

IMINTDYN81 (Version 8.3.0)

Ion Matter Interaction - Dynamic

A versatile and fast Monte Carlo Binary Collision Approximation simulation software

Command Parameters

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1 Program Structure

Main program IMINTDYN:

1. Default_init
2. Init-parallel_environment
3. Read namelist imint.inp
4. Setup simulation mode and free flight path option
5. Check_series_mode
6. Zero_init
7. Default_init
8. Read namelist imint.inp
9. Check symbols
10. Set series_mode
11. Read_all_tables
12. Set scattering_species
13. Set nra_species
14. Read_r33 cross section files
15. iba_init
16. main braodcast
17. set_matrix_ranges
18. initialize random, number generator
19. set max. atomic fraction
20. initialize vacancy mode
21. initialize noblegas treatment
22. angle_init
23. energy_init
24. chemical_init
25. outgas_init
26. diff_ko_init
27. set OR_loss electronic energy loss
28. target_init
29. setup_isotopes
30. main broadcast
31. create_layer_profile
32. sbv_init – surface binding energy model
33. create free_flight path tables
34. create energy_loss tables
35. create sublimation enthalpy table
36. allocate memory
37. setup particle info details
38. setup trajectory details
39. create angle vs impact parameter tables
- 40. call histories, projectile and recoil**
41. gsum_statr, gsum_fleucne, gsum_mat
42. out_particles, out_trajectories
43. outp
44. deinit_parallel_environment

Subroutines history, projectile and recoil:

1. Seed random number generator
2. init_dynamic_load_balancer
3. initialize surface binding energies and models
4. initialize parallel processing
5. loop for each processor core and number of histories
 - a. call projectile
 - b. call recoil
 - c. call oligomer_sputtering
6. end loop
7. update target structure and composition and update output data files after every history

2 Global parameters

Most of the commands were replaced by self-explaining commands. Therefore there is an incompatibility between different version of INIMTDYN. For example the term “nqx” was replaced by “numslices” and later by “target_slices”, because “target_slices” best describes the meaning of splitting the target given by its “target_thickness” (formerly “ttarget”) into a number of slices. Therefore, old script files of previous versions of IMINTDYN have to be modified and updated. This can be done by using an old script file and running IMINTYN in test mode. Evry incompatible command line will appear as error and the corresponding command can be corrected.

The following changes have been made in the different version of the program:

Imintdyn83 changes

The new interaction potential provides by Nordlund Lehtola, Hobler was implemented, The potential is used als default and is selected by specifying potential = "NLH" in the input script file

Imintdyn82.1 changes

numslices > target_slices

ioutput_part > ioutput_particles

subroutine setup_histories updated

../bin/linux.PRO/hostfile on new computer updated

Bug in tableread corrected

Separate programs histogram_erda and histogram_nra created and updated.

Erda treatment updated, in particular confinement of azimuth angles

makefile and makefilepost updated

only file_r33 for non-RBS and NRA

Imintdyn82 changes

numslicesm > slicesmax

numslices > numslices

ncpm > speciesmax

ncp > numspecies

e0 > energy0

e0_inc > energy0_inc

dns0 > atomic_density

name changed: param.F90 -> parameters.F90

imint.sh script file updated

imintdyn 81 changes

script batch file updated

output of atomic concentrations into logbook

postprocessing updated and minor bugs eliminated

layer-input modified, so that a too high number of layers is not possible

identifier for coincidence events in post processing updated

option compound_dns_mode removed

Update of oligomer_sputtering. Some bugs were corrected

Position of Subroutine OR_loss was changed to occur after the setup of isotopes, because the latter one changes ncp and thus the size of various arrays. After that a call to mainbroadcast occurs.

Subroutine setup_isotopes was moved before iba_init

Postprocessing programs were updated and small bugs corrected.

Main script file imint.sh was updated for the test mode

Imintyn80 changes

New subroutines set_vac_stopping, get elements contributing to vacancy stopping

New subroutine get_AtNr_tmax, get atomic numbers of elements for vacancy stopping

Vacancy_stopping_fraction introduced, default = 1.0

Vacancy_species introduced. Up to ncp elements used for electronic stopping for vacancies. The contribution is based on the atomic fractions.

Vacancy stopping introduced for stopping mode 6 and 7 (SRIM and ZBL)

Target_init moved into tableread.F90 subroutine before loading of stopping power data

MPI broadcast extended for string vectors

The weighting in subroutine profile was corrected

parts of imintdyn are shifted in to subroutines in init_all

timestamp routines are updated to show total time for a series simulation

postprocessing with faulty line detection implemented

oligomer_sputtering is now a subroutine and part. of sub.F90; call from recoil

trajec_out(i) und partic_out(i) are defined as character vectors

Andock model removed from recoil

MPI_filewrite(fh,buffer,nchars) und MPI_filewrite1(fh,buffer(:),nlines,nchar) implemented

definition of allocatable array is done more flexible

mpi_wait added in routine mpi_filewrite and mpi_filewrite1 to ensure thread safety

MPI_Bcast extended for string vectors

trajectory_all.dat removed

variable "trajek" replaced by "traject"

for MPI writing we need the linefeed character: linefeed = char(10)

Imoments set als .false. default

no more sequential mode

only intel and oneapi compilers are supported

no debug option in makefile supported

only machtype=x86_64 is supported

debugging is now possible using the commands

debugging_pid=pid1,pid2 and

debugging_history= h1,h2

sbv and sbvtt not anymore a matrix but a 1D array

chemical sputtering and diffusion package taken from SDTrimSP 6.05

Imintdyn72 changes

NRA postprocessing corrected

Several routines were put into lba_init.F90

Buffered output to out_particles implemented.

Bufsize=nnn, default 100 (lines of output)

Particles_per core=nn, default 10

Out_particles optimized for new oneAPI Compiler from INTEL for AMD Ryzen Threadripper of Intel I9 processors

Program can now run with all cores on these processors, time for data output is now negligible

Dimer_sputtering, Dimer compound, table-dimers.txt

series_mode: New: "angle_lin", "angle_posneg", "energy_lin", "energy_log", "angle&energy", "history_log", "angle_random", "none"

energy_mode: "input_file", "3D_Maxwell", "1D_Maxwell", "fixed", "none", Works only for series_mode="none"

angle_mode: "input_file", "cosine", "cosine_2pi", "random", "posneg", "fixed", "none"; Works only for series_mode="none"

Case_e0: only used as internal parameter

Case_alpha: only used as internal parameter

"two_comp" replaced by "binary_compound"

"i_two_comp=" replaced by "compound_dns_mode"

e_bulkb(1:num_species) is now obsolete

"Kijflag" replaced by "curvature_flag"

"proj_per_history" replaced by "proj_per_history"

Imintdyn71 changes

File_r33_nra as namelist input

rra_target_species(ncpm) array

nra_product_species(ncpm) array

several different nra reactions and different products can be specified

nra(5, ncpm) as array

heavy(ncpm) array

Update of random number generator

More efficient handling of nraflag and enforced scattering, weight factor includes concentration. Output only if weightfactor > 0.

Read_r33 separated from read_all_tables

Matrix output: small changes and in commands and some minor bugs fixed.

Imintdyn70 changes:

- New scattering mode "nra"
- Read nra IBANDL R33 files
- New postprocessing option for ebs_scattering: "esa"
- New postprocessing option for erda_scattering: "nra"
- New subroutine "interpolate_fractions" in program projectile and recoil
- fix_layer_density

option Intel Fortran random numbers, use IFPORT, result = RAND([iflag]), result= RANDOM(iflag), RANDOM_INIT,RANDOM_NUMBER, RANDOM_SEED, SRAND

Imintdyn65: changes:

- Testmode, get command argument
- Particle data output also for series
- Post processing also for series output
- Cleaning up the series mode simulations:
- Vacancy_mode = 1,2,3,...
- Vacancy_sequence
- Call pvfktn(l,cx) calculates vacancy formation probability
- Fluence steps out
- No restart option
- shth = sheath_potential
- Ltime_red not used anymore
- use_isotopes
- atomic_fraction
- atomic_fraction
- max_atomic_fraction
- max_atomic_fraction_flag
- target_thickness
- cutoff_from_sbe
- post processing program update
- free path display update
- layer definition file .def
- layer output file .dat
- lorentz_flag

- matric output extended! `echo 0 | sudo tee /proc/sys/kernel/yama/ptrace_scope`
- `add_bulk_layers`
- `main_broadcast` as new subroutine
- `iba_init` as new subroutine
- `use_isotope` option was corrected. Now `setup_isotopes` subroutine appears after `target_init` subroutine. After that all parameters are broadcast again. This is necessary because `setup_isotopes` reorders all arrays of size `ncpm` because the value `ncp` changes
- `allocate_memory` as a new subroutine
- improved output for `logbook.dat`
- `set_series_mode` as new subroutine
- `set_scattering_species` as new subroutine
- `set_matrix_ranges` as new subroutine
- `create_layer_profile` as new subroutine
- `dsf` changed to `surface_thickness`
- postprocessing output files now all have 6 head lines and 3 subheader lines
- Estimate of dynamic thickness has been upgraded (`or_loss` now before `target_init`)
- postprocessing output has been modified for 5 decimal digit output

2.1 How to compile and run the program

The IMINTDYN package is stored in folder `imintdyn821` with subfolders “bin”, “case”, “post”, “src”, “tables”, “template inp.files”, and “template layer.def”. The subfolder “bin” contains a folder “linux.PRO” and a folder “testnml”.

The subfolder “src” contains all F90 source files as well as the files `Makefile` and `Makefile_testnml.mak`. It also contains a folder “SRIM-sources” with source programs to create the SRIM stopping data base for IMINTDYN.

To compile the program

- (i) open a terminal screen and change to `imintdyn821/bin/linux.PRO`
- (ii) execute the `Makefile` using the command “`sh mk`”. `Makefile` automatically identifies the INTEL compiler and then compiles and generates the program `imintdyn.exe`.
- (iii) To make sure that all sub-programs are newly compiled, you should execute the command “`make clean`” in advance

To compile the postprocessing programs

- (i) open a terminal screen and change to imintdyn82/post
- (ii) execute the Makefile using the command "sh mkpost". Makefile will then newly create all postprocessing programs.

The folder "tables" contains all required tables for imintdyn, as well as a folder SRIM_tables and SRIM_raw_data. In folder SRIM_raw_data is the application "read_srim_stopping" which reads all stopping .txt files and created the SRIM2013-nn stopping data files. These should then be copied into the SRIM_tables folder.

The folder "case" contains the simulation data, each simulation has its own subfolder. In this subfolder there is the simulation script file "imint-filename1.inp" and optional script files "filename2.inp" and "layerdefinition.def". To run a simulation one needs to start the script file "imint.sh" using the command "bash imint.sh". After the simulation is finished, the results are stored in folders with name "filename1-filename2".

In the file "imint.sh" one can define several scriptfiles and support files. Before a simulation is started, one should set the script file to testmode. In testmode only the setup part of imintdyn is executed and one gets detailed error response in case the input script files *.inp contain some errors. Only if the testmode finishes without errors, one should deactivate the testmode and start the simulation.

The simulation uses a temporary directory in the folder case.

2.2 Parameters defined in module param.F90:

ncpm = 15 maximum number of elements (projectiles and target elements) [up to 999]
numslicesm = 2000 maximum number of depth intervals
thick_min = 1.0 minimum target thickness [A] (sufficient for a 1D layer of 2D material)
default cutoff_fraction = 0.333 default fraction to calculate the cutoff energy from the sublimation enthalpy

2.3 Parameters for random numbers

If ART_RAND is selected, then
Else

iwpi = 4 long integer format for random seed number
iwpi = 8 double long integer format for random seed number

2.4 Other parameters defined in module param.F90 (and used in dlb.F90):

pemax = 128	maximum number of Processors
nalpha = 180	max number of data in angular input file (a precision of 0.5 deg for angles between 0 and 90 deg seems sufficient)
nemax = 200	max number of data in energy input file
max_generation = 1000	maximum number of recoil generations
non_RBS = 1024	number of elements in a non_RBS cross section file
yzlost = 115.	Factor to find lost particles (y,z) >= yzlost*ttarget

2.5 Parameters defined in module default_init, used in module task_description.F90 and dlb.F90 (only MODE: PAR)

dntmax=100000	memory size of global task queue
ntqmax=100000	memory size of local task queue

Allocated memory: $(ntqmax+dntqmax) \times (13 \times (4 \text{ Bytes}) + 18 \times (8 \text{ Bytes})) / 1024. / 1024.$ (in Mbytes) = 37.38 MBytes

MEMORY for trajectory: assumption
 Integer = 4 byte
 Real = 8 byte

one trajectory -> $(ncpm+13) \times (4) + 18 \times (8) = 256$ Bytes for ncpm=15

2.6 Parameter ptime is a variable calculated in subroutine "inelast.F90"

It describes the flight time between two collisions. It is calculated from the pathlength divided by velocity and is given in units of femtosceonds.

3 Input file 'imint.inp' structure for an imintdyn.exe simulation

IMINTDYN uses one or two input files to define a simulation. The first and primary file has the name "imint-nameX.inp" with the precursor "imint". It contains all necessary information of a simulation, for example imint-ArSi1keV.inp to simulate sputter erosion of Si with 1 keV Ar ions. Upon execution of the simulation, this file is copied to a temporary directory under the name "imint.inp".

The second file has the name "subnameX.inp" (without a precursor "imint") and contains specific details of the simulation which can be changed easily without modifying the primary file. The content of "subnameX.inp" overwrites the previous data from file "imint-nameX.inp". For example, n001.inp to define a negative curvature of the sample surface. Upon execution of the simulation this file is copied to a temporary directory under name "imintb.inp"

Within the execution script file "imint.sh" one can define several primary and secondary input files, for each primary file a secondary file is required.

The form of imint-nameX.inp file is that for an input file (header + namelist):

The 1st line which does not start with a '!' character is the Headline (Text), ASCII format.

The following lines are defined by the namelist IMINT_INP (see subroutine WORK.F90) . The variables and values follow the initial line &IMINT_INP and close with the line " / ". The script file may contain FORTRAN Comments starting with " ! " as 1st character . The head line appears before &IMINT_INP it is a line which does not start with a "!" character.

Namelist is a list of variables or arrays, **separated by commas**

The Namelist in program imintdyn.exe is IMINT_INP . The structure is start line "&IMINT_INP" , end line "/" .

```
&IMINT_INP
<variables and values>
/
```

The sequence of the input variables and values in the input file is arbitrary (namelist).

3.1 Testmode

To test if the input script file contains nor errors, one can run IMINTDYN in testmode. In this mode I)NIMTDYN setup is processed, but no simulation is started. Furthermore, only one processor is used. The testmode can be activated with a command line extension “ -testmode”. For example a simulation using MPI routine and a number \$np of processors can be tested using the following line in the bash script file imint.sh.

```
mpirun -np "$NP" ../../bin/linux.PRO/imintdyn.exe -testmode
```

The bash script file has a variable TM defined at the beginning of the file as TM="-testmode" or TM="-off". The variable is added to the program execution command line as

```
mpirun -np "$NP" ../../bin/linux.PRO/imintdyn.exe „$TM”
```

4 Necessary input variables (in some cases no default values exist)

Command	Default value	Description
alpha0(1:num_species)	none	See: series_mode="angle", see: angle_mode=
alpha0_inc(1:num_species)	none	angle increment (deg) for series_mode="angle"
energy0(1:num_species)	none	See: series_mode="energy", see: energy_mode=
energy0_inc(1:num_species)	none	energy increment (eV) for series_mode="energy" and for projectiles (beam_fraction > 0.)
fluence	0.1	incident fluence in units 10^{16} atoms/cm ² or 1 atom/Å ² in case of a dynamic simulation default is 0.1 atoms/cm ² or $1 \cdot 10^{15}$ atoms/cm ²
potential	"NLH" ASCII code Keeps the old ipot variable internally	selects the interaction potential "NLH" Nordlund, Lehtola, Hobler interaction potential (default) This potential has an individual data set for each projectile target combination ! "KrC", well suited for sputter erosion and low energy collisions "Moliere", "ZBL", "Nakagawa" = Nakagawa-Yamamura "Si-Si", "power"
projectiles	None	If defined with value > 0, then histories and proj_per_history are calculated based on num_cores and dynamic or static simulation mode.
histories	24	number of histories (projectiles) each history consists of proj_per_history individual projectiles. The total number of projectiles is then histories*proj_per_history. In dynamic mode : target update occurs after each history proj_per_history(default) =24 is the default value, but may be changed by the program based on the number of processors used. This ensures that all processors have the same computational load.

Command	Default value	Description
		<p>proj_per_history should not be too high to avoid significant incremental changes of the stoichiometry</p> <p>proj_per_history should be \leq num_cores in dynamic mode. Then it is: $\text{proj_per_history} = \text{proj_per_history}(\text{default}) + \text{num_cores} - \text{mod}(\text{proj_per_history}(\text{default}), \text{num_cores})$</p>
target_thickness =	none	<p>total target thickness in Angstrom (\AA)</p> <p>The minimum default value is 3 \AA, set in "param.F90"</p>
target_slices =	100	<p>number of slices of the target (discretization). The maximum value maxslices = 2000 (default) set in param.F90 .If target_slices < 0, then the positive absolute value of target_slices is used.</p> <p>Target_slices = 0 is not allowed.</p> <p>If a file e.g. "layer.def" defines the number of target slices, then target_slices is set from this layer.def file.</p> <p>Target_slices should not be too large, or a layer thickness should not be smaller than an atomic spacing or 1 \AA. In this case the concentration of atoms in a layer may get zero in case of a dynamic simulation.</p>
add_bulk_layers=	.true.	<p>If the number target_slices of target layers decreases below the initial number specified in the input script file, then a layer with the composition and thickness of the last layer is added. In this way the target gets an infinitely thick bulk substrate and target_slices does not decrease below its initial value.</p> <p>This option prevents a target from being completely eroded.</p>
beam_fraction(1:num_species)	none	<p>projectile atomic fractions (in the incident beam) of num_species species. Num_species is determined automatically based on the symbols specified.</p> <p>For a projectile: beam_fraction > 0. ,</p> <p>Note: it is not necessary to normalize beam_fraction values to $\text{sum}(\text{beam_fraction}(1:\text{num_species})) = 1$! This is done by the program.</p>

Command	Default value	Description
		For target atoms: beam_fraction = 0.
atomic_fraction(1:num_species)	none	initial target atomic fractions of num_species species in case of a homogenous initial composition. Note: sum(qu(1:num_species)) = not necessary to be normalized. Example: Ar on Ta ₃ O ₇ : qu(1:num_species) = 0., 3., 7.
symbol(1:num_species)	none	numspecies chemical symbols of elements according to table1. Symbol(1:numspecies) is a character*5 string. (special symbols: H2, D, T, He3, He, C_a, C_g, C_d, ta-C, C_f, Sp2, Sp3, Sp3H) (other special symbols: Vac (vacancy) , Ar-l, Ar-s, Ar-g (liquid,solid,gas), same for other noble gases)

5 Optional variables:

These values have default values defined by subroutines default_init and zero_init (see default_init.txt). If values different from the default values are needed, then these values have to be given explicitly in the input file. Other choices are provided in the description column

5.1.1 Definition of input data file names and directories and output file options

Command	Default values	Description
dir_angleinp =	'./'	directory of inputfile 'angle.inp' as default name (or another specified name), default is the current directory (see also: dir_layerinp, dir_tableinp, dir_energyinp)
dir_energyinp=	'./'	directory of inputfile 'energy.inp' as default name (or another specified name) , default is the current directory (see also: dir_layerinp, dir_tableinp, dir_angleinp)
dir_layerinp=	'./'	directory of inputfile 'layer.inp' as default name (or another specified name); the default directory is './' . (see also: dir_tableinp, dir_angleinp, dir_energyinp). example: home/imintdyn/case/specific_case/
dir_tableinp=	'../tables'	directory of inputfile for tables; (see also: dir_layerinp, dir_angleinp, dir_energyinp)
file_layerinp=	'layer.def'	filename of layered target structure defintions: The output files have names "layeroutput-000.dat" and "layeroutinp-000.def" The number is filled in case of a series simulation with the series index. The file layeroutinp-000.def has an additional empty 1 st column for a projectile element an can be directly used as input file for e.q. a SIMS analysis simulation using Cs O or other ions as target species.
file_angleinp=	'angle.def'	filename of angular input data defintions. maximum 180 angular values (set in subroutine "work" the file has 2 header lines and the data pairs of angle[deg], probability (not normalized)
file_energyinp=	'energy.def'	filename of energy input data definitions maximum 200 values (set in subroutine "work", only energies 0.1 eV < E < 10 ⁹ eV are accepted, 2 header lines, pairs of energy[eV], probability (not normalized)

Command	Default values	Description
fluence_steps_out =	100	Determines the output for Number of fluence steps. The old parameter idout is used internally and is calculated as idout=histories/fluence_steps_out A large value of fluence_steps_out causes output files of very large size
ioutput_trajectories (1:6)=	10	number of traced trajectories for: stopped, backscattered and transmitted projectiles, stopped, backspattered, transmission sputtered recoils (see also: ltraj_p, ltraj_r) The value represents the number per processor core should not exceed 5-10 because trajectory calculation require very much allocated memory
ioutput_particles (1:6)=	10000	number of traced particles for: stopped, backscattered and transmitted projectiles, stopped, backspattered, transmission sputtered recoils (see also: lparticle_p, lparticle_r). The file size for 100,000 particles may reach 30 Mbytes.
layer_input =	.false.	initial target composition flag if layer_input=.true., then read the layer structure from layer input file see also dir_layerinput and file_layerinput true : initial depth dependent composition taken from layer input file (e.g. layer.def) false : (default) initial composition is homogeneous, target_slices layers with constant layer interval thickness layer input file starts with 2 text lines, followed by several lines: j, xxx, atomic_fractionx(1:num_species) j > 0 : number of layers ; xxx layer thickness in Å j = 0 : duplicate previous structure until target_slices defined in imint.inp

Command	Default values	Description
		j < 0 ; end reading input file. In the case the other input values in the line are ignored calculates also new values for target_slices and new ttarget !
lenergy_distr=	.false.	output of the energy distribution in the target (stop energy ,electronic loss and elastic nuclear loss) E_distr_all.dat E_distr_stop.dat E_distr_inel.dat E_distr_nucl.dat
dist_nx=60	60	x-size of the matrix of energy distribution in target
dist_ny=60	60	y-size of the matrix of energy distribution in target
dist_nz=60	60	z-size of the matrix of energy distribution in target
dist_delta=2.0	2.0	distance between the matrix points of energy distribution in target
elosstables=	.false.	[default: .false.], if true, then electronic energy loss data tables are created. The default data file name in elosstable(n).dat with n=1,...num_species The table consist of 301 energy values in the range 10 eV to 2 GeV
impactflag	.false.	If .true., then an output file of scattering angle as function of impact parameter for the incident energy energy0 and p _{impact} in the range 10 ⁻⁶ to 10 Å is created. This table allows to adjust the “enforce_scattering” option.
shtable=	.false.	If .true., then create a table of sbe[eV] versus concentration (for binary compounds)
lorentz_flag	.false.	If .true. then a relativistic correction is included for energy straggling and relativistic mass of projectiles
dimer_sputtering	.false.	.true. : activate dimer sand oligomer sputtering calculation
dimer_compound		Taken from table-dimer.txt Results are stored in dimer-sputtering.dat
dimer_efficiency	1.0	Efficiency of dimer formation 0 - 1
sbe_dimer_p	.true.	default momentum driven SBE barrier for dimers

5.1.2 Matrix output distribution commands:

Command	Default values	Description
lmatrices=	.false.	.true. : output of matrices for emitted particles, only for static simulations .false. : no matrix output
matrix_transmitted =	.false.	create matrices for transmitted particles
matrix_backscattered =	.true.	create matrices for back-emitted particles
matrix_cumulative =	.true.	cumulative or differential data collection, only for dynamic simulations
matrix_log_energy =	.false.	Output type of energy in matrix energy vs depth for emitted particles .false. : linear energy intervals .true. : logarithmic energy intervals
matrix_cos_angle =	.false.	Output of emission angle in a matrix for emitted particles .false. : angle in degree intervals .true. : cosine intervals
matrix_e_min_r=	0	minimum of lin. energy distribution of recoils in matrices [eV]
matrix_e_max_r=	none	max(energy0) maximum of lin. energy distribution of recoils in matrices [eV]
matrix_e_min_p=	0	minimum of lin. energy distribution of projectiles in matrices [eV]
matrix_e_max_p=	none	max(energy0) maximum of lin. energy distribution of projectiles in matrices [eV]
matrix_e_steps_lin_r=	10	Number of recoil energy steps between min and max value-linear scale
matrix_e_steps_log_r=	-1	Number of recoil energy steps between min and max value-log scale, 9 steps per decade
matrix_e_steps_lin_p=	10	Number of projectile energy steps between min and max value-linear scale
matrix_e_steps_log_p=	-1	Number of projectile energy steps between min and max value-log scale, 9 steps per decade

Command	Default values	Description
File names for matrix output:		meagb_p_file ='matrix_energy_angle_back_proj.dat' meagt_p_file ='matrix_energy_angle_trans_proj.dat' meagb_r_file ='matrix_energy_angle_back_sputt.dat' meagt_r_file ='matrix_energy_angle_trans_sputt.dat' mepb_p_file ='matrix_energy_path_back_proj.dat' mept_p_file ='matrix_energy_path_trans_proj.dat' mpe_ex_p_file='matrix_start energy_vs max_penetration_back.dat' polar and azimuthal angular_matrix_file_br ='meagb_r_000000.dat' polar and azimuthal angular_matrix_file_bp ='meagb_p_000000.dat' polar and azimuthal angular_matrix_file_tr ='meagt_r_000000.dat' polar and azimuthal angular_matrix_file_tp ='meagt_p_000000.dat'

5.1.3 Output of detailed projectile and recoil information:

Command	Default values	Description
Imoments=	.false.	output of moments for energy distributions (linear and logarithmic) of projectiles and recoils and for range distributions (linear) of projectiles. .true. : moments are written in to a output file .false. : moments are not written
lparticle_p=	.false.	.true. : output of projectile information .false. : no output of projectile information (see also: ioutput_particles)
lparticle_r=	.false.	.true. : output of recoil information .false. : no output of recoil information (see also: ioutput_particles)

Command	Default values	Description
out_angle_pt =	0.,90.	Defines minimum and maximum emission angle of projectiles for output data Forward transmitted: $0 \leq \text{angle} \leq 90$ deg
out_angle_pb =	90.,180.	Back scattered : $90 \leq \text{angle} \leq 180$ deg
out_angle_rt =	0.,90.	Defines minimum and maximum emission angle of recoils for output data Forward transmitted: $0 \leq \text{angle} \leq 90$ deg
out_angle_rb =	90.,180.	Back sputtered : $90 \leq \text{angle} \leq 180$ deg
out_energy_p =	0.,	Defines minimum energy of emitted projectiles for output data
out_energy_r =	0.,	Defines minimum energy of emitted recoils for output data
ltableread =	.true.	Default strongly recommended is .true. .true. : read from table_elements, table_compounds, table_isotopes, table_Hstopping table_Hestopping1 .false. : no table read, a_num_z, a_mass, dns0, e_surfb, e_displ have to be given tables: table_elements .txt: chemical symbol (symbol), nuclear charge (a_num_z), atomic mass (a_mass), mass density, atomic density (dns0), surface binding energy (e_surfb), displacement energy (e_displ), cutoff energy (e_cutoff) table_compounds.txt : symbol of binary cvcompound target and physical values see also: nm, tabe.compound, binary_compound, dimer_sputtering, rho_mean table_isotopes.txt : chemical symbol, nuclear charge, isotope mass, atomic weight (in amu), natural abundance table_Hstopping.txt : inelastic stopping coefficients for hydrogen: symbol, nuclear charge, inelastic stopping coefficients a1 to a12 (ch_h), ck

Command	Default values	Description
		table_Hestopping1.txt , table Hestopping2.txt: inelastic stopping coefficients for helium: symbol, nuclear charge, inelastic stopping coefficients a1 to a9 (ch_he)
ltraj_p=	.false.	.true. : output of projectile trajectories .false. : no output of projectile trajectories
ltraj_r=	.false.	.true. : output of recoil trajectories .false. : no output of recoil trajectories

5.1.4 Definition of ion species and target elements

Command	Default values	Description
symbol(1:num_species)	none	num_species chemical symbols of elements according to “table1-elements.txt” (special symbols: H2, D, T, He3, He, C_a, C_g, C_d, ta-C, C_f, Sp2, Sp3, Sp3H, a-C1D). Other special symbols: Vac, Vac1D (vacancy) , Ar-l, Ar-s, Ar-g (liquid,solid,gas), same for other noble gases).
beam_fraction(1:num_species)	0.	projectile atomic fractions (in the incident beam) of num_species species. For a projectile: beam_fraction > 0. , Note: sum(beam_fraction (1:num_species)) = 1 is normalized by the program. For target atoms: beam_fraction = 0.
atomic_fraction(1:num_species)	none	initial target atomic fractions of num_species species in case of a homogenous initial composition (iq0 = 0) Note: sum(atomic_fraction (1:num_species)) = 1 is normalized by the program
max_atomic_fraction(1:num_species)	none	maximum atomic fractions in the target for num_species species, if dynamic simulation. Example: for SiO ₂ one should set max_atomic_fraction = 1.0 ; 0.667 so that excess oxygen diffuses out
max_atomic_fraction_flag	.false.	.true. , then calculate max_atomic_fraction = max_atomic_fraction (0 deg) * cos(alpha0) only if case_apha=5 and only for species with beam_fraction > 0. This is an option, that adjusts the retained noble gas concentration for varying incidence angles using an empirical cosine law. .false., default
charge(1:num_species)=	0.	elementary charge of ion species if case_e0 = 2 or 3 and sheath_potential > 0 (plasma). Otherwise it is not used. >= 1. for beam_fraction > 0 (projectiles) charge state should at least be 1+ . = 0. for beam_fraction = 0 (target atoms)

Command	Default values	Description
alpha0(1:num_species)	none	angle of incidence (in deg) of num_species species used depending on angle_mode and series_mode alpha0 is measured in neg y-direction with respect to the surface normal (-x) if alpha0 < 0, then azimuth angle phi is set to 180 deg
alpha0_inc(1:num_species)	none	angle increment (deg) for linear angle series
energy0(1:num_species)	none	Incident ion energy
energy0_inc(1:num_species)	none	energy increment (eV) for projectiles (beam_fraction > 0.) in case of linear energy series
series_mode	"none"	"angle_lin", "angle_posneg", "energy_lin", "energy_log", "angle&energy", "history_log", "angle_random", "none"
angle_mode	"none"	"input_file", "cosine", "cosine_2pi", "random", "posneg", "fixed", "none" Works only for series_mode="none"
energy_mode	"none"	"input_file", "3D_Maxwell", "1D_Maxwell", "fixed", "none" Works only for series_mode="none"
series_steps=	1	number of calculations if a series of calculations is carried out (for a given selected series mode) Histories = log scale with steps 5,16,50,160,500,1600,5000,.... energy log scale: 3 values per decade: 1,2,5,10,20,50, etc
case_e0 only used as internal parameter	0	flag for the choice of the incident energy: = 0 : fixed incident energies (eV) of projectiles (beam_fraction > 0.) = 1 : input of a given energy distribution from file energy.inp = 2 : temperature (eV) of plasma of a 3D Maxwellian velocity distribution of projectiles, particle energy e_part comes from subroutine mxvelo = 3 : temperature (eV) of plasma of a 1D Maxwellian energy distribution of projectiles, particle energy e_part comes from subroutine energ = 4 : does not yet exist = 5 : series of calculations with different projectile energies

Command	Default values	Description
		<p>linear series with fixed incremental energy $energy = energy0 + (i-1) * eenergy0_inc ; i = 1, series_steps)$ if also case_alpha = 5, then the program ignores case_e0 !! output: output???.dat with ??? = i default setting: lmatrices = .false. ; ltraj_p = .false. ; ltraj_r = .false. ; lparticle_r = .false. lparticle_p = .false. ; case_alpha = 0 note: all *.dat files from last calculation</p> <p>= 6 : series of calculations with different angles of incidence and different energies (also sets case_alpha0 =6) INPUT OF ANGULAR AND ENERGY DISTRIBUTION FROM FILE see subroutine ene_ang_init in init_all</p> <p>= 7 : series of calculations with different projectile energies logarithmic series given by 3 values per decade: 1,2,5,10,20,50, etc $kk = 1 + \text{mod}(iseries_steps - 1, 3)$ $energy0(:) = eenergy0(1) * 10^{((iseries_steps - 1) / 3) * \text{int}(10^{(kk * 0.35)})}$ example: energy0=20, series_steps=6 : 20,40,100,200,400,1000 eV</p> <p>output: output???.dat with ??? = i default setting: lmatrices = .false. ; ltraj_p = .false. ; ltraj_r = .false. ; lparticle_r = .false. , lparticle_p = .false. ; case_alpha = 0 (note: all *.dat files from last calculation)</p>
case_alpha = only used as internal parameter	0	flag for the choice of the angle of incidence : = 0 : angle of incidence (degree) counted from the surface normal alpha0 = 0... 90 (starting above surface)

Command	Default values	Description
		<p>alpha0 = 90...180 (starting in solid) (azimuthal angle phi = 0)</p> <p>= 1 : input of a given incident angular distribution from file angle.inp</p> <p>= 2 : cosine distribution of angles of incidence (only from above surface)</p> <p>= 3 : cosine distribution of angles of incidence alpha=0...Pi/2,max: at 0 phi= 0...2Pi</p> <p>= 4 : random distribution of angles of incidence (only above surface) (alpha and phi random)</p> <p>= 5 : series of calculations with different angles of incidence (alpha = alpha0 + (i-1) * alpha_inc; i = 1, series_steps) output : output.*dat default set :lmatrices = .false. ltraj_p = .false. ltraj_r = .false. lparticle_r = .false. lparticle_p = .false. case_e0 = 0 (note: all *.dat output files from last calculation) if also case_e0 = 5, then the program ignores case_e0 !!</p> <p>= 6 : series of calculations with different angles of incidence and different energies (also sets case_e0 = 6) INPUT OF ANGULAR AND ENERGY DISTRIBUTION FROM FILE see subroutine ene_ang_init in init_all</p> <p>= 10 : fixed polar angle of incidence (same as case_alpha = 0) but alternating opposite azimuthal angle phi=0 deg or phi=180 deg</p> <p>= 15 : series polar angles of incidence (same as case_alpha = 5) but alternating opposite azimuthal angle phi=0 deg or phi=180 deg</p>

Command	Default values	Description
phi0 = (1:num_species)=	0	azimuth angle of ion incidence in deg; 0 deg = along y-axis; 90 deg = along z-axis
atomic_mass(1:num_species)=	> = 0 < 0: use table_elements	mass (in amu) of num_species elements; default from "table_elements.txt" atomic_mass = 0 is a vacancy, then also atomic_number = 0.
atomic_number(1:num_species)	See table_elements	atomic number of num_species elements; default from "table_elements.txt"
use_isotopes(1:num_species)=	0	flag for isotope mass = 0 : natural isotope mixture (mass from "table_elements.txt") = 1 : isotope masses and natural abundances from "table_isotopes" (valid for projectiles as well as for target species) the masses are set by a_mass(1:num_species) or taken from "table_elements.txt" use_isotopes(:) = 1 increases the number of species to account for the number of isotopes. Each isotope is included as a new target species. Make sure that ncpm (default 15) is not exceeded !

Command	Default values	Description
compound_flag =	.false.	.false.: not a binary molecular target If .true. then simulation is for a binary compound defined by "binary_compound" and rho_mean, the number nm of atoms in a molecule is taken from " table.compound " There may be more than one projectile, but only two target elements which appear last, i.e. num_species-1 and num_species !!!!! Thjer compound sublimation enthalpy is defined in the table, but it is not used by default(the table contains a correction factor set to zero. If a compound formation enthalpy shall be use, the correction factor should be set > 0. Then, for a binary compound a new model is used to calculate the sublimation enthalpy automatically for bvariable stoichiometry of the compound!
binary_compound =	none	symbol of binary compound target according to " table_compounds " (e.g. binary_compound ="Ta2O5"). binary_compound is a character*10 string Note: only selected compounds in " table_compounds " see also: nm, " table_compounds ", binary_compound, rho_mean There may be more than one projectile, but only two target elements which appear last, i.e. num_species - 1 and numspecies !!!!!
rho_mean =	none	mass density of a binary compound target; default from " table_compounds " [g/cm**3) see also: nm, " table_compounds ", binary_compound, rho_mean There may be more than one projectile, but only two target elements which appear last, i.e. num_species-1 and num_species !!!!!
delta_hf_corr =	0.	Correction factor to scale the compound formation enthalpy $0 < \text{delta_Hf_corr} < 1$ This overrides the factor from the compound table
target_thickness =	none	total target thickness in Angstrom (Å) The minimum default value is 3 Å, set in "param.F90"

Command	Default values	Description
target_temperature =	300.	target temperature, only of interest at high temperatures, it reduces the surface binding energy according to a Maxwellian energy distribution
x0(1:num_species)=	0.	starting position (impact) of projectile (see also curvature Kij) in Å ≤ 0 : outside the surface at x=x _c > 0 : inside the solid In a surface curvature is defined using the curvature parameters Kij, then the value of x0 should be set to typically + 100 Å.
deltahd(1:num_species)	See table_elements	heat of dissociation (eV) of elements dns0(1:num_species) of a molecular target
deltahf	See table_elements	heat of formation (eV) of a molecular target
atomic density (1:num_species) =	> 0 < 0 : use table_elements	atomic density in units (atoms/Å ³ or 10 ²⁴ atoms/cm ³) of num_species elements; Example: 0.05 for Si if atomic_density ≤ 0 or not specified, then the table_elements values is taken
fix_layer_density	.false.	If .true., then keep the initial density of each target layer fixed. If layers are dynamically added or deleted, then the new layer density of the affected layers are interpolated. This option avoids drastic density changes if a very low density gas atom is implanted into a dense solid (example: O into SiO ₂)

5.1.5 Definition of specific energy values

Command	Default values	Description
e_cutoff(1:num_species)=	1.	cutoff energy (eV) of target species; defaults from table1 (1 eV for noble gases; 1 eV for H, D, T; Normally a value of 1 eV is sufficiently low. If values of zero are specified, then a value of 1.0 eV is chosen as default.
cutoff_from_sbe	.true.	.true., then take cutoff_energy = cutoff_fraction * sbe, if simulation is in SBE_mode. If .false., the use values from table or imint.inp
cutoff_fraction	0.333	Fraction of the sublimation enthalpy to calculate the cutoff energy in sublimation mode 0 < cutoff_fraction <= 1 ; in case of another value the default value 0.333 is chosen in sublimation mode, the cutoff is taken as fraction of the sublimation enthalpy. For some gases, the sublimation enthalpy is zero. In this case, the cutoff is set to the corresponding e_cutoff value to ensure that a particle comes to rest.
e_displ(1:num_species)	0.	displacement energy (eV); default from table1. The displacement energy can be set to zero, but the simulation speeds up if a larger value is chosen. Note, that a displacement energy is only defined in a crystalline matrix and is used to estimate the amount of damage created (similar to SRIM software)
e_thres_r(1:num_species)	0.	threshold energy (eV); The threshold energy is used to define a minimum energy transfer to the recoil for bookkeeping of projectile and recoil data for CERDA

Command	Default values	Description
e_surfb(1:num_species)	See table1	<p>surface binding energy (eV) (sublimation enthalpy) for standard conditions, RT and standard pressure; default from table1</p> <p>Imintdyn.exe uses the tabulated sublimation enthalpies from table1 (new: table1-elements.txt). Or the values specified in the .inp file. This was the previous option isbv=1 in SDTrimSP and TRIDYN.</p> <p>In case of a binary compound with compound_flag=.true., the surface binding energy is modified based on the compound formation enthalpy tabulated in table.compound (new: table-compound.txt). In dynamic simulation and for sputtered recoils, the sublimation enthalpy is calculated based on the local concentration values in up to 10 layers to the front or back surface. For projectiles a sublimation enthalpy is set to zero, because they are never bound to target atoms.</p> <p>If a value > 0 is specified, then the correction enthalpy for 6000 K is scaled based on the tabulated value in table1. This scaling ensures, that the correction enthalpy remains smaller than the sublimation enthalpy at standard RT.</p>
temp_cascade=	298.15.	<p>Assumed temperature of the collision cascade in K. This value is used to reduce the surface binding energy (the sublimation enthalpy) and the binary compound formation enthalpy according to a linear approximation of the Shomate equation between 298K and 6000 K.</p> <p>If temp_cascade <= 0., then the temperature is approximated dependent of the projectile energy by $T = RT + (6000-RT) \cdot \exp(-\text{energy0(keV)}/2.)$</p>

Command	Default values	Description
sbe_mode =	.true.	If .true., then the surface binding energies are used in the usual way: A particle is only sputtered, if its kinetic energy normal to the surface is larger than the sublimation energy. If .false., then the sublimation energy is subtracted from the kinetic energy of recoils immediately after the recoil is created. To be sputtered, an energy > 0 is sufficient. A particle whose energy was reduced initially gets the sublimation enthalpy added before it comes to rest.
refraction =	.true.	Switches the refraction effect on and off
temp_auto =	.false.	.true.: Automatic calculation of the cascade temperature after 100 histories, or update every 100 histories for a dynamic simulation
tilt_flag =	.false.	If .false., then Sigmund's Gaussian ellipsoid is calculated in the x,y,z coordinate system. If .true., then Sigmund's Gaussian ellipsoid is calculated in the tilted coordinate system with x' along the beam direction.

Command	Default values	Description
stopping_mode(1:num_species) =	3	<p>inelastic loss model - how to calculate the electronic energy loss</p> <p>= 1 : Lindhard-Scharff; necessary condition: $E < 25 * Z^{4/3} * M$ (in keV), where E, Z, M are the energy, the atomic number and the atomic mass of the moving particle, respectively</p> <p>= 2 : Oen-Robinson; necessary condition: $E < 25 * Z^{4/3} * M$ (in keV)</p> <p>= 3 : Equipartition of 1 and 2 [DEFAULT]</p> <p>= 4 : high energy hydrogen (H,D,T) ($25 \text{ keV} < E < 475 \text{ MeV}$), values from table3</p> <p>= 5 : high energy helium (He3,He4) ($E > 100 \text{ keV}$), values from table4</p> <p>= 6 : Ziegler-Biersack stopping based on the tables scoef95A and scoef95b. This stopping model option is very similar to the data used in SRIM-2013</p> <p>= 7 : SRIM-2013 electronic stopping power data. This option may require to extract stopping data from the SRIM Software. Stopping powers are interpolated from a logarithmic table with values in the range 0 eV to 2 GeV. Raw stopping data files are stored in subdirectory tables in a folder "SRIM_raw_data". The program "read_srim_stopping.exe" reads the raw data files and creates the SRIM2013-nn.dat files, where nn is the atomic number of the projectiles. Each file contains up to 92 stopping tables for each target element. The stopping unit is eV per 10^{15} at/cm^2 and is independent of the actual target density.</p> <p>Imintdyn selects the stopping data for the num_species specified target atoms form the SRIM2013-nn.dat files. The SRIM2013-nn.dat files contain stopping powers for target atoms with Z=1 to 92. If a vacancy with Z=0 is specified as a target atom using symbol "Vac" the program set the electronic loss for such a "target atom" to zero.</p>

Command	Default values	Description
sheath_potential =	0.0125	= 0.0125 eV : weak sheath potential sheath_potential= $k_B T_e / 2e$ for $T=300K$ > 0 : sheath potential (eV), usually = $3 * energy0 $, only if case_e0=2,3 (Maxwellian distribution, plasma)
surf_inel_loss =	0	= 0 : no inelastic energy loss outside the target surface ($x \leq 0$, $x > ttarget$) = 1 : inelastic energy loss outside the target surface ($-suu > x > 0$. And $ttargets < x < ttarget + suu$) If (kijflag) is set, then the lower value $x=0$ is shifted to $x0(jin)$

5.1.6 Definition of simulation options

Command	Default values	Description
dynamic_simulation= static_simulation= cascade_simulation=	.false. .true. .true.	mode of simulation, parameter idrel= full dynamic calculation (TRIDYN), internally: dynamic_flag=.true., cascade flag =.true. cascade_flag=.false : only projectiles (no recoils) are followed. internally: static_flag=.true., Typical parameter for a dynamic simulation: dynamic_flag=.true., cascade flag =.true. Typical parameter for a static simulation is static flag =.true., cascade flag =.true.

Command	Default values	Description																																																
proj_per_history=	12,24	<p>number of projectiles between two target updates (histories) for dynamic simulation. In a static simulation, the total number of simulated ions is histories * proj_per_history.</p> <p>each history consists of proj_per_history individual projectiles. The total number of projectiles is then histories* proj_per_history . In dynamic mode: target update occurs after each history</p> <p>proj_per_history (default) =12 is the default value, but may be changed by the program based on the number of processors used. This ensures that all processors have the same computational load.</p> <p>proj_per_history should not be too high (<=12) to avoid significant incremental changes of the stoichiometry</p> <p>proj_per_history should be <= num_cores. Then it is: $\text{proj_per_history} = \text{proj_per_history}(\text{default}) + \text{num_cores} - \text{mod}(\text{proj_per_history}(\text{default}), \text{num_cores})$</p> <table border="0"> <thead> <tr> <th># of processors</th> <th>num_cores</th> <th>proj_per_history</th> <th>proj_per_history(default)=12</th> </tr> </thead> <tbody> <tr> <td>1,2,3,4</td> <td></td> <td>12</td> <td></td> </tr> <tr> <td>5</td> <td></td> <td>10</td> <td></td> </tr> <tr> <td>6</td> <td></td> <td>12</td> <td></td> </tr> <tr> <td>7</td> <td></td> <td>14</td> <td></td> </tr> <tr> <td>8</td> <td></td> <td>16</td> <td></td> </tr> <tr> <td>9</td> <td></td> <td>18 == > 9</td> <td></td> </tr> <tr> <td>10</td> <td></td> <td>20 == > 10</td> <td></td> </tr> <tr> <td>11</td> <td></td> <td>22 == > 11</td> <td></td> </tr> <tr> <td>12</td> <td></td> <td>12</td> <td></td> </tr> <tr> <td>13</td> <td></td> <td>13 == > warning</td> <td></td> </tr> <tr> <td>.....</td> <td></td> <td></td> <td></td> </tr> </tbody> </table>	# of processors	num_cores	proj_per_history	proj_per_history(default)=12	1,2,3,4		12		5		10		6		12		7		14		8		16		9		18 == > 9		10		20 == > 10		11		22 == > 11		12		12		13		13 == > warning				
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Command	Default values	Description
particles_per_core	10	Number of particles distributed to each core. This value is generally used for static simulations. In dynamic simulations the value depends on the number of cores_num_cores and the desired proj_per_history.
bufsize	100	100 line of buffered output to out_particles. 100 is sufficient for storing projectile and sputter data. For stopping of recoils the number may be even larger.
surface_thickness =	3.	Defines and average thickness of a surface layer (Å). This layer is used to calculate the surface composition as function of fluence
flux=	1.0	flux of incident atoms (atoms/Å ² /s = 10 ²⁰ atoms/m ² /s)
lchem_ch=	. false.	calculation with chemical erosion for H on C(SP ² ,SP ³ ,SP ³ H), D on C(SP ² ,SP ³ ,SP ³ H) using a mix of methods by Hopf, Roth and Mech
atomic_fraction_interpolate =	.false.	If .true. then linear interpolation of atomic fractions between the depth intervals. Only required if larger depth intervals and significant concentration gradients
integration_mode =	1	integration method (default=1) = 0 : MAGIC, only valid for KrC, ZBL, Moliere = 1 : Gauss-Mehler quadrature, ipivot >= 8 recommended = 2 : Gauss-Legendre quadrature, ipivot <= 16 = 3 : Gauss-Legendre quadrature 8 (double precision), ipivot <= 16
ipivot=	8	number of pivots in the Gauss-Mehler and Gauss-Legendre integration, the minimum number is 4 (larger numbers increase the computing time)
case_layer_thick=	0	mixing scheme of target = 0 : mixing layer with neighbor layer, if thickness is 150% or 50% = 1 : mixing to constant layer thickness
ca_scre(1:num_species,1:num_species)=	1.0	correction factor for the screening length in the interaction potential (not applicable for KrC and ZBL potentials)
ck_elec(1:num_species,1:num_species)=	1.0	correction factor for the inelastic energy loss; correction factors for hydrogen (below 25 keV) are given in table_Hstopping.txt
diff_koeff1(1:num_species)=	0.	transport coefficient if loutgas=.true. [Å ³ /ion]
diff_koeff2(1:num_species)=	0.	diffusions coefficient if loutgas=.true. [Å ⁴ /ion]

Command	Default values	Description
imcp=	0	flag indicating whether (flib)-moments of depth distributions are calculated = 0 : no moment calculation = 1 : calculate moments of depth distributions for all projectiles with (qubeam>0.)
subthr_rec_bound	.true.	flag for subthreshold recoil atoms .false. : subthreshold recoil atoms free >= 0 : subthreshold recoils bound
weak_coll_p=	2	number of ring cylinders for weak simultaneous collisions for projectiles ; for high energies (MeV H or He) weak_coll_p can be reduced to 1 or 0 to reduce computing time. The inner ring with $p=r1*pmax$ ($r1$ random number) calculates the primary collision. The first ring has $p=pmax*\sqrt{r1+1}$. The 2 nd ring has $p=pmax*\sqrt{r1+2}$, etc. For each ring the number $r1$ and also the collision partner is individually determined. For 2 rings, p may reach a value of $pmax*\sqrt{3}$, so that the tails of the screened potential becomes relevant.
weak_coll_r=	2	number of ring cylinders for weak simultaneous collisions for recoils ; for high energies (MeV H or He) weak_coll_p can be reduced to 1 or 0 to reduce computing time. The inner ring with $p=r1*pmax$ ($r1$ random number) calculates the primary collision. The first ring has $p=pmax*\sqrt{r1+1}$. The 2 nd ring has $p=pmax*\sqrt{r1+2}$, etc. For each ring the number $r1$ and also the collision partner is individually determined. For 2 rings, p may reach a value of $pmax*\sqrt{3}$, so that the tails of the screened potential becomes relevant.
Kij =	0.,0.,0.	defines a surface curvature by $h(y,z) = h_0 - 1/2 K_{11} y^2 - K_{12} yz - 1/2 K_{22} z^2$ Kij is given in units 1/Angstrom. A positive curvature describes a valley-like surface, note that the surface normal is along -x. If Kij not equal 0, then one has to set proper x_0 values, i.e. the curved surface at any point should be positioned below $x_0 = 0$ (values of $x_0 > 0$). Kij should be small enough, at most $ K < 0.002$

Command	Default values	Description
curvature_flag	.false.	If any of the Kij values is non-zero, the curvature is set .true. The flag can be set to .true. in the input file to enforce a shift of the surface to $x_0 > 0$, even if no curvature is defined. This is only used for test purpose. .Not.curvature_flag also influences the inelastic loss calculation (parameter surf_inel_loss) by shifting the target limits from $x=0$ to $x=x_0$.
debugging=	0	a parameter to help debugging program errors with different levels of debugging output 0: debugging is off 1: output from subroutines imintdyn.F90 and histories.F90 (logical flag "debugging1") 2: only some output from subroutine histories.F90 (logical flag "debugging2") 3: only output from subroutines imintdyn.F90 and projectile.F90 (logical flag "debugging3") 4: only output from subroutines imintdyn.F90 and recoil.F90 (logical flag "debugging4") 5: only debugging of subroutines set_sublenergy in sub.F90 6: only debugging of subroutines set_temp_cascade in sub.F90 7: only debugging of subroutine free_path in sub.F90 8: only debugging for gsum_fluence and gsum_fluence2 Implemented by H. Hofsaess 06-5-2021
debugging_pid=	0,num_cores	Only then debugging output is created (imintdyn80)
debugging_hist=	1,histories	Only then debugging output is created (imintdyn80)
craterflag=	.true.	Activates calculation of crater function moments and writing of crater function output files. See files: craterfile, cratermoments, balancefile, impl_crater_moments, zero_moments

Command	Default values	Description
monolayer=	.true.	true: (default) uses the shortest mean free path length $l_m=1/(dn_s0)^{1/3}$.false: uses $-l_m*\ln(1 - \text{random number})$. Can be used for a larger target thickness This option can be circumvented by free_path_length !!

5.1.7 Definitions for ion beam analysis simulations

Command	Default values	Description
scattering_mode =	'none'	'none', or 'cerda', 'ercs', 'ebs', 'erda', 'nra' cerda: coincidence ERDA: coincident scattering of identical projectile and target species ercs: elastic-recoil-coincidence-spectroscopy: coincident scattering/recoil of light projectiles and heavier targets species ebs: elastic backscattering: Rutherford or non-Rutherford backscattering erda: detection of light recoils from heavy projectiles emitted under small recoil angles in backscattering condition. ERDA is usually done in static_simulation but requires to set cascade_simulation=.true. nra: detection of nra reaction products only if one of the scattering modes is specified, then enforce_scattering remains active. Otherwise enforce_scattering is set to .false.

Command	Default values	Description
free_flight_path	.false.	If true, then calculate a free path length based on the scattering cross section and the fractional electronic energy loss. Basis is a minimum deflection angle (default 1 deg) between collisions and a minimum relative electronic energy loss (default 1%) between collisions
ffp_accuracy	0.01	Set the accuracy of the free path length approximation. A value of 0.01 sets a max deflection angle of 1 deg and a max. rel energy loss of 1%. The value should not exceed 0.05(or 5deg and 5%)
huge_angle =	15.	(default value) threshold for counting events with large scattering angle. The number of events with scattering angle above "huge_angle" are listed in particle*.dat files Note: for ERDA simulations, the scattering angles are usually small, then the value for huge angle should be to a reasonable small value.
small_angle =	2.	(default value) threshold for counting events with small scattering angle. The number of events with scattering angle larger than "small_angle" and below "huge_angle" are listed in particle*.dat files as medium angles Note: for ERDA simulations, the scattering angles are usually small, then the value for huge angle should be to a reasonable small value.
enforce_scattering =	.false.	If .true., then enforced scattering is activated. The program calculates the average number of collisions for transmitted and backscattered projectiles. The value num_scattering defines the number of large angle scattering events each projectile shall undergo. Bases on the average number of collisions and num_scattering a random number is selected, defining the probability to reduce the max. impact parameter. The reduction factor is set by pmax_coeff, which may be as low as 5E-6. A scattering_mode must be selected

Command	Default values	Description
single_scattering =	.true.	true: a random number r is chosen to find a target position $x_s = r * t_{\text{target}} / \text{ffp}$ where enforced scattering takes place. Only one scattering event is considered false: plural scattering mode is active. The probability for enforced scattering $p = \text{num_scattering} * \text{ffp} / t_{\text{target}}$. A random number r is chosen and if $r < p$, then enforced scattering is activated. There may be more than one scattering event, so that plural scattering takes place. With proceeding path step n , the probability for a projectile to survive is $(1-p)^n$.
enforce_rec_scatt =	.false.	If true, then enforced scattering also applies to recoils. This option is only used in C-ERDA simulations like HH- scattering or Li-Li scattering and only when single_scattering=.false.
cerda_angle cerda_angle2 =	0. 90.	the minimum and maximum scattering angle can be defined in the range $0 < \theta < 89$ deg. Used for identical projectile and scatter atom mass !! C-ERDA like H-H, He-He or Li-Li scattering. From that the impact parameters for enforced scattering are calculated
ercs_angle ercs_angle2 =	0. 90.	the minimum and maximum scattering angles can be defined in the range $0 < \theta < 89$ deg. Used for light projectiles and heavier scatter atom mass !! ERCS like He-B, He-C, He-N, He-O scattering. From that the impact parameters for enforced scattering the recoils are calculated Example: He projectile typically 60-80deg, recoils angles are typically 40-50deg
ebs_angle ebs_angle2 =	0. 180.	the minimum and maximum scattering angle can be defined in the range $0 < \theta < 179$ deg. Used for light projectiles and heavy scatter masses mainly for backscattering. From that the impact parameters for enforced scattering the projectiles are calculated

Command	Default values	Description
erda_angle erda_angle2 =	0. 90.	the minimum and maximum recoil angle can be defined in the range $0 < \theta < 90$ deg. Used for heavy projectiles and light scatter masses mainly for ERDA simulations. From that the impact parameters for enforced scattering of the recoils and projectiles are calculated. For $M_{\text{recoil}} < M_{\text{projectile}}$, there exists a maximum scattering angle. Small impact parameter events occur for recoil angles between the corresponding recoil angle and recoil angle of 0 deg. erda_angle2 is replaced by the recoil angle corresponding to the max. scattering angle
erda_azimuth azimuth_range	0° 180°	center and range of ERDA emission azimuth angles are given for the desired recoil emission. internally the azimuth angles are related to the projectiles. Erda_azimuth can be within -90 to 90 deg, azimuth range is 0 – 180 deg Therefore, the center 0 deg is converted internally to 180 deg (the sign of $\cos(\text{erda_azimuth})$ and $\sin(\text{erda_azimuth})$ are changed. The weight factor is increased by $\pi/\text{azimuth}/\text{range}$
no_e_dependence =	.false.	if true, then p_{impact} is adjusted with a factor energy_0/E . This eliminates an energy dependence in cross section for CERDA and EBS. If false, then max. p_{impact} is chosen assuming up to 70% residual projectile energy after passing the target.
num_scattering =	1.0	Number of large angle scattering events per path of a projectile through the target
scatt_species(1:num_species) =	'---'	num_species ASCII characters identical to Symbol, specifying for which collision partner the scattering is enforced. Example: "H" selects collisions with H-Atoms e.g. in a compound TiH ₂ . If the first string is set to "all", then large angle scattering applies to a target elements Isotopes must no be specified, just the name of the elemental species
non_rutherford	.false.	If .true., then non Rutherford cross sections will be used

Command	Default values	Description
file_r33	none	An IBANDL non-RBS cross section or NRA files up to 3 files per target species can be used, total 3*ncpm files
nra_species = nra_product =		Up to max number of element species, number of target and corresponding product elements for NRA simulations. Example: nra_species = "Li7","Li6" nra_product="He4","He3" The nra products must also be specified in the list of target elements, symbol= ... !
l_threshold=	.false.	(default: .false.) if true, then all events with energy above e_thres_r are ignored, i.e. only recoils with energies above the threshold are considered. Also projectile data output can be selected according to e_thresh If .false, then E_displ is only used to reset recoils to its initial position if Ekin < E_displ.
lpart_r_ed=	.false.	if true, then do not write recoil particle info data to file partic*.dat if initial recoil energy < E_displ. This avoids storage of too many low energy collision events
lpart_pb_ed=	.false.	if true, then do not write backscattered projectile (pb) info data to file partic*.dat if energy transfer to recoil is DE < e_thresh_r. This avoids storage of too many low energy collision events
lpart_pt_ed=	.false.	if true, then do not write transmitted projectile (pt) info data to file partic*.dat if energy transfer to recoil is DE < e_thresh_r. This avoids storage of too many low energy collision events
loutgas=	.false.	calculation with outgassing transport and diffusion (see also: diff_koeff1, diff_koeff2)

5.1.8 Treatment of vacancies and noble gas atoms

Command	Default values	Description
vacancy_mode	1	<p>How to calculate p_{vac} with $0 \leq q_{vac} \leq 1; 0 \leq c_{vac} \leq 1$:</p> <p>0: vacancy_mode = 0 no vacancy generation at all, also no vacancies are annihilated</p> <p>1: default quadratic - $p_{vac} = q_{vac} \cdot (1 - c_{vac})^2$</p> <p>2: quadratic + $p_{vac} = q_{vac} + (1 - q_{vac}) \cdot (1 - (1 - c_{vac})^2)$</p> <p>3: linear - $p_{vac} = q_{vac} \cdot (1 - c_{vac})$</p> <p>4: linear + $p_{vac} = q_{vac} + c_{vac} (1 - q_{vac})$</p> <p>5: cubic - $p_{vac} = q_{vac} \cdot (1 - c_{vac})^3$</p> <p>6: cubic + $p_{vac} = q_{vac} + (1 - q_{vac}) \cdot (1 - (1 - c_{vac})^3)$</p> <p>7: sine - $p_{vac} = q_{vac} \cdot \frac{1}{2} (1 + \sin(\pi(c_{vac} + 0.5)))$</p> <p>8: sine + $p_{vac} = q_{vac} + (1 - q_{vac}) \cdot \frac{1}{2} (1 - \sin(\pi(c_{vac} + 0.5)))$</p> <p>9: constant $p_{vac} = q_{vac}$</p> <p>Other values: mode 0</p>
q_vacancy (1:num_species) =	1.0	<p>characteristic probability for vacancy formation. $0 < q_{vacancy} < 1$. Formation of vacancies given by the probability and the local vacancy concentration c_v. Default mode vacancy_mode = 1:</p>
i_vac_coord=	4	<p>coordination number of atom stop positions. If one of the neighboring atoms is a vacancy, then a stopped particle annihilates the vacancy. Annihilation probability saturates at $\min(1, \text{atomic_fraction} \times (\text{vacancy}) * i_vac_ccord)$</p>

Command	Default values	Description
vacancy_stopping_fraction	1.0	Fraction of the elemental stopping power contributing to electronic stopping of vacancies
vacancy_species=	none	Target elements used for electronic stopping calculation for vacancies. Up to ncp elements can be specified and are selected based on the global atomic fraction and the value q_vacancy. If nothing is specified, then only the major target element is used for vacancy stopping calculation.
vacancy_sequence	0	Anti coincidence_mode: -1 , then not any two vacancies are chosen in a sequence, except there is a layer filled with vacancies to larger than 80%. This mode is used to satisfy the stacking sequence of graphene layers Sequence_mode: = n integer number. Then up to n vacancies are selected in a sequence, except there is a layer which does not contain vacancies. This mode is used tp satisfy the stacking of Metal-Dichalcogenide triple-layers.
lgaserf=	.true.	activates a surface error function profile for qmax of vacancies ensures that qmax for vacancies goes to zero near the surface. Default value is .true.
gaserf_sigma =	2.	width of error function in unit of layers for noble gas concentration profile (default 2 layers)
gaserf_pos =	4.	layer position of error function center for noble gas concentration profile (default: layer 4)
lvacerf=	.true.	activates a surface error function profile vor qmax of vacancies and ensures that qmax for vacancies goes to zero near the surface. Default value is .true.
vacerf_sigma =	2.	width of error function in unit of layers for vacancy concentration profile (default 2 layers)
vacerf_pos =	4.	layer position of error function center for vacancy concentration profile (default: layer 4)

6 Default filenames and directories

These names are set in subroutine default_init.F90. filenames have a maximum default length of 50 characters and directory names a default maximum length of 100 characters

```
! SET MAIN PROGRAM NAME and PROGRAM VERSION
!
  programname='imintdyn821'
  programversion='Ver 8.2100'
!
! -----
! SET input and output filenames and directories
!
  imint_out_target='imint_output_target-000.dat'
  imint_out_moments='imint_output_moments-000.dat'
  imint_out_moments1='imint_output_momentsA-000.dat'
  imint_out_sputter='imint_output_sputter-000.dat'
  trajec_all='trajec_all.dat'
  trajec_out(1)='trajec_stop.p.dat'
  trajec_out(2)='trajec_back_p.dat'
  trajec_out(3)='trajec_tran_p.dat'
  trajec_out(4)='trajec_stop_r.dat'
  trajec_out(5)='trajec_back_r.dat'
  trajec_out(6)='trajec_tran_r.dat'

  partic_out(1)='partic_stop_p.dat'
  partic_out(2)='partic_back_p.dat'
```

```
  partic_out(3)='partic_tran_p.dat'
  partic_out(4)='partic_stop_r.dat'
  partic_out(5)='partic_back_r.dat'
  partic_out(6)='partic_tran_r.dat'
```

```
  series_filename='serie.dat'
  energy_analysis='energy_analysis.dat'
  timestamp='time.dat'
  time_run='time_run.dat'
  rf_filename1='Rstart_1.inp'
  rf_filename2='Rstart_2.inp'
```

```
  output_filename='output-000.dat'
```

```
  craterfile='craterfunction.dat'
  cratermoments='cratermoments.dat'
  balancefile='balancefile.dat'
  impl_crater_moments='crater-moments-implantation.dat'
```

```
  zero_moments='zero-moments-file.dat'
```

vacancyfile='vacancyfile.dat'

file_elosstable='energy_loss_table0.dat' ! 0 = 1,num_species
file_pimpact_angle='pimpact_vs_angle.dat'

layerinput='layer.inp' ! not used at the moment
layeroutput='layer_profile.dat'
layeroutinp='layer_profile.def
dir_layerinp = './'
file_layerinp = 'layer.def'
dir_energyinp = './'
file_energyinp = 'energy.def'
file_energyout_test='energyout_test.dat'
dir_angleinp = './'
file_angleinp = 'angle.def'
file_angleout_test='angleout_test.dat'
dir_tableinp = '.././tables/'

! SRIM data are in default subdirectory 'SRIM-tables'

e_act_HCW = 'e_act_HCW.inp'
a0_HCW = 'a0_HCW.inp'

scoef95a = 'SCOEf.95A'
scoef95b = 'SCOEf.95B'

SRIMfile='SRIM2013-nn.dat'
dir_SRIMinp='SRIM_tables/'

! filenames used in subroutine "output"

meagb_p_file = 'matrix_energy_angle_back_proj.dat'
meagt_p_file = 'matrix_energy_angle_trans_proj.dat'
meagb_s_file = 'matrix_energy_angle_back_sputt.dat'
meagt_s_file = 'matrix_energy_angle_trans_sputt.dat'
mepb_p_file = 'matrix_energy_path_back_proj.dat'
mept_p_file = 'matrix_energy_path_back_proj.dat'
morigin_ex_bs = 'matrix_energy_depthorig_back_sputt.dat'
morigin_ex_ts = 'matrix_energy_depthorig_trans_sputt.dat'
E_distr_stop = 'energy_distribution_stop'
E_distr_inel = 'energy_distribution_electronic'
E_distr_nucl = 'energy_distribution_nuclear'
E_distr_all = 'energy_distribution_all'
depth_proj = 'depth_distribution_projectiles.dat'
depth_recoil = 'depth_distribution_recoils.dat'

!

! matrix filename in subroutine "histories"

!

angular_matrix_file = 'meagb_s_000000.dat'

!

! =====

io numbers used for different input and output files:

io = 6 !6 Standard output

io7 = 7 !6 time output

io10 = 10 !
io21 = 21 ! --- angle_input,energy_input
io22 = 22 ! --- table_read
io22 = 22 ! --- read table.compound
io23 = 23 ! --- ioSCOEf_95A Ziegler Biersack stopping power table
io24 = 24 ! --- ioSCOEf_95B Ziegler Biersack stopping power data
io29 = 29 ! ---general output
io30 = 30 ! ---imint.inp or imintb.inp
io32 = 32 ! ---crater function file output
io31 = 31 ! ---general output, series_filename, imint_out_target
io33 = 33 ! ---balance file output
io34 = 34 ! ---crater function moments output
io35 = 35 ! ---implantation crater function output

io35 = 35 ! --- imint_out_moments1,
! --- E_distr_stop.dat, _distr_inel.dat,E_distr_nucl.dat,E_distr_all.dat

io36 = 36 ! -- zero-moments-file output
io37 = 37 ! -- dimer-sputtering file output

io41 = 41 ! ---energy analysis, depth proj, depth recoil
io50 = 50 ! --- output file for energy loss tables
io55 = 55 ! ---layer input and output data

io56 = 56 ! ---layer output data for layer.inp
io57 = 57 ! ---matrix output
io67 = 67 ! ---matrix output
io67 = 76 ! ---matrix output

! these 7 io values should be in a row:
io80 = 80 ! --- only as dummy for io80 + j

io81 = 81 ! --- trajec_stop_p
io82 = 82 ! --- trajec_back_p
io83 = 83 ! --- trajec_tran_p
io84 = 84 ! --- trajec_stop_r
io85 = 85 ! --- trajec_back_r
io86 = 86 ! --- trajec_tran_r

! these 6 io values should be in a row:

io90 = 90 ! --- only as dummy for io90 + j
io91 = 91 ! --- partic_stop_p
io92 = 92 ! --- partic_back_p
io93 = 93 ! --- partic_tran_p
io94 = 94 ! --- partic_stop_r
io95 = 95 ! --- partic_back_r
io96 = 96 ! --- partic_tran_r

io321 = 321 !
io331 = 331 ! vacancyfile, imin_out_sputter
io341 = 341 ! imint_out_moments
io351 = 351 ! imint_out_moments1
io100 = 100 ! --- time_run.dat, rw restart

7 Sample input file (imint-template.inp)

```
1 keV Ar -> Si + Vac test simulation
&IMINT_INP

!      debugging= nn          ! set debugging flag to level nn 0,..10, to print output infos
!      debugging_history = 1,10
!      debugging_pid= 1,4
!      impactflag=.true.    ! create a file impact parameter vs. scattering angle

energy_mode = "none"          ! use a constant energy
angle_mode = "none"          ! us a constant incidence angle
series_mode = "none"
series_steps = 1
curvature_flag=.false.
!
!      set projectile and target element definitions:

target_thickness = 1000E+0    ! target depth (Angstroems)
target_slices    = 500E+0    ! number of target intervals

symbol =          "Ar",      "Si",      "Vac"    ! names of projectile and target elements
atomic_density = 0.05,      0.05,      0.05    ! set atomic densities
energy0 =         1000,      0.00,      0.00    ! ion energy (eV)
!energy0_inc =    0.,        0.,        0.,
alpha0 =          0.0,        0.0,        0.0    ! incidence angle (deg)
!alpha0_inc =     5.0 ,       0.0,        0.0
!phi0 =           0.0 ,       0.0,        0.0    ! azimuthal angle (deg)
!x0 =             0.,        0.,        0.0    ! origin of impact [A]
```

```
stopping_mode =      7,          7,          7,
atomic_fraction =   0.0,          1.0,          0.0      ! initial concentration in the target
beam_fraction =     1.000,        0.000,        0.000      ! fraction of incident beam

max_atomic_fraction =0.2,          1.0,          1.0      ! max concentration
charge =             1. ,          0.0,          0.00      ! electrical charge
e_cutoff =           1.0,          1.0,          1.0      ! cut off energies [eV]
e_surfb =            0.0,          4.7,          0.0      ! surface binding energies [eV]
e_displ =            0.0,          0.0,          0.0      ! displacement energies [eV]

q_vacancy =          0.7,          0.7,          0.7,      ! characteristic probability for vac formation

!fil_angleinp='angle_test.inp'

Kij = 0.00, 0., 0.      ! define the surface curvature [1/A]

Projectiles = 2.eE5
! histories = 10000,      ! number of histories
! Projectiles_per_history = 24
fluence_steps_out = 100
fluence = 1      ! maximum fluence in units 1E16/cm²

dynamic_simulation =.true.  ! SBE model
! static_simulation=.false.
! cascade_simulation=.false. ! always true in dynamic mode
surf_inel_loss = 0
potential="KrC"      ! "KrC", "Moliere", "ZBL", "Nakagawa"-Yamamura, "Si-Si", "power"
integration_mode = 2      ! 0 = Magic, 1 = Gauss-Mehler, 2=Gauss-Legendre
! 3 = Gauss-Legendre 8 (double precision)
```

```
subthr_rec_bound = .true.! sub threshold atoms are bound

weak_coll_p = 2,          ! nr of weak collisions for projectiles
weak_coll_r = 2,          ! number of weak collisions for recoils

lpart_r_ed =.true.       ! store only events for E > E_displ

!enforce_scattering=.true. ! enforce large angle scattering
!num_sacattering = 1.     ! scattering events per impact
!pmax_coeff = 1E-4       ! reduction factor for pmax value
!scatt_species ="Si"     ! restrict large angle scattering to this element

!free_path_length=.true. ! enable larger free_path length
!ffp_accuracy =0.01      ! accuracy is 1% rel energy loss or 1 deg max deflection
!
i_vac_coord = 1         ! coordination number around stopped atoms
vacany_stopping_fraction = 1.0
lvacerf = .true.       ! erf profile for qumax(num_species_vacancy)
lgaserf = .true.       ! erf profile for qumax(num_species_noblegas)
vacerf_pos = 6.        ! center layer position of error function profile
vacerf_sigma = 3.      ! width of error function profile
gaserf_pos = 6.        ! center layer position of error function profile
gaserf_sigma = 3.      ! width of error function profile

lmatrices = .false. ! matrix output

ltraj_p = .false. ! projectile trajectories
ltraj_r = .false. ! recoil trajectories
```

```
! number of traced trajectories for:
! stopped, backscattered and transmitted projectiles, stopped, backspattered,
! transmission sputtered recoils
ioutput_trajectories = 5, 5, 0, 5, 5, 0,
lparticle_r = .false.    ! create projectile data files
lparticle_p = .false.    ! create recoil data files

! number of traced particles for:
! stopped, backscattered and transmitted projectiles, stopped, backspattered,
! transmission sputtered recoils
!
ioutput_particles = 100000, 100000, 0, 100000, 100000, 0,

/
```

8 Example for a shell script file (imint-template.sh)

```
#!/bin/bash
DIRECTORY=$(pwd)
ERRCODE=0

TM='-off'
#TM='-testmode' # if set, then IMINTDYN is run in testmode

# list of input files imint-namex.inp
INPUT=(name1 name2) # primary input script file, default: "imint-name1.inp"
INPUTB=(subname1 subname2) # 2nd input script file, overwrites data from imint.inp, default: "subname1.inp"
# list of input files namey.def !!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!
SUPPORT=(support1 support2) # example: layer definition file "sample_layer.def"

POST="none" # ebs, hrrbs, hrebs, esa, tofrbs, ercs, erda, toferda, cerda, matrix or none post processing flag
BINSIZE=2048 # binning size of postprocessing spectra
SMOOTH=2 # smoothing of postprocessing spectra

WORK=temp # temporary directory

if [ $TM = "-testmode" ]; then
WORK=temptest
else
WORK=temp # temporary directory
fi # - use a special temp folder for testmode not to mess up with data in the temp folder

# set number NP of processors and simulation mode parallel PAR, or sequential SEQ
echo =====
if test -d /opt/intel/oneapi; then
echo "Intel OneAPI compiler"
COMPILER=oneapi
NP=24
```

```

    PPN=48
    if [ $TM = "-testmode" ]; then
    NP=1
    PPN=48
    fi
else
    echo "Older Intel Compiler"
    COMPILER=intel
    NP=4
    PPN=8
    if [ $TM = "-testmode" ]; then
    NP=1
    PPN=8
    fi
fi
MODE=PAR # PAR,SEQ

echo =====
echo start new simulation

echo case directory is $DIRECTORY
echo input script files: ${INPUT[*]}
echo inputB script files: ${INPUTB[*]}
echo support files: ${SUPPORT[*]}.def
echo post processing programs: ${POST[*]}
echo simulation mode is "$MODE"
echo number of processors "$NP"
echo ----- setup finished -----

# process all input files
index=-1
for name in ${INPUT[*]} # -----i---
do
    index=$((index+1))
    nameb=${INPUTB[$index]}
    cd $DIRECTORY # change to specific case directory
    file=imint-${name}.inp
    fileb=${nameb}.inp

```

```

echo =====
if [ -f "$file" ]; then # if file exists, then
  echo input script file $file exists, start processing !
  nameout=${name}
  echo =====

  if [ -f "$fileb" ]; then # if file exists, then
    echo input script file $fileb also exists !
    nameout=${name}-${nameb}
    echo =====
  fi
  echo output directory name is: $nameout
  cd .. # go back to case directory
  CASEDIR=$(pwd)
  # create directory temp-namout or temptest if it does not exist
  if [ $TM = "-testmode" ]; then
    mkdir -p "$WORK"
    cd $WORK
  else
    mkdir -p "$WORK"- "${nameout}"
    cd $WORK-${nameout}
  fi

  # clean up working directory if not empty
  if [ -z "$(ls -A )" ]; then # -----if--
    echo directory $WORK-${nameout} is empty
  else
    echo delete als files in directory $WORK
    rm --force *.* # force delete files
  fi # -----end if --
  DEFAULT=$(pwd) # now use this temporarydirectory as the default one
  echo working directory is: $DEFAULT
  cd $DIRECTORY # change to specific case directory

cp $file $DEFAULT/imint.inp
# copy related angle, energy and layer definition files
# file extensions are typically .inp or.def
if [ -f "$fileb" ]; then # if file exists, then

```



```

    cp $fileb $DEFAULT/imintb.inp
    echo 2nd input file $fileb was copied
    fi

for j in ${SUPPORT[*]} # -----j-----
do
cp ${j}.def $DEFAULT/${j}.def
done # -----j-----
echo support files copied if available
cd $DEFAULT
# run parallel on NP processors , or sequential
if [ $MODE = "PAR" ] # -----if ---
then
echo ----- start mpirun in parallel mode -----
    if [ $COMPILER = "oneapi" ]; then
        mpirun -n "$NP" -ppn "$PPN" -f ../../bin/linux.PRO/hostfile ../../bin/linux.PRO/imintdyn.exe "$TM"
        ERRCODE=$?
    elif [ $COMPILER = "intel" ]; then
        mpirun -n "$NP" ../../bin/linux.PRO/imintdyn.exe "$TM"
        ERRCODE=$?
    fi
else
echo ----- start mpirun in sequential mode -----
    ../../bin/linux.SEQ/imintdyn.exe "$TM"
    ERRCODE=$?
fi # -----end if---
echo ===== imintdyn program execution finished =====
echo error code = $ERRCODE
    if [ $TM != "-testmode" ]; then
        mkdir -p $DIRECTORY/$nameout
        cp *.* $DIRECTORY/$nameout
        rm *.*
        echo ===== data copied to $DIRECTORY =====
        cd $CASEDIR
        rmdir "$WORK"-"${nameout}"
        echo ===== temporary directory deleted =====
        cd $DIRECTORY
    else

```

```

        echo directory $WORK contains test output data !!
    fi

    if [ $TM != "-testmode" ]; then
    if [ $ERRCODE = "0" ]; then # -----if---
        echo ===== start postprocessing program read_imintdyn_target.exe =====
# goto results directory and start the post processing programs
        cd $DIRECTORY
        cd ${nameout} # goto sub-directory for post processing
# create all files of postprocessing_data

        ../../../../post/read_imintdyn_target.exe

# create all matrix file of angular and energy distributions of emitted particles
        if [ $POST = "matrix" ] ; then
            echo ===== start postprocessing program read_imintdyn_matrix.exe =====
            ../../../../post/read_imintdyn_matrix.exe
            elif [ $POST = "ebs" ] || [ $POST = "rbs" ] ; then
# create the outebs.dat file of energy histograms for RBS or non-Rutherford BS
# options: -s s1 size of histogram; -a a1,a2 angluar regime
            echo ===== start postprocessing program histogram_ebs.exe =====
            ../../../../post/histogram_ebs.exe -b $BINSIZE -s $SMOOTH
            elif [ $POST = "hrrbs" ] || [ $POST = "hrebs" ] || [ $POST = "esa" ] ; then
# create the outebs.dat file of energy histograms for ESA detector resolution default 0.005
# options: -s s1 size of histogram; -a a1,a2 angluar regime -esamode 0.005
            echo ===== start postprocessing program histogram_ebs.exe with ESA mode =====
            ../../../../post/histogram_ebs.exe -esamode 0.005 -b $BINSIZE -s $SMOOTH
            elif [ $POST = "tofrbs" ] ; then
# create the outebs.dat file of energy histograms for TOF detector resolution default 300ps 1m
# options: -s s1 size of histogram; -a a1,a2 angluar regime -tofmode 300.,1.
            echo ===== start postprocessing program histogram_ebs.exe with ESA mode =====
            ../../../../post/histogram_ebs.exe -tofmode 300 1.0 -b $BINSIZE -s $SMOOTH
            elif [ $POST = "cerda" ] ; then
# create the outcerda.dat file of energy histograms of coincidence CERDA events
# options: -s s1 size of histogram; -a a1,a2 angluar regime
            echo ===== start postprocessing program histogram_cerda.exe =====
            ../../../../post/histogram_cerda.exe -b $BINSIZE -s $SMOOTH -c $NP
            elif [ $POST = "ercs" ] ; then

```

```

# create the outerecs.dat file of energy histograms of coincidence ERCS events
# options: -s s1 size of histogram; -a a1,a2 angluar regime
      echo ===== start postprocessing program histogram_ercs.exe =====
      ../../../../post/histogram_ercs.exe -b $BINSIZE -s $SMOOTH -c $NP
      elif [ $POST = "erda" ] ; then
# create the outerda.dat file of energy histograms of ERDA events
# options: -s s1 size of histogram; -a a1,a2 angluar regime
      echo ===== start postprocessing program histogram_erda.exe =====
      ../../../../post/histogram_erda_nra.exe -b $BINSIZE -s $SMOOTH
      elif [ $POST = "toferda" ] ; then
# create the outerda.dat file of energy histograms of TOF-ERDA events 300ps, 1m
# options: -s s1 size of histogram; -a a1,a2 angluar regime
      echo ===== start postprocessing program histogram_erda.exe =====
      ../../../../post/histogram_erda_nra.exe -tofmode 300., 1.0 -b $BINSIZE -s $SMOOTH
      elif [ $POST = "nra" ] ; then
# create the outerda.dat file of energy histograms of NRA events
# options: -s s1 size of histogram; -a a1,a2 angluar regime
      echo ===== start postprocessing program histogram_erda.exe =====
      ../../../../post/histogram_erda_nra.exe -nramode 0
      fi # -----endif-----
echo ===== all done without error =====

      elif [ $ERRCODE > "0" ]; then # -----if---
      echo ==== Because of ERRCODE no post processing done ====
      fi # -----end if ---
      fi # --- endif testmode
      cd $DIRECTORY
else
      echo =====
      echo $file does not exist!!
      echo =====
fi
done

```

9 Example of an input file for an angular distribution (template-angle.inp)

```
template for angular distribution as input (may be generated with ORIGIN)
angle[deg] (0 < alpha < 90deg), probability(not normalized), up to 180 lines
28 0
28.5 0
29 0
29.5 0
30 1
30.5 1
31 1
31.5 2
32 3
32.5 4
33 5
33.5 7
34 8
34.5 11
35 13
35.5 16
36 19
36.5 23
37 27
37.5 32
38 37
38.5 42
39 48
39.5 54
40 60
40.5 66
```

41	72
41.5	78
42	83
42.5	88
43	92
43.5	95
44	98
44.5	99
45	100
45.5	99
46	98
46.5	95
47	92
47.5	88
48	83
48.5	78
49	72
49.5	66
50	60
50.5	54
51	48
51.5	42
52	37
52.5	32
53	27
53.5	23
54	19
54.5	16
55	13
55.5	11

56 8
56.5 7
57 5
57.5 4
58 3
58.5 2
59 1
59.5 1
60 1
60.5 0
61 0
61.5 0
62 0
62.5 0
63 0
63.5 0
64 0
64.5 0
65 0

10 Example of an input file for an energy distribution (template-energy.inp)

```
template for energy distribution input (may be generated with ORIGIN
energy [eV] (0.1 eV < E > 1E9 eV), probability (not normalized), up to 200 lines
50.  1.
100. 2.
120. 5.
150. 7.
400. 12.
500. 17.
800. 22.
1000. 27.
1200. 35.
1500. 54.
1800. 76.
2400. 81.
2500. 74.
2800. 52.
2900. 34.
3000. 26.
3100. 15.
3400. 12.
3500. 5.
3600. 2.
3700. 1.
```

11 Example of an input file for a layered structure (template-layer.def)

This is an example of a multilayer where each layer has a thickness of Å. We define a structure for three species 1,2,3. We start with two layers of species 2, followed by 3 layers of species 3, followed by 2 layers of species 2, etc. At the end we have 50 layers of species 2.

We have an alternating composition of 100% for element species 2 and 3. Element species 1 is supposed to be the incident ion and has initial composition 0.

Note that the number of layers must be given as Integer and the other values as floating point values (which must contain a decimal ".").

In total there are 65 layers distributed over the target thickness $\text{target_thickness} = 65 \times 3 \text{ \AA} = 195 \text{ \AA}$. -1 in the last line indicates the end of the layered structure. If this is replaced by 0, then the previous layered structure is repeated until the maximum of target_slicesmax layers are filled. Based on the input, the new target thickness target_thickness with the sum target_slices of the specified layers is calculated.

```
number n of      thick-  composition of target (1:num_species); sum( qu(1:num_species)) = 1
layers          ness [A]  1    ... num_species ; n<0: end, n=0: copy until target_slicesmax
  2             3.00     0.00  1.00  0.00
  3             3.00     0.00  0.00  1.00
  2             3.00     0.00  1.00  0.00
  3             3.00     0.00  0.00  1.00
  2             3.00     0.00  1.00  0.00
  3             3.00     0.00  0.00  1.00
  50            3.00     0.00  1.00  0.00
  -1
```

In contrast to SDTrimsP, the line with "-1" finished the layer profile input. No further values are required in that line. A value "0" in the last line copies the last layer until the maximum number of layers is reached.

12 Compiling the program

The Make file mk.sh:

```
#!/bin/sh -x
#
# bourne shell compatible script. We can't be sure that it's a bourne shell,
# it can be any shell depending on where /bin/sh soft link is pointing to usually.
# For instance, With the latest Ubuntu "sh" by default pointing to "dash".
# -x
#
SRC_DIR=../../src
make clean
# if makefile does not exist as symbolic link, then make a link to src directory
if [ ! -L Makefile ]; then
    ln -s $SRC_DIR/Makefile Makefile
fi
#
#           OSTYPE= linux
#           RAND=RAND_INTEL or RAND=RAND_CRAY or RAND=NR_RANDQ1 random number generation
#           COMPILER=INTEL or COMPILER=ONEAPI (new intel compiler) select the FORTRAN compiler

echo =====
if test -d /opt/intel/oneapi; then
    echo "Intel OneAPI compiler"
    FC='ONEAPI'
else
    echo "Older Intel Compiler"
    FC='INTEL'
fi
echo =====

make OSTYPE=linux SRC_DIR=$SRC_DIR RAND=NR_RANDQ1 COMPILER=$FC -e -f $SRC_DIR/Makefile
# -e overrides environment variables
# -f specifies the location of Makefile
```

Makefile:

```
EXEC = imintdyn.exe

# Versions can be compiled for 64 bit machines from the source code:
# MODE = PAR parallel processing
# Furthermore random number grnerators can be chosen:
# RAND = ( CRAY | NR_RANDQ1 | RAND_INTEL |
# Possible communication libraries are:
# COMM = ( MPI )
# Possible compiler for linux are:
# COMPILER = ( INTEL | ONEAPI)
# OSTYPE: LINUX: linux

MODE = PAR
COMM    = MPI
SRC_DIR = .
VPATH  = $(SRC_DIR) # use this path to find certain files
#
# catch error if OSTYPE is not defined:
#
FF      = @echo "makefile not configured for OSTYPE \"$(OSTYPE)\"; false

### =====
### ---  Macros for Linux PC -----
ifeq ($(OSTYPE), linux)

# COMPILER = INTEL
# COMPILER = ONEAPI (New Intel Compiler)
```

```

#---INTEL ONEAPI ---
  ifeq ($(COMPILER), ONEAPI)
    #---openMPI
    FF= mpiifx # New OneAPI Intel Fortran Compiler

    ##FFLAGS = -v 9.0 -w -FR -r8 -O2 -ip -tpp7 -xN
    ##FFLAGS = -v 9.0 -w -FR -r8 -O2 -ip -tpp6 -tune pn4 -pg -pc64 -pc80 -check all
    ## -pg -check all
    ## -Fr source file free format
    ## -w disables all warnings
    ## -axN optimisation for all computer
    ## -xN optimisation for computer where compile
    ## -check none
    ## -check noarg_temp_created
    ## -r8 use double precision as default, replaced by -double-size
    ## -real-size 64 8 byte real an complex declarations

FFLAGS =-w -FR -real-size 64 -O3 -align=all -ansi -axcore=CORE-AVX2,SSE4.2,SSSE3 \
  -falign-functions -fast-transcendentals -finline -no-inline-factor -fp-model=fast=2 -ip \
  -ipo -mcode=avx2,sse4.2,ssse3 -march=x86-64-v3 -mtune=core-avx2 -qopt-matmul \
  -qopt-mem-layout-trans=3 -qopt-multi-version-aggressive -qopt-prefetch=4 -pad -rcd -safe-cray-ptr \
  -scalar-rep -unroll-aggressive -parallel -qopenmp -g -traceback \
  -I$(SRC_DIR) -I/usr/local/lib -D$(OSTYPE) -D$(COMPILER)

#   FFLAGS =-w -FR -real-size 64 -O3 -align=all -ansi-alias -assume buffered_io -axcore=CORE-AVX2,SSE4.1,SSSE3 \
#   -falign-functions -fast-transcendentals -finline -no-inline-factor -fp-model=fast=2 -ip \
#   -ipo -mcode=avx2,sse4.1,ssse3 -march=core-avx2,sse4.1,ssse3 -mtune=core2 -qopt-matmul \
#   -qopt-mem-layout-trans=3 -qopt-multi-version-aggressive -qopt-prefetch=4 -pad -rcd -safe-cray-ptr \
#   -scalar-rep -unroll-aggressive -xSSSE3 -parallel -qopenmp -g -traceback \
#   -I$(SRC_DIR) -I/usr/local/lib -D$(MODE) -D$(OSTYPE) -D$(DEBUG)

endif # endif of compiler OneAPI INTEL

```

```
#---INTEL---
ifeq ($(COMPILER), INTEL)

    #-openMPI
        FF= mpiifort # New Intel Fortran Compiler
#        FF= mpif90 # (Obelix and Old Intel Fortran compiler)
#        FF= mpifort # (GWDG Intel compiler)

##FFLAGS = -v 9.0 -w -FR -r8 -O2 -ip -tpp7 -xN
##FFLAGS = -v 9.0 -w -FR -r8 -O2 -ip -tpp6 -tune pn4 -pg -pc64 -pc80 -check all
## -pg -check all
## -Fr source file free format
## -w disables all warnings
## -axN optimisation for all computer
## -xN optimisation for computer where compile
## -check none
## -check noarg_temp_created
## -r8 use double precision as default, replaced by -double-size
## -real-size 64 8 byte real an complex declarations

FFLAGS =-w -FR -real-size 64 -O3 -align=all -ansi-alias -assume buffered_io -axcore=CORE-AVX2,SSE4.1,SSSE3 \
        -falign-functions -fast-transcendentals -finline -no-inline-factor -fp-model=fast=2 -ip \
        -ipo -mcode=avx2,sse4.1,ssse3 -march=core-avx2,sse4.1,ssse3 -mtune=core2 -opt-matmul \
        -opt-mem-layout-trans=3 -opt-multi-version-aggressive -opt-prefetch=4 -pad -rcd -safe-cray-ptr \
        -scalar-rep -unroll-aggressive -xSSSE3 -parallel -openmp -g -traceback \
        -I$(SRC_DIR) -I/usr/local/lib -D$(OSTYPE) -D$(COMPILER)

endif # endif of compiler INTEL
```

```
# -- set some compiler flags -----  
  
ifeq ($(RAND), NR_RANDQ1)  
    FFLAGS += -D$(RAND)  
endif  
ifeq ($(RAND), RAND_INTEL)  
    FFLAGS += -D$(RAND)  
endif  
ifeq ($(RAND), RAND_CRAY)  
    FFLAGS += -D$(RAND)  
endif  
ifeq ($(COMM), MPI)  
    FFLAGS += -D$(COMM)  
endif  
  
#-----OneAPI INTEL-----  
ifeq ($(COMPILER), ONEAPI)  
  
    LDR = $(FF) $(FFLAGS)  
    #LIBS = -static  
  
endif  
  
#-----INTEL-----  
ifeq ($(COMPILER), INTEL)  
  
    LDR = $(FF) $(FFLAGS)  
    #LIBS = -static  
  
endif
```

```

endif #$(OSTYPE), linux)

### -----
MODULES = MPIfunc.o parameters.o random_numbers.o task_description.o particle_description.o \
          trajectory_description.o work.o dlb.o zb_stoppingtable_data.o zb_stopping_calc.o \

BINS = imintdyn.o history.o recoil.o projectile.o vfmt.o output.o sub.o \
       sub_chem_diff.o inelast.o assign.o zero_init.o interpolate_fractions.o \
       pots.o integrations.o tableread.o read_r33.o cputim.o pimpactscan.o \
       default_init.o gsum_fluence.o gsum_stat.o gsum_mat.o \
       out_particles.o out_trajectories.o out_datfile.o init_all.o \
       iba_init.o main_broadcast.o allocate_memory.o set_matrix_ranges.o t3e.o\

default: $(EXEC)

$(EXEC): $(BINS)
        $(LDR) -o $(EXEC) $(MODULES) $(BINS) $(LIBS)

sub.o: work.o random_numbers.o
sub_chem_diff.o: work.o MPIfunc.o
random_numbers.o: work.o MPIfunc.o
imintdyn.o: work.o parameters.o random_numbers.o alloc_types.o zb_stoppingtable_data.o MPIfunc.o \
           trajectory_description.o particle_description.o
zero_init.o: work.o parameters.o
default_init.o: work.o parameters.o
history.o: work.o random_numbers.o parameters.o dlb.o MPIfunc.o trajectory_description.o \

```

```
particle_description.o sub_chem_diff.o
pimpactscan.o: work.o MPIfunc.o
recoil.o: work.o random_numbers.o parameters.o dlb.o MPIfunc.o
projectile.o: work.o random_numbers.o parameters.o dlb.o MPIfunc.o
gsum_fluence.o: work.o MPIfunc.o parameters.o
gsum_stat.o: work.o MPIfunc.o parameters.o
gsum_mat.o: work.o MPIfunc.o parameters.o
out_particles.o: work.o MPIfunc.o
out_trajectories.o: work.o MPIfunc.o output.o: work.o
task_description.o: MPIfunc.o parameters.o
particle_description.o: parameters.o
work.o: particle_description.o task_description.o trajectory_description.o vfmt.F90
dlb.o: task_description.o MPIfunc.o
inelast.o: work.o zb_stopping_calc.o
out_datfile.o: work.o MPIfunc.o
tableread.o: work.o MPIfunc.o zb_stoppingtable_data.o
read_r33.o: work.o MPIfunc.o
init_all.o: work.o parameters.o MPIfunc.o random_numbers.o
iba_init.o: work.o MPIfunc.o parameters.o
main_broadcast.o: work.o MPIfunc.o parameters.o
allocate_memory.o: work.o parameters.o MPIfunc.o dlb.o
set_matrix_ranges.o: work.o parameters.o MPIfunc.o
interpolate_fractions.o: work.o random_numbers.o dlb MPIfunc.o
pots.o: work.o

.SUFFIXES:
.SUFFIXES: .o .F90 .f90 .c
.F90.o:
$(FF) -c $(FFLAGS) $<
.f90.o:
```

```
$(FF) -c $(FFLAGS) $<
.PHONY: clean update
clean:
    rm -v -f *.[fio] *.mod $(EXEC) aus
```

13 Running the program

IMINTDYN simulations are defined in a sub-folder called “case” in the main folder “imintdyn81”. There each simulation is stored in a specific subfolder, e.g. “1 keV Ar in Si”. This subfolder contains all simulation specific information, such as the main simulation script file “imint-namex” and 2nd order script file “namey”. In addition there might be supporting files such as angular profiles, energy profiles or a file defining a layered target structure. These supporting files are text files and have the extension “.def.”. The script files “imint-namex.inp” and “namey.inp” are read using the FORTRAN namlist feature, which is a easy but rough way to initialize a simulations. All variables defined in the .inp files must be correctly specified in a way they are defined in the imintdyn subroutine work, or specified in the imintdyn command parameter description. Any other variable definitions or typing errors lead to an error when reading the namelist.

A script file “imint.sh” contains all information on how to run the program and how to do the post-processing of output files. Within imint.sh one defines the script files “imint-namex” and “namey” as well as the supporting files. Several files “imint-namex” and “namey” can be defined which are then processed as a batch process. One also defines the type of post processing such as “ebs”, “nra”, “erda” etc.

Some parameters for postprocessing can be defined in “imint.sh”, such as

```
POST="ebs" # ebs, hrrbs, hrebs, esa, tofrbs, ercs, erda, toferda, cerda, nra and matrix or none post processing flag
BINSIZE=2048 # binning size of postprocessing spectra
SMOOTH=2 # smoothing of postprocessing spectra
```

POST defines the post processing requested

BINSIZE defines the size of the created postprocessing energy spectra.

SMOOTH= 2 is required to calculate the correct statistical error in the post precessing energy spectra. This is by default 2, but can be set a bit larger if necessary.

The script file "imint.sh" also contains information about the number of cores which are used for the simulation.

In the first few lines of "imint.sh" one can and should activate the "testmode". Then the program runs through its initialization process but does not start a simulation. If an error in the namelist occurs the program shows the line in the ".inp" script file where an error occurred. In this way it is easy to check and correct the consistency of the ".inp" script files.

When using the INTEL Oneapi compiler, one must specify computer ID in a text file called "hostfile". The command which is executed by "imint.sh" is

```
mpirun -n "$NP" -ppn "$PPN" -f ../../bin/linux.PRO/hostfile ../../bin/linux.PRO/imintdyn.exe "$TM"
```

with number of processors NP, number of Threads PPN, location of the file "hostfile" and flag TM if testmode is activated

"hostfile" is here located in the imintdyn82/bin/linux.PRO directory where the also "imintdyn.exe and the compilation script files are located.

When using the older INTEL compiler, the following command is executed:

```
mpirun -n "$NP" ../../bin/linux.PRO/imintdyn.exe "$TM"
```

with number of processors NP and flag TM if testmode is activated.