delta energy in derivatives |
| UseCoulombDirectly | logical | .true. | Whether to use coulomb force directly in propagation or not. (If switched off while coulomb is switched on in module coulomb, the effect of the coulomb potential comes in via the gradient of the potentials. With this flag you can not switch on/off coulomb, you just select, how it is treated.) |
| UseHadronic | logical | .true. | Whether to use hadronic potentials in propagation |

| | | | |
|--------------------|---------|---------|--|
| FreezeNonint | logical | .false. | If switched on, the real particles which did not interact will have zero velocities, i.e. will be "frozen". This is important for stability of the nuclear ground state in real particle simulations. Note that this flag influences only when freezeRealParticles=.false. |
| RungeKuttaOrder | integer | 1 | Order of Runge-Kutta in derivatives: <ul style="list-style-type: none"> • 1 = first order Runge-Kutta • 2 = second order Runge-Kuttay |
| Mode | integer | 2 | define the type of propagation: <ul style="list-style-type: none"> • 0: Cascade • 1: Euler • 2: PredictorCorrector |
| dh_dp0_switch | logical | .true. | Switch which decides whether we use dh_dp0. |
| offShellInfoDetail | logical | .false. | print out detailed offShellInfo |
| tachyonDebug | logical | .false. | ... |

propagation_RMF_input

code/propagation/propagation_RMF.f90

| | | | |
|--------------------|---------|--------|--|
| predictorCorrector | logical | .true. | Switch for predictor-corrector method in the propagation. If .false. then simple Euler method is used (i.e. only predictor step is done) |
|--------------------|---------|--------|--|

pythia

code/collisions/twoBodyReactions/HiEnergy/DoCollTools.f90

| | | |
|------|---------------------------------|-----------------|
| MSEL | integer | Pythia variable |
| MSTU | integer, dimen- sion(200) | Pythia array |
| MSTJ | integer, dimen- sion(200) | Pythia array |
| MSTP | integer, dimen- sion(200) | Pythia array |

| | | |
|------|---------------------------------------|--------------|
| MSTI | integer, dimen- sion(200) | Pythia array |
| PARU | real, dimen- sion(200) | Pythia array |
| PARJ | real, dimen- sion(200) | Pythia array |
| PARP | real, dimen- sion(200) | Pythia array |
| PARI | real, dimen- sion(200) | Pythia array |
| CKIN | real, dimen- sion(200) | Pythia array |
| PMAS | real, dimen- sion(500, 4) | Pythia array |
| MDCY | integer, dimen- sion(500, 3) | Pythia array |

| | | | |
|----------------------|--|---------|---|
| residue_input | code/analysis/sourceAnalysis/residue.f90 | | |
| DetermineResidue | logical | .false. | If .true., then the determination of the target residue properties for every event and their output in file TargetResidue.dat at the end of time evolution will be done |

| | | | |
|-------------------------------|---|---------|---|
| ResonanceCrossSections | code/collisions/twoBodyReactions/baryonMeson/resonanceCrossSections.f90 | | |
| fullPropagator | logical | .false. | Includes also the real parts in the resonance propagator. In former works (i.e. in the old Effenberger code) this has been neglected. It should be set to .true. only if mediumSwitch_coll=.true. in the namelist width_Baryon. |

| | | | |
|------------------|------------------|---------|--|
| RMF_input | code/rmf/RMF.f90 | | |
| RMF_flag | logical | .false. | If .true. then use relativistic mean fields. |

| | | | |
|-----------------|---------|---------|--|
| N_set | integer | 1 | Select which parameter set to use: <ul style="list-style-type: none"> • 1 — NL1 from G.A. Lalazissis et al., PRC 55, 540 (1997), (K=211.29 MeV, $m^*/m=0.57$) • 2 — NL3 from G.A. Lalazissis et al., PRC 55, 540 (1997), (K=271.76 MeV, $m^*/m=0.60$) • 3 — NL2 set from A. Lang et al., NPA 541, 507 (1992), (K=210 MeV, $m^*/m=0.83$) • 4 — NLZ2 set from M. Bender et al., PRC 60, 34304 (1999), (K=172 MeV, $m^*/m=0.583$) • 5 — NL3* set from G.A. Lalazissis, private communication, (K=258.28 MeV, $m^*/m=0.594$) • 6 — Same as N_set=3, but including the rho meson. • 7 — NL1 set from S.J. Lee et al., PRL 57, 2916 (1986), (K=212 MeV, $m^*/m=0.57$) • 8 — NL2 set from S.J. Lee et al., PRL 57, 2916 (1986), (K=399 MeV, $m^*/m=0.67$) • 9 — Set I from B. Liu et al., PRC 65, 045201 (2002), (K=240 MeV, $m^*/m=0.75$) |
| grad_flag | logical | .false. | If .true. then include space derivatives of the fields. |
| lorentz_flag | logical | .true. | If .false. then the space components of the omega field are put to zero. |
| fourMomDen_flag | logical | .false. | If .true. then compute the four-momentum density field (not used in propagation). |
| kaonpot_flag | logical | .false. | This switch turns on the Kaon potential in RMF mode. |
| fact_pbar | real | 1. | Modification factor for the antiproton coupling constants. |
| fact_hyp | real | 1. | Modification factor for the hyperon coupling constants. |
| fact_antihyp | real | 1. | Modification factor for the antihyperon coupling constants. |
| fact_Xi | real | 1. | Modification factor for the Xi and XiStar coupling constants. |
| fact_antiXi | real | 1. | Modification factor for the antiXi and antiXiStar coupling constants. |
| fact_kaon | real | 0. | Modification factor for the Kaon and antikaon coupling constants. |

| selfenergy_realPart code/spectralFunctions/selfenergy_baryons.f90 | | | |
|--|---------|---------|---|
| rel_accuracy | real | 0.05 | Relative accuracy for resonance self energy |
| intSolver | integer | 1 | Decide on the numerical package to be used for the Cauchy integral: <ul style="list-style-type: none"> • 1=quadpack routine • 2=cernlib routine |
| makeTable | logical | .true. | Switch on/off the usage of an input tabulation |
| noDispersion | logical | .false. | Switch on/off the usage dispersion relations |
| maxRes | integer | 100 | |
| minRes | integer | -100 | |
| extrapolateAbsP | logical | .false. | if(true) then set absP to maxAbsP if absP is larger |
| writeLocal | logical | .false. | <ul style="list-style-type: none"> • Tables are outputted to local directory, not to buuinput |

| selfEnergyMesons code/spectralFunctions/selfenergy_mesons.f90 | | | |
|--|---------|---------|---|
| dispersion | logical | .false. | Use dispersive real parts of the self energy. |

| SMM_input code/analysis/sourceAnalysis_Main.f90 | | | |
|--|---------|---------|---|
| SMM_Flag | logical | .false. | if .true. then source analysis is switched on |
| rho_cutoff | real | 100. | density cutoff (in units of the saturation density "rhoNull") which defines "emitting" particles |
| spectator_cutoff | real | 1. | min. value of number of collisions which defines "spectator"-matter |
| A_cutoff | integer | 2 | min. value of the source mass number |
| SelectionMethod | integer | 0 | defines the selection method of spectators and fireball. Can be used in high energy Hadron-Nucleus events. |
| betaChoice | integer | 0 | Defines the way to calculate the source velocity in RMF mode. Has no influence in calculations with Skyrme potential. |
| MaxTimePrinting | integer | 10 | Indicates how many times the results are printed into files. NOTES Set MaxTimePrinting to a very big value, i.e. 1000, if you wish that the BUU-run develops until time=time_max. |
| DetailedHyperonOutput | logical | .true. | print more informations for Hyperons and pions. |

| | | | |
|-------------|---------|---------|--|
| hyperSource | logical | .false. | If true, the Lambda and Sigma0 hyperons will be included into source |
|-------------|---------|---------|--|

spectralFunction code/spectralFunctions/spectralFunc.f90

| | | | |
|-----------------|---------|--------|--|
| which_nuclwidth | integer | 1 | This flag decides what is used for the nucleon width. Note: The correct normalisation has not been included here!! Choose between: <ul style="list-style-type: none"> • which_nuclwidth=1 - use constant width given in const_nuclwidth • which_nuclwidth=2 - use width increasing linear with density; $\Gamma = \text{const} \cdot \rho / \rho_0$ with const given in nuclwidth_dens • which_nuclwidth=3 - use toy model (constant NN cross section) • which_nuclwidth=4 - use realistic width (cf. diploma thesis of D. Kalok) • which_nuclwidth=5 - use realistic width: width based on our collision term |
| nuclwidth | real | 0.001 | • if which_nuclwidth=1, nuclwidth gives the width used in the Breit-Wigner for the nucleon |
| nuclwidth_dens | real | 0.006 | • if which_nuclwidth=2, nuclwidth_dens gives the width used in density dependent width |
| nuclwidth_sig | real | 5.5 | • 6 MeV are motivated in F. Froemel dissertation • if which_nuclwidth=3, nuclwidth_sig gives the NN cross section in fm ² |
| relativistic | logical | .true. | • Use either relativistic or non relativistic spectral functions. |

spectralFunctionMesons code/spectralFunctions/spectralFunc_mesons.f90

| | | | |
|--------------|---------|--------|---|
| relativistic | logical | .true. | • Use either relativistic or non relativistic spectral functions. |
|--------------|---------|--------|---|

target code/density/nucleus.f90

| | | | |
|-------------|---------|--------|--|
| Target_A | integer | 0 | Mass A of target nucleus (= number of nucleons). If zero, a default isotope is chosen for the given target_Z. |
| Target_Z | integer | 20 | Charge Z of target nucleus (= number of protons). |
| fermiMotion | logical | .true. | Determines whether particles in target nucleus feel Fermi motion or not. |

| | | | |
|-------------------------|---------|---------|--|
| densitySwitch_static | integer | 3 | <p>This switch is important, because it decides, which static density is used to set up the testparticles in the nuclei before the first time-step. Possible values:</p> <ul style="list-style-type: none"> • 0 : density=0.0 • 1 : Static density uses Woods-Saxon according to H. Lenske • 2 : Static density according to NPA 554 • 3 : Static density according to Horst Lenske, implements different radii for neutrons and protons • 4 : Static density according oscillator shell model • 5 : Density distribution is a sphere with density according to the input value of "fermiMomentum_input". • 6 : Static Density based on LDA, implemented by Birger Steinmueller • 7 : Static Density based on LDA + Welke potential • 8 : Static Density prescription according Relativistic Thomas-Fermi (Valid only in RMF-mode) <p>Possible nuclei for the different prescriptions:</p> <ul style="list-style-type: none"> • 1 : $A > 2$ (only $A > 16$ makes sense) • 2 : • 3 : 6→C(12), 8→O(16),O(18), 13→Al(27), 20→Ca(40),Ca(44), 79→Au(197) 82→Pb(208) • 4: 2→He(4), 4→Be(9), 5→B(11), 6→C(12), 8→O(16) |
| fermiMomentum_input | real | 0.225 | Input value of the fermi momentum for densitySwitch_static=5 (in GeV). |
| ReAdjustForConstBinding | logical | .false. | <p>If this flag is set to true, we use the selected density distribution only for a preliminary step, where we calculate the baryonic potential as function of r (which depends on the density distribution). From the condition, that the binding energy has to be constant, we deduce the distribution of the fermi momentum and thus the 'new' density distribution. The tabulated density distribution is replaced via the 'new' one and all behaviour is as usual.</p> |
| ConstBinding | real | -0.008 | if 'ReAdjustForConstBinding' equals true, we a trying to readjust the fermi momentum and the density such, we quarantee this value for the binding energy. |

| TransportGivenParticle <small>code/init/initTransportGivenParticle.f90</small> | | | |
|---|----------------------|--------------|---|
| particle_ID | integer | 1 | Determines what kind of particle is initialized (see idTable) |
| charge | integer | 1 | Determines what charge |
| position | real, dimension(1:3) | (/0.,0.,0./) | Determines the position. |
| threemomentum | real, dimension(1:3) | (/0.,0.,1./) | Determines the three-momentum. |
| mass | real | -1. | Determines the mass (if negative, choose mass according to spectral function). |
| maxmass | real | 1.5 | Determines the maximum mass (if mass is chosen according to spectral function). |
| perweight | real | 1. | Determines the weight. |
| frequency | integer | 10 | after this amount of time steps a new output file is generated |
| initRandomRadiativeDelta | logical | .false. | intended use: radiativeDelta decay. chooses position,threemomentum,mass of Delta randomly; charge is chosen either 0 or 1 |

| W_distributions <small>code/analysis/neutrinoAnalysis.f90</small> | | | |
|--|------|------|---|
| dW_Npi | real | 0.02 | for dsigma/d(InvariantMass); only work if include_W_dist is .true. set the min, max and steps for various W-distributions |
| Wmin_Npi | real | 1.08 | for dsigma/d(InvariantMass); only work if include_W_dist is .true. set the min, max and steps for various W-distributions |
| Wmax_Npi | real | 1.6 | for dsigma/d(InvariantMass); only work if include_W_dist is .true. set the min, max and steps for various W-distributions |
| dW_muipi | real | 0.04 | only work if include_W_dist is .true. set the min, max and steps for various W-distributions |
| Wmin_muipi | real | 0.24 | only work if include_W_dist is .true. set the min, max and steps for various W-distributions |
| Wmax_muipi | real | 1.2 | only work if include_W_dist is .true. set the min, max and steps for various W-distributions |
| dW_muN | real | 0.04 | only work if include_W_dist is .true. set the min, max and steps for various W-distributions |
| Wmin_muN | real | 1.04 | only work if include_W_dist is .true. set the min, max and steps for various W-distributions |
| Wmax_muN | real | 2.12 | only work if include_W_dist is .true. set the min, max and steps for various W-distributions |

| width_Baryon | | | |
|-----------------------------|----------------------------------|---------|---|
| | code/width/baryonWidthMedium.f90 | | |
| mediumSwitch | logical | .false. | Switch on and off the in-medium width of all baryons at once. If .false., the vacuum width are used. |
| mediumSwitch_Delta | logical | .false. | Only meaningful if mediumSwitch=.true.: Switch on and off the in-medium width of the Delta. (.false.=off) Note that in that case the Delta is treated specially: what is used for the in-medium width is determined by the flag in deltaWidth. This switch is not consistent with mediumSwitch_coll! |
| mediumSwitch_proton_neutron | logical | .false. | Only meaningful if mediumSwitch=.true.: Switch on and off the in-medium width of the proton and the neutron. (.false.=off) Note that in that case the nucleons are treated specially. This switch is not consistent with mediumSwitch_coll! |
| mediumSwitch_coll | logical | .false. | Only meaningful if mediumSwitch=.true.: Use in-medium width according to collision term. NOTES if set to TRUE, then also UseOffShellPotentialBaryons (see module offShellPotential) must be .true. |
| verboseInit | logical | .false. | switch on/off informational messages during initialization |
| verboseInitStop | logical | .false. | Stop after informational messages during initialization or not. |

| width_Meson | | | |
|--------------------|---------------------------------|---------|---|
| | code/width/mesonWidthMedium.f90 | | |
| mediumSwitch | integer | 0 | Treatment of In-Medium Widths for mesons: <ul style="list-style-type: none"> • 0: Only vacuum widths are used. • 1: The collisional width is assumed to be constant (only density-dependent). • 2: The full tabulated in-medium width is used, as calculated via the collision term. |
| Gamma_coll_rho | real | 0.150 | Collisional width for the rho meson in GeV. Only used if mediumSwitch = 1. |
| Gamma_coll_omega | real | 0.150 | Collisional width for the omega meson in GeV. Only used if mediumSwitch = 1. |
| Gamma_coll_phi | real | 0.030 | Collisional width for the phi meson in GeV. Only used if mediumSwitch = 1. |
| verboseInit | logical | .false. | switch on/off informational messages during initialization |
| allowMix | logical | .false. | switch on/off linear interpolation between bins in density while returning the tabulated values for MassAssInfo. |

| XsectionRatios_input code/collisions/phaseSpace/XsectionRatios.f90 | | | |
|---|---------|---------|---|
| flagScreen | logical | .false. | If .true. – The in-medium screening is applied to the input cross section. If .false. – No cross section modification . |
| flagInMedium | logical | .false. | If .true. – In-medium ratios are used to decide whether an event is accepted or not. If .false. – The event is always accepted |
| flagTabulate | logical | .false. | If .true. – in-medium ratios are tabulated. If .false. – in-medium ratios are read-in. (This flag is important only if flagInMedium = .true.) |
| shift0 | real | 0. | Mass shift $m-m^*$ (GeV) for using in elementary particle collision mode. |

| YScalingAnalysis code/analysis/yScalingAnalysis.f90 | | | |
|--|---------|---------|--|
| analyze | logical | .false. | Determines wether the y-scaling analysis is performed |
| optionalOutput | logical | .false. | Determines wether in addition to the standard 'scaling_analysis.dat' other histograms will be generated. E.g. * 'single_nucleon.dat' - a table for comparing nucleon-knockout with fully inclusive <pre> jpre¿ x sections * 'scaling_info.dat' - general parameters of the analysis, to be used for quick jpre¿ analysis * 'scaling_delta.dat' - output to be used for analysis of scaling function in jpre¿ resonance excitation region </pre> |
| variable | integer | 1 | determines which kind of scaling variable will be used (cf. Donnelly, Sick 1999): <ul style="list-style-type: none"> • 1) RFG full variable Psi • 2) RFG approximation Psi • 3) PWIA full Upsilon (y/kf) • 9) evaluation will be done for all variables, output written to separate files |
| kFermi | real | 0.2251 | Nucleon Fermi momentum in nucleus. If none specified 0.2251 will be used, except if densitySwitch_static is set to 5, then fermiMomentum.input is used. The 0.225_1_ aims at preventing confusion whith delibaretely set differences between kFermi and fermiMomentum.input |
| E_shift | real | 0.020 | Energy correction to account for binding effects, otherwise neglected in RFG model |

| Yukawa code/potential/yuka.f90 | | | |
|---------------------------------------|---------|---------|----------------------------------|
| yukawaFlag | logical | .false. | Switches Yukawa potential on/off |

| | | | |
|-----|------|-------|--|
| smu | real | 2.175 | Yukawa mass in fm ^{**(-1)} . (range of potential) |
|-----|------|-------|--|
