### Python Rapid Artificial Intelligence Ab Initio Molecular Dynamics

**User Manual** 



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- 2. Jingbai Li, Rachel Stein, Daniel Adrion, Steven A. Lopez, "Machine-learning photodynamics simulations uncover the role of substituent effects on the photochemical formation of cubanes", *J. Am. Chem. Soc.* **2021**, 143, 48, 20166–20175. DOI:10.1021/jacs.1c07725
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# 1. What is PyRAI<sup>2</sup>MD

Python Rapid Artificial Intelligence Ab Initio Molecular Dynamics (PyRAI<sup>2</sup>MD) is a suite of Python scripts for nonadiabatic molecular dynamics simulation using machine-learning (ML) potentials. The primary aim of this project is to leverage the present nonadiabatic molecular dynamics (NAMD) techniques enabling nanosecond-scale simulations for medium-size molecular systems at high-level quantum chemical methods e.g., complete active space self-consistent field (CASSCF) with extended multistate second-order perturbative corrections (XMS-CASPT2).

PyRAI<sup>2</sup>MD is designed as a user-friendly platform that integrate the trajectory surface hopping algorithms, and the state-of-the-art Neural Networks (NNs) models. PyRAI<sup>2</sup>MD aims to simplify the job preparation procedures for newcomers of ML and NAMD.

PyRAI<sup>2</sup>MD integrates a NAMD kernel and an ML kernel via an internal communication in memory. In turn, new features in NAMD simulations and ML models can be developed simultaneously.

# 2. Features

## 2.1. Nonadiabatic molecular dynamics

NVE, NVT, center of mass velocity removal, excessive kinetic energy FSSH, ZNSH, NOSH

## 2.2. Machine-learning models

NNs

Model selection

## 2.3. External quantum chemical programs

Molcas

Local, slurm, customized basis set

**BAGEL** 

Local, slurm

**ORCA** 

Local, slurm

GFN-xTB

Local, slurm

**MNDO** 

In the future

## 3. Installation

PyRAI<sup>2</sup>MD is tested on Python 3.7–3.9.

First, download the codes.

git clone https://github.com/mlcclab/PyRAI2MD-hiam.git

Go to the PyRAI<sup>2</sup>MD folder and install. After installation, it creates a command pyrai2md to run calculations.

cd ./PyRAI2MD-hiam pip install .

Compile fssh library using pyrai2md command.

pyrai2md update

To run PyRAI<sup>2</sup>MD, simply use the command following by the input file.

pyrai2md input

PyRAI<sup>2</sup>MD contains some test calculations to verify the code and dependencies. Go to the test folder.

cd ./test

Edit test\_case.py and choose the test job by setting test\_\$job = 1. Modify the environment variables in the run script file, run\_test.sh. The run the script.

bash run test.sh

# 4. Getting started with PyRAI<sup>2</sup>MD

### 4.1. Input structure

PyRAI<sup>2</sup>MD reads a plain text file and does not require a specific extension. An input file looks like below:



The content is case insensitive, but each keyword (*blue*) must take one to read the input value (*red*) properly. The '&' defines a keyword section (*black*) and the empty line will be automatically skipped. Current available keyword sections include:

CONTROL	This section	reads general	information	to set	up	calculations.	lt	also
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controls the parameters used in adaptive sampling for the neural network

active learning.

MOLECULE This section reads molecular specifications including configuration

interaction space, spin multiplicities. It also defines the interstate couplings,

multiscale regions, periodic conditions, and external constrains.

**MOLCAS** This section reads environment variables for setting up Molcas calculations.

**BAGEL** This section reads environment variables for setting up BAGEL calculations.

**ORCA** This section reads environment variables for setting up ORCA calculations.

XTB This section reads environment variables for setting up GFN2-xTB

calculations.

MD This section reads (nonadiabatic) molecular dynamics parameters. It

controls the cutoff of the trajectories for the neural network active learning.

**NN** This section reads the model information of neural networks. It trains

PyRAI<sup>2</sup>MD native MLP models.

MLP This section reads the model information of neural networks. It trains MLP

models using pyNNsMD library.

**SCHNET** This section reads the model information of neural networks. It trains

SchNet models using pyNNsMD library.

**E2N2** This section reads the model information of neural networks. It trains E2N2

models using GCNNP library (E2N2 is currently under development and

not available yet).

**SEARCH** This section reads the parameters used in grid search for optimizing neural

network hyperparameters. Currently, it only support PyRAI<sup>2</sup>MD native MLP

models.

**EG** This section reads the hyperparameters for energy+gradient model. It is

required when NN or MLP is set.

**NAC** This section reads the hyperparameters for nonadiabatic coupling model.

It is required when NN or MLP is set.

**SOC** This section reads the hyperparameters for spin-orbit coupling model. It is

required when NN or MLP is set.

**EG2** This section reads the hyperparameters for the second energy+gradient

model. It is required when NN or MLP is set.

NAC2 This section reads the hyperparameters for the second nonadiabatic

coupling model. It is required when NN or MLP is set.

**SOC2** This section reads the hyperparameters for the second spin-orbit coupling

model. It is required when NN or MLP is set.

**SCH EG** This section reads the hyperparameters for energy+gradient model. It is

required when **SCHNET** is set. SchNet models do not have many parameters to tune, thus the second set of hyperparameters are not used.

SCH NAC The current SchNet model does not support NAC prediction

**SCH\_SOC** This section reads the hyperparameters for spin-orbit coupling model. It is

required when **SCHNET** is set. SchNet models do not have many parameters to tune, thus the second set of hyperparameters are not used.

**E2N2\_EG** This section reads the hyperparameters for energy+gradient model. It is

required when **E2N2** is set. E2N2 models do not have many parameters to

tune, thus the second set of hyperparameters are not used.

**E2N2\_NAC** This section reads the hyperparameters for nonadiabatic coupling model.

It is required when **E2N2** is set. E2N2 models do not have many parameters to tune, thus the second set of hyperparameters are not used.

**E2N2\_SOC** This section reads the hyperparameters for spin-orbit coupling model. It is

required when **E2N2** is set. E2N2 models do not have many parameters to

tune, thus the second set of hyperparameters are not used.

**FILE** This section reads molecular information to use PyRAI<sup>2</sup>MD tool for

training data extraction.

# 5. Keyword sections

# 5.1. CONTROL

\$CONTROL					
title	None	name for the output, user defined			
ml_ncpu	1	number of cpu used for ml jobs			
qc_ncpu	1	number of cpu used for qc jobs			
gl_seed	1	random number seed			
jobtype	sp	type of PyRAI <sup>2</sup> MD job			
qm	nn	neural networks as the electronic property calculator			
key	words below	w are used for adaptive sampling			
abinit	molcas	molcas as the ab initio calculator			
load	1	load existing model for adaptive sampling			
pop_step	200	save average population for the first 200 steps			
refine	0	refine data collected near the state-crossing region, the default value skips this procedure			
refine_num	4	number of data collected near the state-crossing region for refinement			
refine_end	200	the range to search structures near the state- crossing region, the default searches the first 200 steps in the recorded structures in MD			
refine_gap	0.3	the energy gap to select the crossing region structures			
maxiter	1	maximum number of iterations in the adaptive sampling			
maxsample	1	Maximum number of sampled structures per trajectory			
dynsample	0	use dynamically weighted thresholds, the default value uses constant thresholds to sample structures			
maxdiscard	0	maximum discarded snapshots before adjusting thresholds			
maxenergy	0.05	maximum energy threshold to stop trajectories, the unit is Hartree			

minenergy	0.02	minimum energy threshold to record snapshots of a trajectory
dynenergy	0.1	weights to increase or decrease the current energy threshold according to the distance between the minimum and maximum energy threshold
inienergy	0.3	initial value of the maximum energy threshold
fwdenergy	1	number of iterations delayed before increasing the current energy threshold
bckenergy	1	number of iterations delayed before decreasing the current energy threshold
maxgrad	0.15	maximum gradient threshold to stop trajectories, the unit is Hartree·Bohr <sup>-1</sup>
mingrad	0.06	minimum gradient threshold to record snapshots of a trajectory
dyngrad	0.1	weights to increase or decrease the current gradient threshold according to the distance between the minimum and maximum gradient threshold
inigrad	0.3	initial value of the maximum gradient threshold
fwdgrad	1	number of iterations delayed before increasing the current gradient threshold
bckgrad	1	number of iterations delayed before decreasing the current gradient threshold
maxnac	0.15	maximum nac threshold to stop trajectories, the unit is Bohr <sup>-1</sup>
minnac	0.06	minimum nac threshold to record snapshots of a trajectory
dynnac	0.1	weights to increase or decrease the current nac threshold according to the distance between the minimum and maximum nac threshold
ininac	0.3	initial value of the maximum nac threshold
fwdnac	1	number of iterations delayed before increasing the current nac threshold
bcknac	1	number of iterations delayed before decreasing the current nac threshold
maxsoc	50	maximum soc threshold to stop trajectories, the unit is cm <sup>-1</sup>
minsoc	20	minimum soc threshold to record snapshots of a trajectory
dynsoc	0.1	weights to increase or decrease the current soc threshold according to the distance between the minimum and maximum soc threshold

inisoc	0.3	initial value of the maximum soc threshold
fwdsoc	1	number of iterations delayed before increasing the soc energy threshold
bcksoc	1	number of iterations delayed before decreasing the soc energy threshold

title sets the name of the calculation, all temporary and logfiles will be named

according to this value.

ml\_ncpu sets the number of cpu that will be used to run ML-related jobs using python

multiprocessing. ML-related jobtype are train, adaptive, search.

qc\_ncpu sets the number of cpu that will be used to run QC-related jobs using python

multiprocessing. QC-related jobtype is adaptive.

ms\_ncpu sets the number of cpu that will be used to run multiscale calculations using

python multiprocessing. Currently disabled.

gl\_seed sets the global seed for random number generator. It affects the

reproducibility of the surface hopping calculations during NAMD and

adaptive sampling.

jobtype sets the type of PyRAI<sup>2</sup>MD job. Available options are:

sp single-point calculations,

md NAMD simulation,

hop surface hopping calculation.

adaptive adaptive sampling,train training NNs,

predictionpredicting electronic properties using trained NNs,searchNN hyperparameter optimization with grid search.

chooses the electronic property calculator. Available options are:

nn uses PyRAI2MD native MLP model,

mlp uses pyNNsMD MLP model, schnet uses pyNNsMD SchNet model, e2n2 uses GCNNP E2N2 model,

molcas uses OpenMolcas for CASSCF calculations,

mlctkr uses OpenMolcas/Tinker for QM/MM calculations,

bagel uses BAGEL, for CASSCF and XMS-CASPT2 calculations orca uses ORCA for DTF (only ground-state), TD-DFT, or Spin-

flip TDDFT calculations

xtb uses GFN2-xTB for ground-state calculations

specifying a method followed with xtb will enable ONIOM-type QM/QM2 calculation. e.g, qm molcas xtb. The QM region is defined in &MOLECULE section. Three-layer ONIOM calculation can be invoked by qm molcas xtb xtb. Besides the QM and QM2 calculations, It uses GFN-FF for the third layer (MM) calculation without further keywords.

abinit

chooses the reference QC electronic property calculator. Available options are the same as **qm** except for **nn**. The chosen program will be used to recompute the QC-data for the collected structures during adaptive sampling.

load

reads a pretrained NNs for adaptive sampling. When it is set to 0, it will first training NNs before running the adaptive sampling.

pop\_step

sets the number of MD steps to compute the average population over all trajectories propagated during adaptive sampling. Note that the step size depends on both the timestep and checkpointing frequency, which can be specified by size in &MD section.

refine

turns on additional structural sampling around the state-crossing region during adaptive sampling. Available options are:

- skip additional sampling, which is the default;
- always do additional sampling, this will search structure even trajectories complete with no other uncertain structures;
- do additional sampling until trajectories complete without uncertain structures;
- do additional sampling only if trajectories do not complete and do not find uncertain structures.

refine\_num

sets the number of structures that will be collected near the state-crossing regions during adaptive sampling. The selection start from the structure with the smallest energy gap.

refine end

sets the range to search the structures near the state-crossing region in the recorded trajectories. Note that the adaptive sampling only records the last a few MD steps to reduce the memory usage. The number of recorded MD steps can be adjusted by **record\_step** in &**MD** section.

refine\_gap

sets the energy gap threshold to select the structures near the state-crossing region. The default value is 0.3 eV.

maxiter

sets the maximum number of iterations for adaptive sampling. The adaptive sampling will stop when it reach the maximum value or no longer find new structures.

maxsample

set the number of structures to be collected during the adaptive sampling. Note that this number does not include the number of structure refinement from refine\_num.

dynsample

turns on the dynamically weighted adaptive sampling. The threshold values will be dynamically adjusted according to the numerical distance between the minimum and the maximum value. It is turn off in default.

maxdiscard

set the maximum number of discard structures in a trajectory. A structure will be discarded if it contains a non-physical bond length shorter than the sum of the van der Waals radius of each atom multiplied by 0.7. When the number of discarded structures exceed **maxdiscard**, the current threshold will be decreased to limit the exploration region of adaptive sampling. Otherwise, the current threshold will be increased to expand the exploration region of adaptive sampling. Note that the threshold adjustment can be delayed by fwd\* and bck\* keywords for the forward and backward direction.

maxenergy

sets the maximum value of the energy threshold to stop a trajectory.

minenergy

sets the minimum value of the energy threshold to record a trajectory.

dynenergy

sets the weights of the to increase or decrease the current energy threshold according to the distance between the minimum and maximum energy threshold. The adjustment is weights \* (max - min) but the adjusted values will not exceed the minimum or maximum values.

inienergy

set the initial value of the energy threshold to be dynamically adjusted.

**fwdenergy** 

set the number of delayed iterations to increase the current threshold.

bckenergy

set the number of delayed iterations to decrease the current threshold.

maxgrad

sets the maximum value of the gradient threshold to stop a trajectory.

mingrad

sets the minimum value of the gradient threshold to record a trajectory.

dyngrad

sets the weights of the to increase or decrease the current gradient threshold according to the distance between the minimum and maximum gradient threshold. The adjustment is weights \* (max - min) but the adjusted values will not exceed the minimum or maximum values.

inigrad set the initial value of the gradient threshold to be dynamically adjusted. fwdgrad set the number of delayed iterations to increase the current threshold. bckgrad set the number of delayed iterations to decrease the current threshold. maxnac sets the maximum value of the nac threshold to stop a trajectory. minnac sets the minimum value of the nac threshold to record a trajectory. dynnac sets the weights of the to increase or decrease the current nac threshold according to the distance between the minimum and maximum nac threshold. The adjustment is weights \* (max - min) but the adjusted values will not exceed the minimum or maximum values. ininac set the initial value of the nac threshold to be dynamically adjusted. fwdnac set the number of delayed iterations to increase the current threshold. bcknac set the number of delayed iterations to decrease the current threshold. maxsoc sets the maximum value of the soc threshold to stop a trajectory. minsoc sets the minimum value of the soc threshold to record a trajectory. dynsoc sets the weights of the to increase or decrease the current soc threshold according to the distance between the minimum and maximum soc threshold. The adjustment is weights \* (max - min) but the adjusted values will not exceed the minimum or maximum values. inisoc set the initial value of the nac threshold to be dynamically adjusted. fwdsoc set the number of delayed iterations to increase the current threshold. bcksoc set the number of delayed iterations to decrease the current threshold.

#### 5.2. MOLECULE

&MOLECULE		
ci	1	definition of the configuration interaction space for

		each spin state
spin	0	definition of the spin multiplicity for each spin state
coupling	None	definition of the interstate couplings
highlevel	None	definition of the high level atoms
midlevel	None	defination of the middel level atoms
embedding	False	embed surrounding charge in high level region
read_charge	False	read charge from a .charge file
freeze	None	definition of frozen atoms
constrain	None	definition of constrained atoms
cbond	None	definition of restrained bonds
cangle	None	definition of restrained angles
cdihedral	None	definition of restrained dihedrals
tbond	None	target value of restrained bonds
tangle	None	target value of restrained angles
tdihedral	None	target value of restrained dihedrals
fbond	10.0	prefactor of the biasing potential on bond
fangle	0.005	prefactor of the biasing potential on angle
fdihedral	1e-6	prefactor of the biasing potential on dihedral
shape	ellipsoid	definition of constraining potential
factor	40	exponential factor of the constraining potential
scale	1.0	prefactor of the exponential potential
cavity	None	constraining radius along x, y, and z-axis
center	None	center of the constraining potential
center_type	xyz	set type of center for constraining potential
groups	None	define molecule groups to apply constrain
compress	None	compress shape of potential
track_type	None	track geometric changes in given type of parameter
track_index	None	atom indices to compute geometrical parameters
track_thrhd	None	threshold of geometrical changes to stop MD

ci

sets configuration interaction space for each spin state, i.e., the number of states in each spin multiplicity, 2 means two states of the first spin, i.e., S0, S1. It can take multiple integers if multiple spin states are iinvolved, e.g. 2 means two states in spin 1 and two states in spin 2. the spin multiplicities are defined by spin.

spin

sets the total spin number for each spin state, 0 is singlet, 1 is triplet. It follows the same order as ci.

coupling

reads pairwise indices to define the coupling between two states. Each pair should be separated by ','. The following example,

ci 2 2 spin 0 1

**coupling** 1 2, 2 3, 2 4, 3 4

defines that state 1 and 2 are singlet and state 3 and 4 are triplet. It includes the nac between state 1 and 2 (singlet) and state 3 and 4 (triplet) as well as the soc between state 2 and 3 (singlet-triplet) and state 2 and 4 (singlet-triplet). The order of index pairs does not matter and the coupling of the non-defined pairs (e.g, state 1 and 4) will be treated as zero.

highlevel

reads the atom indices in QM region. The indices can be written individually, or in a range, e.g., 1 2 3 5 6, 1-3 5-6 or 1-2 3 5-6.

midlevel

reads the atom indices in QM2 region. The indices can be written individually, or in a range, e.g., 1 2 3 5 6, 1-3 5-6 or 1-2 3 5-6. The rest of atoms will be placed in the outer region (MM).

embedding

embed middle-level surrounding charge in the high-level region if set true.

read charge

read middle level surrounding charge from a .charge file if set true. This option will keep the same charge throughout the dynamics, suitable for rigid crystal environment. Turn it false to dynamically update the embeding charge during dynamics for flexible environment like solvent. Note the ML models in PyRAI<sup>2</sup>MD currently do not have charge embedding function. You must set it true to use the same embedding charge throughout the NN training. Otherwise, the energies and forces are not learnable.

freeze

reads the indices to freeze atoms during dynamics

constrain

reads the indices to apply constraints on atoms during dynamics. All atoms will be included If no indices are provided.

cbond

reads the indices to apply biasing potential on bonds during dynamics. No bond will be restrained If the keyword is omitted. 1 2, 3 4 represent two bonds, one between atom 1 and 2 and the other between atom 3 and 4.

cangle

reads the indices to apply biasing potential on angles during dynamics. No angle will be restrained If the keyword is omitted. 1 2 3, 4 5 6 represent two angles, one between atom 1, 2, and 3, centered on atom 2 and the other between atom 4, 5, and 6, centered on atom 5.

cdihedral

reads the indices to apply biasing potential on dihedrals during dynamics. No dihedral will be restrained If the keyword is omitted. 1 2 3 4, 5 6 7 8 represent two dihedrals, one over atom 1, 2, 3, and 4, where the axis is defined by atom 2 and 3; the other over atom 5, 6, 7 and 8, where the axis is defined by atom 6 and 7.

tbond

define the target value for the restrained bonds. Multiple values are acceptable for multiple restrained bonds. For example, 1.5 1.6 will restrain the two bonds to 1.5 and 1.6 Angstrom. If only one value is given, it will be applied to all restrained bonds. If this keyword is omitted, the original values of the restrained bonds will be used as target values.

tangle

define the target value for the restrained angles. Multiple values are acceptable for multiple restrained angles. For example, 100 90 will restrain the two angles to 100 and 90 Degree. If only one value is given, it will be applied to all restrained angles. If this keyword is omitted, the original values of the restrained angles will be used as target values.

tdihedral

define the target value for the restrained dihedrals. Multiple values are acceptable for multiple restrained dihedrals. For example, 100 90 will restrain the two dihedrals to 100 and 90 Degree. If only one value is given, it will be applied to all restrained dihedrals. If this keyword is omitted, the original values of the restrained dihedrals will be used as target values.

fbond

define the prefactor for the biasing potential on bonds. Default is 10.0 Hartree.

fangle

define the prefactor for the biasing potential on angles. Default is 0.005 Hartree.

fdihedral

define the prefactor for the biasing potential on dihedrals. Default is 1e-6 Hartree.

shape define the shape of the constraining potential. Available options are

ellipsoid and cuboid.

factor define the exponential factor of the constraining potential. The larger the

value is, the shaper the potential wall is. Default is 40.

scale define the prefactor for the exponential potential. Default is 1.0 Hartree.

cavity reads constraining radius along x, y, and z-axis. If no value is provided, the

constraining potential will be turned off.

center reads the atom indices to define the center of the constraining potential.

**center\_type** choose the type of center to apply the constraining potential.

mass use the center of mass xyz use geometrical center origin set the center to (0, 0, 0)

**groups** define the molecule groups to apply the constraining potential.

In default, each atom is considered in the individual group, so the constraining potential will be applied to each of them according to their distance to the center.

Alternatively, we can define a group of atoms that their constraining potential will be determined by the distance between the center of mass and potential center. For example,

8 3, 4 5 represent the first group has 8 molecules with 3 atoms, the second group has 4 molecules with 5 atoms. Note the total number of atoms (8\*3+4\*5=44) must equal to the number of constrain atoms.

reads the target ratio and step to compress the shape of the constraining potential. For example,

 $0.75\ 1000$  will compress the constraining potential from the original shape defined by cavity to 0.75 of them along x, y, and z-axis in 1000 steps. This option is useful to tune the density of the system.

track\_type set the type of geometrical parameter used to early stop the trajectories.

Available options are:

frag track the distance between two fragments.

dist track the distance between two atoms.

track index reads the atom indices to define the fragments of interatomic distances.

To define fragments, the format follows as 1 2 3 4, 5 6 7 8, where the first and second four indices, separated by a ',' punctuation, correspond to the atoms defining the first and second fragment. For tracking interatomic distances, the format follows as 1 2, 3 4, 5 6, 7 8, where the four pairs of indices are separated by ','. Each of them corresponds to a distance between two atoms.

track\_thrhd

reads the threshold to early stop the trajectories if the distances exceed the thresholds. For tracking fragments, only one value is needed. For tracking interatomic distances, more values are supported. If only one value is given, it will be used for all distances. If multiple values are provided, each of them will be used to check the distance defined by track\_index accordingly. In this case, the number of values should match the number of tracked distances. The unit is Angstrom.

track\_stop

decide what task to be stopped when the tracking geometric parameters meet the threshold.

Available options are:

- 0 has no effects, which is the default.
- 1 stop the trajectory immediately
- 2 stop surface hopping and continue the trajectory at the current state until the tracking parameters become smaller than the threshold.

#### 5.3. MOLCAS

The Molcas calculation also needs an input template and guess orbital named with .StrOrb in the current folder. See X for examples of running Molcas calculations.

&MOLCAS		
molcas	None	path to Molcas executable
molcas_nproc	1	number of cpu for OpenMP parallelization
molcas_mem	2000	number of memories for calculation
molcas_print	2	logfile printing level
molcas_project	None	project name
molcas_calcdir	\$PWD	path to the temporary calculation folder
molcas_workdir	None	path to Molcas scratch folder

basis	2	additional basis set information
omp_num_threads	1	number of threads for OpenMP parallelization
use_hpc	0	submit calculation to remote cluster
keep_tmp	1	keep the temporary calculation folder

molcas sets the path to Molcas executable.

molcas nproc sets \$MOLCAS NPROC environment variable, the default value is 1.

molcas\_mem sets \$MOLCAS MEM environment variable, the default value is 2000 MB.

molcas\_print sets \$MOLCAS PRINT environment variable, the default value is 2.

molcas\_project sets \$MOLCAS\_PROJECT environment variable, the default value is

taken from title in &CONTROL section

molcas\_calcdir sets the path to a temporary folder for Molcas calculation. The temporary folder will be named as tmp\_MOLCAS. If no path is provided, the tmp\_MOLCAS will be created in the current folder. Note this is the folder to run Molcas calculations, but not necessary to be the Molcas scratch folder,

which is set by molcas\_workdir.

molcas workdir sets \$MOLCAS WORKDIR environment variable. If no path is provided, it

will be the same path as the tmp\_MOLCAS folder set by molcas\_calc. Note that Molcas is input/output intensive, the temporary files could be large and the calculation running in SLURM's /scratch could be slower than in a local disk. It is recommended to use a local folder such as /tmp or /srv/tmp. If you are not sure which folder to use, a shortcut is AUTO, which

needs to be upper-case.

basis reads atom annotation to use different basis sets if it is set to 1. It is turned off in default (2). To use different basis sets, you need to prepare a xyz file.

off in default (2). To use different basis sets, you need to prepare a xyz file following the same atom order and annotate the atom with '\_', e.g. "C\_ X Y Z". The coordinates can be random. Then add the basis set in &GATEWAY in the Molcas input template, e.g. "ANO-S-MB, C\_.ANO-S-VDZP", which

will use ANO-S-VDZP for annotated atoms but ANO-S-MB for others.

omp\_num\_threads sets OpenMP parallel threads for OpenMolcas, the default value is 1. Note

that not all Molcas functions are parallelized.

use hpc

submits the Molcas calculation to the job scheduler. It is turned off in default, thus the calculation is running as a subprocess in the current machine. For single calculation, it is recommended to run the Molcas calculation without <code>use\_hpc</code> because it does not have to wait in the queue. However, if there are more Molcas calculations than available cpus or the disk space for all calculations is not enough, e.g. in adaptive sampling, it is better to use <code>use\_hpc</code> to distribute the calculations to all available nodes via a job scheduler. To use this function, you need to prepare a submission script template with the same name as <code>title</code> in &CONTROL section, e.g. job title.slurm and specify the all necessary #SBATCH variables.

keep tmp

keep the temporary Molcas calculation folder. It is turned on in default. Set to 0 to turned off.

#### 5.4. BAGFI

The BAGEL calculation also needs an input template and orbital archive in the present folder. See X for examples of running BAGEL calculations.

&BAGEL		
bagel	None	path to BAGEL executable
bagel_nproc	1	number of cpu for BAGEL parallelization
bagel_project	Npne	project name
bagel_workdir	\$PWD	path to BAGEL calculation folder
bagel_archive	None	name of BAGEL orbital archive
mpi	None	path to the MPI library
blas	None	path to BLAS library
lapack	None	path to LAPACK library
boost	None	path to BOOST library
mkl	None	path to MKL library
arch	None	cpu architecture
omp_num_threads	None	number of threads for OpenMP parallelization
use_mpi	0	use MPI for parallelization

use_hpc	0	submit calculation to remote cluster
keep_tmp	1	keep the temporary calculation folder

bagel sets the path to BAGEL executable.

bagel\_nproc sets the number of cpu for BAGEL calculation with OpenMP parallelization

bagel\_project sets the name of BAGEL calculation, the default value is taken from title in

&CONTROL section

bagel\_workdir sets the path to a temporary folder. It creates a sub folder tmp BAGEL for

BAGEL calculation. BAGEL is mainly running in memory. Therefore, it does

not suffer from the input/output overhead issue.

bagel archive sets the name of BAGEL orbital archive if the orbital archive has a different

name from title in &CONTROL section. In default, the name is taken from

title in &CONTROL section.

mpi sets the path to MPI. For the latest (2022) Intel's OneAPI, the environment

variables of mkl and mpi can be initialized together by sourcing the setvar.sh in the OneAPI's folder. PyRAI2MD will use mkl to find the source

file.and this keyword can be left to empty.

blas sets the path to BLAS library.

lapack sets the path to LAPACK library.

**boost** sets the path to BOOST library.

mkl sets the path to Intel MKL library. For the latest (2022) Intel's OneAPI, the

environment variables of mkl and mpi can be initialized together by sourcing the setvar.sh in the OneAPI's folder. Thus, this keyword needs to

be set to the OneAPI's folder that contains the setvar.sh.

arch specifies the cpu architecture, the previous default value is intel64. For the

latest (2022) Intel's OneAPI, the environment variables of mkl and mpi can be initialized together by sourcing the setvar.sh in the OneAPI's folder.

Thus, this keyword needs to be left emtyp.

omp num threads sets OpenMP parallel threads for BAGEL, the default value is 1.

use hpc

submits the BAGEL calculation to the job scheduler. It is turned off in default, thus the calculation is running as a subprocess in the current machine. For single calculation, it is recommended to run the BAGEL calculation without use hpc because it does not have to wait in the gueue. However, if there are more BAGEL calculations than available cpus or the disk space for all calculations is not enough, e.g. in adaptive sampling, it is better to use use hpc to distribute the calculations to all available nodes via a job scheduler. To use this function, you need to prepare a submission script template with the same name as title in &CONTROL section, e.g. job title.slurm and specify the all necessary #SBATCH variables.

keep tmp

orca

keep the temporary BAGEL calculation folder. It is turned on in default. Set to 0 to turned off.

#### 5.5. ORCA

The ORCA calculation only needs an input template the present folder. See X for examples of running ORCA calculations.

The keywords, default values, and short descriptions are listed below.

&BAGEL		
orca	None	path to ORCA executable
orca_project	None	project name
orca_workdir	\$PWD	path to ORCA calculation folder
dft_type	tddft	type of DFT calculation
mpi	\$PWD	path to the OpenMPI library
use_hpc	0	submit calculation to remote cluster
keep_tmp	1	keep the temporary calculation folder

Full descriptions for all available keywords are summarized below.

sets the path to ORCA executable. It only supports ORCA 5.0 orca project sets the name of ORCA calculation, the default value is taken from title in &CONTROL section orca workdir sets the path to a temporary folder. It creates a sub folder tmp ORCA for ORCA calculation.

**dft\_type** sets the type of DFT calculation.

dft ground-state DFT calculation.

tddft TDDFT calculation.

sf\_tddft Spin-flip TDDFT calculation. It only supports 1-particle-1-

hole operator, it could be hard to converge more than 3

singlet states. Must be used with cautions.

mpi sets the path to OpenMPI

use\_hpc submits the ORCA calculation to the job scheduler. It is turned off in default,

thus the calculation is running as a subprocess in the current machine. For single calculation, it is recommended to run the ORCA calculation without <code>use\_hpc</code> because it does not have to wait in the queue. However, if there are more ORCA calculations than available cpus or the disk space for all calculations is not enough, e.g. in adaptive sampling, it is better to use <code>use\_hpc</code> to distribute the calculations to all available nodes via a job scheduler. To use this function, you need to prepare a submission script template with the same name as <code>title</code> in <code>&CONTROL</code> section, e.g.

job title.slurm and specify the all necessary #SBATCH variables.

keep\_tmp keep the temporary ORCA calculation folder. It is turned on in default. Set

to 0 to turned off.

### 5.6. OpenQP

The OpenQP calculation only needs an input template the present folder. See X for examples of running OpenQP calculations.

&BAGEL		
openqp	None	path to OpenQP root folder
openqp_project	None	project name
openqp_workdir	\$PWD	path to OpenQP calculation folder
threads	1	Number of threads for OpenMP
use_hpc	0	submit calculation to remote cluster
keep_tmp	1	keep the temporary calculation folder

opengp sets the path to OpenQP root folder.

opengp project sets the name of OpenQP calculation, the default value is taken from title

in &CONTROL section

openqp\_workdir sets the path to a temporary folder. It creates a sub folder tmp\_OpenQP

for OpenQP calculation.

threads sets number of threads for OpenMP parallelization.

use hpc submits the OpenQP calculation to the job scheduler. It is turned off in

default, thus the calculation is running as a subprocess in the current machine. For single calculation, it is recommended to run the OpenQP calculation without <a href="use-hpc">use-hpc</a> because OpenQP is internally linked with PyRAI<sup>2</sup>MD for efficient data communication. However, if there are more OpenQP calculations than available cpus or the disk space for all calculations is not enough, e.g. in adaptive sampling, it is better to use <a href="use-hpc">use-hpc</a> to distribute the calculations to all available nodes via a job scheduler. To use this function, you need to prepare a submission script template with the same name as <a href="title">title</a> in &CONTROL section, e.g.

job title.slurm and specify the all necessary #SBATCH variables.

keep tmp keep the temporary OpenQP calculation folder. It is turned on in default.

Set to 0 to turned off.

### 5.7. XTB

The GFN2-xTB calculation does not needs any input template in the present folder. See X for examples of running GFN2-xTB calculations.

&XTB		
xtb	None	path to xTB executable
xtb_project	None	project name
xtb_workdir	\$PWD	path to xTB calculation folder
xtb_nproc	1	Number of OMP threads
gnfver	-2	version of GFN-xTB

mem	1000	Memory for OMP stack size
use_hpc	0	submit calculation to remote cluster
keep_tmp	1	keep the temporary calculation folder

**xtb** sets the path to GFN-xTB executable.

xtb\_project sets the name of GFN-xTB calculation, the default value is taken from title

in &CONTROL section

xtb\_workdir sets the path to a temporary folder. It creates a sub folder tmp XTB for

GFN-xTBcalculation.

xtb\_nproc sets the number of threads for parallel GFN-xTB calculation

**gnfver** sets the version of GFN-xTB calculation. Available options are:

-2 default GFN version of the installed GFN-xTB

-1 use GFN FF

0 use GFN0

1 use GFN1

2 use GFN2

xmem sets the memory for OMP STACKSIZE in MB.

use\_hpc submits the GFN2-xTB calculation to the job scheduler. It is turned off in

default, thus the calculation is running as a subprocess in the current machine. For single calculation, it is recommended to run the GFN2-xTB calculation without **use\_hpc** because it does not have to wait in the queue. However, if there are more ORCA calculations than available cpus or the disk space for all calculations is not enough, e.g. in adaptive sampling, it is better to use **use\_hpc** to distribute the calculations to all available nodes via a job scheduler. To use this function, you need to prepare a submission script template with the same name as **title** in &CONTROL section, e.g.

job title.slurm and specify the all necessary #SBATCH variables.

keep the temporary ORCA calculation folder. It is turned on in default. Set

to 0 to turned off.

5.8. MD

&MD		
initcond	0	sample initial condition
excess	0	excess kinetic energy in Hartree
scale	1	scale kinetic energy by a factor
target	0	set a target kinetic energy in Hartree
graddesc	0	gradient descent mode (zero velocity)
reset	0	remove center of mass velocity
resetstep	0	center of mass velocity reset interval
ninitcond	20	number of sampled initial conditions
method	wigner	initial condition sampling method
format	molden	frequency file format
randvelo	0	Initialize random velocity
temp	300	temperature in Kelvin
step	10	number of threads for OpenMP parallelization
size	20.67	step size in the atomic unit of time
root	1	initial state
activestate	0	only compute gradients of the current state
sfhp	nosh	surface hopping algorithm
nactype	ktdc	type of nac
phasecheck	0	apply phase correction to nac
gap	0.5	energy gap threshold to compute Zhu-Nakamura surface hopping between the same spin states
gapsoc	0.5	energy gap threshold to compute Zhu-Nakamura surface hopping between the different spin states
substep	20	number of substep in wave function integration in FSSH calculation
integrate	0	accumulate the nuclear amplitude transfer in FSSH calculation *This is only for debug purpose*
deco	0.1	energy-based decoherence correction in Hartree

	_	
adjust	1	adjust velocity at surface hopping
reflect	1	reflect velocity at frustrated hopping
maxh	10	Maximum number of allowed surface hoppings
dosoc	0	compute Zhu-Nakamura surface hopping between the different spin states
thermo	off	apply a thermostat for NVT ensemble
thermodelay	200	delay time for applying a thermostat in the ground- state
silent	1	no output prints on screen
verbose	0	logfile printing level
direct	2000	number of MD steps that will be written in output
buffer	500	number of MD steps that will be skipped in output
record	0	part of the data that will be recorded for adaptive sampling
record_step	0	number of the last MD snapshots that will be recorded for adaptive sampling
checkpoint	0	checkpoint a trajectory for a given number of MD steps
restart	0	restart calculation
addstep	0	add MD steps in a restart calculation

#### initcond

generates initial conditions from a frequency file. It is turned off in default. Thus, it reads coordinates and velocities from .xyz and .velo files. In adaptive sampling, the initial conditions are always generated from a frequency file, no matter it is set to 1 or 0.

#### excess

adds extra kinetic energy beyond the initial kinetic energy then scales the initial velocity isotopically. It is sometimes useful to accelerate the MD and drive the trajectory uphill. The unit is Hartree. This option is the first adjustment to the kinetic energy.

#### scale

scales the initial kinetic energy isotropically by a factor. It is sometimes useful to accelerate the MD and drive the trajectory uphill. This option is the second adjustment to kinetic energy.

target

sets a target kinetic energy to scale the initial velocity isotopically. It is sometimes useful to accelerate the MD and drive the trajectory uphill. This option is the last adjustment to the kinetic energy.

graddesc

propagates a trajectory following the gradient descent by setting the velocities to zero during the MD. It is turned off in default.

reset

removes translation and rotation velocity at the center of mass. It is turned off in default. It helps avoid the "flying ice" artifact, which results from the draining of vibration energy to translation and rotation energy when velocity rescaling (e.g., thermostat) is frequently used.

resetstep

sets the interval of removing translation and rotation velocity at the center of mass. It is usually recommended to reset velocity every 2000 steps with a timestep of 0.5 fs. If it is set to 0, it only reset the initial velocity. This keyword must be used together with reset.

ninitcond

sets the number of initial conditions in sampling. The last condition is used in MD if the value is greater than 1. In adaptive sampling, this value determines the number of trajectories to collect new structures.

method

chooses the method to do initial condition sampling. It is recommended to do Wigner sampling using wigner. The Boltzmann sampling is also available with boltzmann.

**format** 

sets the frequency file format. It supports the Molcas' molden file (\$xxx.freq.molden), BAGEL frequency calculation output file (need to rename as \$xxx.freq.bagel), ORCA frequency calculation output file (need to rename as \$xxx.freq.orca), Gaussian frequency calculation output file and fchk file with "Freq=SaveNormalModes" (need to rename as \$xxx.freq.log and \$xxx.freq.fchk).

temp

sets the temperature in Kelvin for initial condition sampling and thermostat. It is not used in microcanonical ensemble (i.e., NVE).

randvelo

initialize random atomic velocity according to the input temperature.

step

sets the number of MD steps.

size

sets the step size in the atomic unit of time. 1 au = 0.02418884254 fs.

root

sets the initial state in NAMD. It should not be larger than the total number of states defined by **ci** in &MOLECULE.

activestate

only computes the gradients of current state with QC calculations. It is turned off in default. It reduces the cost of FSSH dynamics because the gradients of other states are not used. However, the gradients of all states are needed in Zhu-Nakamura surface hopping. This keyword is not used in ML-NAMD as NNs predict gradients of all states.

sfhp

chooses the surface hopping algorithm. Available options are:

fssh Tully's the fewest switches surface hopping with explicit nac,

gsh Zhu-Nakamura surface hopping,

nosh turn off the surface hopping calculation.

nactype

chooses the type of nac for fssh calculation. Available options are:

nac nonadiabatic coupling vectors, non-weighted by the state energy gap

ktdc curvature driven time-dependent coupling, which approximates nonadiabatic coupling by the first-order derivative of energy in two adjacent MD step.

dcm derivative coupling matrix, computed from the state overlap between two MD steps. Only support OpenQP.

phasecheck

apply phase correction to nonadiabatic coupling by the overlap of nac vectors at two adjacent MD step. It is turned off in default. It is only used when sfhp is set to fssh and nactype is set to nac.

gap

sets the energy gap threshold (in eV) to compute Zhu-Nakamura surface hopping between two states with same spin multiplicity. The Zhu-Nakamura surface hopping calculations are skipped when the energy gap is larger than this value. When **sfhp** is set to **fssh** and **nactype** ktdc, the NACs are considered as 0 if the energy gap is larger than this value.

gapsoc

sets the energy gap threshold (in eV) to compute Zhu-Nakamura surface hopping between two states with different spin multiplicities. The surface hopping calculations are skipped when the energy gap is larger than this value. This keyword is not used when **sfhp** is set to **fssh**.

substep

sets the number of substeps to integrate the electronic wave function in fssh calculation. It is not used when sfhp is set to gsh.

integrate

accumulate the nuclear amplitude in fssh calculation. This is only used for debug purpose and must not be used to produce results for publication.

deco

applies the energy-based decoherence correction in fssh calculation. The unit is in Hartree. It is not used when sfhp is set to gsh.

adjust

scales the velocity at surface hopping events. Available options are:

- 0 do not scale velocity,
- 1 scale velocity isotropically,
- 2 scale velocity along the NAC direction.

reflect

changes the velocity direction when frustrated hopping happens. Available options are:

- directly reflect velocity
- 2 reflect the velocity component along the NAC vectors.

maxh

sets the maximum number of allowed surface hopping events.

dosoc

computes Zhu-Nakamura surface hopping between two states with different spin multiplicities. It requires additional calculations of spin-orbit coupling and is turned off in default.

thermo

controls the ensemble of trajectory. Available options are:

off do not rescale velocity (NVE)

- o rescale velocity to conserve total energy (forced to NVE ensemble)
- 1 rescale velocity using Nóse-Hoover thermostat (NVT ensemble)
- rescale velocity to conserve total energy in the excited state then applying Nóse-Hoover thermostat in the ground-state.

thermodelay

sets the number of MD step delayed for applying a thermostat in the ground-state. It is only used when set thermo is set to 2.

silent

turns off printing output on screen. It is turned on in default.

verbose

controls the printing level.

- only prints energy and state populations,
- 1 prints coordinates, velocities, gradients, and NACs,
- 2 prints more calculations information (screen output only).

direct

sets the number of MD steps to be written in the output file. It starts from the first step.

buffer

sets the number of MD steps to be skipped in output file after direct writing steps.

record

choose the part of data to be recorded in a trajectory during adaptive sampling. Available options are:

whole all data will be recorded;

**qm** only data in qm region will be recorded.

record\_step sets the number of the latest MD steps in a trajectory to be cached in

memory. The cached trajectories are used to sample uncertain data in adaptive sampling. Reduce this number if the molecular dynamics have a huge number of steps or the adaptive sampling does not have enough

memory to proceed.

checkpoint sets the number of MD steps to checkpoint a trajectory. The trajectory is

stored in python pickle file (.pkl) and can be used to restart the calculation.

It is turned off in default.

restart reads the .pkl file to restart a calculation. It is turned off in default.

adds additional MD steps in the restarted calculation. Use this if you want

to continue to propagate a completed trajectory.

### 5.9. NN (MLP, SCHNET, E2N2, DIMENET)

The neural networks in PyRAI<sup>2</sup>MD are implemented with TensorFlow/Keras API and pyTorch. The neural network is built upon fully connected feedforward multilayer perceptron and graph convolutional neural networks. They consist of an input layer, several hidden layers, and an output layer. Each layer is connected by multiple neurons with activation functions. The connection between layers is a linear function including weights and bias.

PyRAI<sup>2</sup>MD offers a convenient interface to train a neural network and load a trained model for the prediction of energies, forces, non-adiabatic couplings, and spin-orbit couplings. PyRAI<sup>2</sup>MD always trains two sets of neural networks, which can have completely different architectures or only different initial weights. This is useful to measure the prediction uncertainty when predicting data out of the training set. The energies and forces are combined in one model and the non-adiabatic couplings and spin-orbit couplings use an independent model. Users can choose to train either one or all of them.

The keywords, default values, and short descriptions are listed below. All types of neural networks share the same keywords in their sections. Here we use &NN section as an example.

&NN (MLP, SCHNET,		
E2N2, and DIMENET)		
modeldir	\$PWD	path to save or load NN
train_data	None	path to load training data
pred_data	None	path to load prediction data
nsplits	10	number of folds to split training data

shuffle	False	shuffle training data every epoch
nn_eg_type	1	number of energy+gradient model
nn_nac_type	0	number of nac model
nn_soc_type	0	number of soc model
eg_unit	si	unit of energy+gradient model
nac_unit	si	unit of nac model
soc_unit	si	unit of soc model
select_eg_out	None	select the output of energy+gradient model
select_nac_out	None	select the output of nac model
select_soc_out	None	select the output of soc model
permute_map	No	path to permutation map for data augmentation
silent	1	no output prints on screen
gpu	0	Use GPU for training, only support E2N2

nsplits

modeldir	sets a path to save or load a NN model. The default location is the present
	folder. The model is saved in a folder named as "NN-\$xxx".

train\_data sets a path to load the training data from a JSON file. See X for the information of data format. If a file name is provided, it assumes that the file is in the current folder.

sets a path to load the prediction data from a JSON file. If a file name is provided, it assumes that the file is in the current folder. It is only used when jobtype is set to prediction.

sets the number of folds to split the training data. The first fold will be used for validation of the first model, and the second fold will be used for validation of the second model. The rest of the data will be used for training model accordingly.

shuffle shuffle the training data every epoch. It helps accelerate the training.

nn\_eg\_type

defines the number of energy+force models with different architectures. Available options are:

- build two neural networks with the same architecture but being initialized with different weights. The hyperparameters are read from &EG.
- build two neural networks with different architecture being initialized with different weights. The hyperparameters are read from &EG and &EG2, respectively.

nn\_nac\_type

defines the number of nac models with different architectures. Available options are:

- 0 skip the nac model.
- build two neural networks with the same architecture but being initialized with different weights. The hyperparameters are read from **&NAC**.
- build two neural networks with different architecture being initialized with different weights. The hyperparameters are read from &NAC and &NAC2, respectively.

nn\_soc\_type

defines the number of nac models with different architectures. Available options are:

- 0 skip the soc model.
- build two neural networks with the same architecture but being initialized with different weights. The hyperparameters are read from **&SOC**.
- build two neural networks with different architecture being initialized with different weights. The hyperparameters are read from &SOC and &SOC2, respectively.

eg unit

set the unit of energy and gradients used in training. Available options are:

- au energy in Hartree and gradient in Hartree Bohr<sup>-1</sup>,
- si energy in eV and gradients in eV· $Å^{-1}$ .

nac unit

set the unit of nac used in training. Available options are:

- au nac in Hartree · Bohr<sup>-1</sup>,
- si nac in  $eV \cdot Å^{-1}$ .

soc\_unit

set the unit of nac used in training. Available options are:

si soc in cm<sup>-1</sup>.

select eg unit

select the output of the energy+gradient model for ML-NAMD simulations. The default option is to use all predicted energy and gradients. The number of states is determined by the training data. If you wish only include the

energy and gradients of a few states instead of all states, you can specify them as the following.

1 2 3 4 will use the data of the first 4 states if more than 4 states are trained. If different spin states are included, for instant, 3 singlet states followed by 3 triplet states, you can use 1 2 4 5 to select the first two singlet states and two triplet states.

select\_nac\_unit

select the output of the nac model for ML-NAMD simulations. The default option is to use all predicted nac. You can set the indices of the nac in the training data to use specific nac values. For instance, if the nac training data are organized for state pair 1 2, 2 3, 3 4, and 4 5, setting **select\_nac\_unit** to 1 2 3 will select the nac of the state pair 1 2, 2 3, and 3 4 (skip the state pair 4 5) for ML-NAMD simulation.

select\_soc\_unit

select the output of the soc model for ML-NAMD simulations. The default option is to use all predicted soc. You can set the indices of the soc in the training data to use specific soc values. For instance, if the soc training data are organized for state pair 1 3, 1 4, 2 3, and 2 4, setting **select\_soc\_unit** to 1 3 will select the soc of the state pair 1 3 and 2 3 (skip the state pair 1 4 and 2 4) for ML-NAMD simulation.

permute\_map

read a text file that defined the permutations of atom indexing. Each line should only include one set of permutation. "1 5 3 2 4 6" means first switch the index of atom 2 and atom 5 then switch the index of atom 4 and the atom 2.

silent

turns off printing output on screen. It is turned on in default.

gpu

use GPU to train E2N2 models. This option does not work for other NN models for the moment.

#### **5.10. SEARCH**

&SEARCH		
keywords below are available for nn		
depth	1	a list to search number of hidden layers
nn_size	20	a list to search number of neurons per hidden layer
batch_size	32	a list to search batch size
reg_l1	1e-8	a list to search I1 factor

reg_l2	1e-8	a list to search I2 factor
dropout	0.005	a list to search dropout ratio
key	words belov	v are available for e2n2
n_features	16	a list to search number of features
n_blocks	3	a list to search number of interaction blocks
I_max	1	a list to search number of rotation order
n_rbf	8	a list to search number of radial basis
rbf_layer	2	a list to search number of RBF layers
rbf_neurons	32	a list to search number of RBF neurons
use_hpc	1	unit of energy+gradient model
retrieve	0	read results from training logfiles

depth	searches a list of parameters for hidden layers, e.g., 2 3 4 5. This keyword only works for nn.
nn_size	searches a list of parameters for number of neurons per hidden layer, e.g., 100 200 300. This keyword only works for nn.
batch_size	searches a list of parameters for batch size, e.g, 64 128. This keyword only works for nn.
reg_l1	searches a list of parameters for I1 factor, e.g., 1e–5 1e–6 1e–7. It is used when use_reg_activ, use_reg_weight, or use_reg_bias is set to I1 or I1_I2 in &EG, &EG2, &NAC, &NAC2, &SOC, and &SOC2 sections. This keyword only works for nn.
reg_l2	searches a list of parameters for I1 factor, e.g., 1e–5 1e–6 1e–7. It is used when use_reg_activ, use_reg_weight, or use_reg_bias is set to I2 or I1_I2 in &EG, &EG2, &NAC, &NAC2, &SOC, and &SOC2 sections. This keyword only works for nn.
dropout	searches a list of parameters for dropout ratio, e.g., 0.001 0.002 0.003. This keyword only works for nn.

n features searches a list of parameters for number of features, e.g., 8 16. This keyword only works for e2n2. n blocks searches a list of parameters for number of blocks, e.g. 34. This keyword only works for e2n2.

I max searches a list of parameters for number of rotation order, e.g., 1 2. In general 1 is good. This keyword only works for e2n2.

n\_rbf searches a list of parameters for number of radia basis, e.g, 16 32. This keyword only works for e2n2.

rbf layers searches a list of parameters for number of RBF layers, e.g, 2 3. This keyword only works for e2n2.

rbf neurons searches a list of parameters for number of RBF neurons, e.g, 32 64. This keyword only works for e2n2.

> submits the NN training to the job scheduler. It is turned on in default, thus the training will be submitted to SLURM as a subprocess in the current machine. For training a few NNs on a node with many cpu, it is not recommended to use use hpc because the job will have to wait in the queue while the current machine is idle. However, if there are hundreds of training in a grid search, it is better to use use hpc to distribute the calculations to all available nodes via SLURM. To use this function, you need to prepare a SLURM template with the same name as title in &CONTROL section, e.g. job title.slurm and specify the all necessary #SBATCH variables. If gpu is used (for using e2n2 mode), the SLURM tempalte need to be named as job\_title.gres. Note the difference in the file extension.

reads the logfiles of NN trainings in a completed grid-search and regenerate a logfile containing a summary of training results. No training calculation is performed. It is used when the grid search completed normally but the failed to print results. It is turned off in default.

#### 5.11. EG and EG2

use\_hpc

retrieve

&EG and &EG2		
invd_index	None	path to inverse distance indices file

depth	4	number of hidden layers
nn_size	100	number of neurons per hidden layer
batch_size	64	number of data in one batch
activ	leaky_softplus	activation function
activ_alpha	0.03	activation function coefficient alpha
loss_weights	11	weights of energy and gradient loss
use_dropout	False	turn on dropout
dropout	0.005	dropout ratio
use_reg_activ	None	turn on regularization on activation function
use_reg_weight	None	turn on regularization on weights
use_reg_bias	None	turn on regularization on bias
reg_l1	1e-5	I1 factor
reg_l2	1e-5	I2 factor
use_step_callback	True	turn on stepwise learning rate schedular
scale_x_mean	False	shift x values to mean
scale_x_std	False	scale x values to std
scale_y_mean	True	shift y values to mean
scale_y_std	True	scale y values to std
normalization_mode	1	normalize hidden layer weights
еро	2000	number of epochs
epostep	10	number of epochs for validation
learning_rate	1e-3	initial learning rate
learning_rate_step	1e-3 1e-4 1e-5 1e-6	stepwise learning rates
epoch_step_reduction	500 500 500 500	number of epochs for stepwise learning rate reduction

invd\_index

sets a path to a file containing the pairwise indices for counting inverse distance. Each line should contain a pair of atom indices. If it is not used, all pairwise distances will be included.

**depth** sets the number of hidden layers.

nn\_size sets the number of neurons per hidden layer.

**batch\_size** sets the number of training data in one batch.

activ sets the activation function. leaky softplus is used in default.

activ alpha sets the alpha coefficient in leaky\_softplus activation function.

loss weights sets the weights of energy and gradient loss in the total loss function. It

reads two values, e.g., 11

**use\_dropout** turn on dropout during the training.

dropout sets the dropout ratio. Note that dropout should not be used together with

use\_reg\_activ, use\_reg\_weight, or use\_reg\_bias.

**use\_reg\_activ** turn on regularization on activation function. Available options are:

I1 I1 regularization,

l2 regularization,

I1 I2 I1 and I2 regularization.

use reg weight turn on regularization on hidden layer weights. Available options are:

I1 I1 regularization

l2 l2 regularization

I1 I2 I1 and I2 regularization

use reg bias turn on regularization on hidden layer bias. Available options are:

I1 I1 regularization

I2 l2 regularization

I1 I2 I1 and I2 regularization

reg\_I1 sets a I1 factor. It is used when use\_reg\_activ, use\_reg\_weight, or

use reg bias is set to 11 or 11 12.

reg\_l2 sets a l2 factor. It is used when use\_reg\_activ, use\_reg\_weight, or

use reg bias is set to 12 or 11 12.

use step callback turn on the stepwise learning rate schedular. It is turned on in default.

scale x mean shift x values to their mean value. It is not recommended because x values

are inverse distances.

scale x std shift x values to their standard deviation. It is not recommended because x

values are inverse distances.

scale y mean shift y values to their mean value. It is used in default to standardize the

target data.

scale y std shift y values to their standard deviation. It is used in default to standardize

the target data.

normalization\_mode normalize the weights of hidden layer to avoid gradient explosion during

the training.

**learning\_rate** sets the initial learning rate.

epo sets the number of epochs.

**epostep** sets the number of epochs to validate the model.

**learning\_rate\_step** sets the stepwise reduced learning rates for each portion of epochs.

epoch\_step\_reduction sets the number of epochs for each portion of learning rates reduction.

#### 5.12. NAC and NAC2

&NAC and &NAC2		
invd_index	None	path to inverse distance indices file
depth	4	number of hidden layers
nn_size	100	number of neurons per hidden layer
batch_size	64	number of data in one batch
activ	leaky_softplus	activation function
activ_alpha	0.03	activation function coefficient alpha
phase_less_loss	False	use phaseless loss for nac
use_dropout	False	turn on dropout
dropout	0.005	dropout ratio
use_reg_activ	None	turn on regularization on activation function

use_reg_weight	None	turn on regularization on weights
use_reg_bias	None	turn on regularization on bias
reg_l1	1e-5	I1 factor
reg_l2	1e-5	I2 factor
use_step_callback	True	turn on stepwise learning rate schedular
scale_x_mean	False	shift x values to mean
scale_x_std	False	scale x values to std
scale_y_mean	True	shift y values to mean
scale_y_std	True scale y values to std	
normalization_mode	1	normalize hidden layer weights
еро	2000	number of epochs
epostep	10	number of epochs for validation
learning_rate	1e-3	initial learning rate
learning_rate_step	1e-3 1e-4 1e-5 1e-6	stepwise learning rates
epoch_step_reduction	500 500 500 500	number of epochs for stepwise learning rate reduction

invd_index	sets a path to a file	containing the pairwise	indices for counting inverse
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distance. Each line should contain a pair of atom indices. If it is not used,

all pairwise distances will be included.

**depth** sets the number of hidden layers.

nn\_size sets the number of neurons per hidden layer.

batch\_size sets the number of training data in one batch.

activ sets the activation function. leaky\_softplus is used in default.

activ\_alpha sets the alpha coefficient in leaky\_softplus activation function.

phase\_less\_loss use phaseless loss for nac.

**use\_dropout** turn on dropout during the training.

dropout sets the dropout ratio. Note that dropout should not be used together with use reg activ, use reg weight, or use reg bias. turn on regularization on activation function. Available options are: use reg activ 11 regularization, 11 12 12 regularization, 11 12 I1 and I2 regularization. use reg weight turn on regularization on hidden layer weights. Available options are: 11 regularization 11 12 12 regularization I1 I2 I1 and I2 regularization use reg bias turn on regularization on hidden layer bias. Available options are: 11 regularization 11 12 12 regularization I1 I2 I1 and I2 regularization sets a I1 factor. It is used when use\_reg\_activ, use\_reg\_weight, or reg\_l1 use reg bias is set to 11 or 11 12. sets a l2 factor. It is used when use\_reg\_activ, use\_reg\_weight, or reg\_l2 use reg bias is set to 12 or 11 12. use step callback turn on the stepwise learning rate schedular. It is turned on in default. shift x values to their mean value. It is not recommended because x values scale\_x\_mean are inverse distances. shift x values to their standard deviation. It is not recommended because x scale x std values are inverse distances. scale\_y\_mean shift y values to their mean value. It is used in default to standardize the target data. scale y std shift y values to their standard deviation. It is used in default to standardize the target data. normalization mode normalize the weights of hidden layer to avoid gradient explosion during the training. learning\_rate sets the initial learning rate. sets the number of epochs. epo

**epostep** sets the number of epochs to validate the model.

learning\_rate\_step sets the stepwise reduced learning rates for each portion of epochs.

epoch\_step\_reduction sets the number of epochs for each portion of learning rates reduction.

# 5.13. SOC and SOC2

&EG and &EG2		
invd_index	None	path to inverse distance indices file
depth	4	number of hidden layers
nn_size	100	number of neurons per hidden layer
batch_size	64	number of data in one batch
activ	leaky_softplus	activation function
activ_alpha	0.03	activation function coefficient alpha
use_dropout	False	turn on dropout
dropout	0.005	dropout ratio
use_reg_activ	None	turn on regularization on activation function
use_reg_weight	None	turn on regularization on weights
use_reg_bias	None	turn on regularization on bias
reg_l1	1e-5	I1 factor
reg_l2	1e-5	I2 factor
use_step_callback	True	turn on stepwise learning rate schedular
scale_x_mean	False	shift x values to mean
scale_x_std	False	scale x values to std
scale_y_mean	True	shift y values to mean
scale_y_std	True	scale y values to std
normalization_mode	1	normalize hidden layer weights

еро	2000	number of epochs
epostep	10	number of epochs for validation
learning_rate	1e-3	initial learning rate
learning_rate_step	1e-3 1e-4 1e-5 1e-6	stepwise learning rates
epoch_step_reduction	500 500 500 500	number of epochs for stepwise learning rate reduction

invd index sets a path to a file containing the pairwise indices for counting inverse

distance. Each line should contain a pair of atom indices. If it is not used,

all pairwise distances will be included.

**depth** sets the number of hidden layers.

nn\_size sets the number of neurons per hidden layer.

batch size sets the number of training data in one batch.

activ sets the activation function. leaky softplus is used in default.

activ\_alpha sets the alpha coefficient in leaky\_softplus activation function.

**use dropout** turn on dropout during the training.

dropout sets the dropout ratio. Note that dropout should not be used together with

use reg activ, use reg weight, or use reg bias.

use reg activ turn on regularization on activation function. Available options are:

I1 I1 regularization,I2 regularization,

I1 I2 I1 and I2 regularization.

**use\_reg\_weight** turn on regularization on hidden layer weights. Available options are:

I1 I1 regularizationI2 regularization

I1 I2 I1 and I2 regularization

use reg bias turn on regularization on hidden layer bias. Available options are:

I1 I1 regularization

l2 l2 regularization

I1 I2 I1 and I2 regularization

reg_ri	use_reg_bias is set to I1 or I1_I2.
reg_l2	sets a I2 factor. It is used when use_reg_activ, use_reg_weight, or use_reg_bias is set to I2 or I1 I2.

use step callback turn on the stepwise learning rate schedular. It is turned on in default.

scale\_x\_mean shift x values to their mean value. It is not recommended because x values are inverse distances.

scale\_x\_std shift x values to their standard deviation. It is not recommended because x

values are inverse distances.

scale\_y\_mean shift y values to their mean value. It is used in default to standardize the

target data.

scale\_y\_std shift y values to their standard deviation. It is used in default to standardize

the target data.

normalization\_mode normalize the weights of hidden layer to avoid gradient explosion during

the training.

**learning rate** sets the initial learning rate.

epo sets the number of epochs.

**epostep** sets the number of epochs to validate the model.

learning rate step sets the stepwise reduced learning rates for each portion of epochs.

epoch\_step\_reduction sets the number of epochs for each portion of learning rates reduction.

# 5.14. SCH\_EG

&SCH_EG		
node_features	128	number of node-embedding feature
n_features	64	number of trainable node features
n_edges	10	maximum number of neighbors

n_filters	64 number of Gaussian filters		
use_filter_bias	True add filter bias		
cfc_activ	shifted_softplus	activation function for the filters	
n_blocks	3	number of interaction blocks	
maxradius	4	maximum radius cutoff	
offset	0.0	offset of Gaussian filter centers	
sigma	0.4	width of Gaussian filters	
mlp	64	neurons per layer in the output MLI	
use_mlp_bias	True	add bias to the output MLP	
mlp_activ	shifted_softplus	activation function for the MLP	
use_output_bias	True	add bias to the output layer	
use_step_callback	True turn on stepwise learni schedular		
loss_weights	1 1 weights of energy and gradie		
еро	2000 number of epochs		
epostep	10 number of epochs for valida		
learning_rate	1e-3	initial learning rate	
learning_rate_step	1e-3 1e-4 1e-5 1e-6	stepwise learning rates	
epoch_step_reduction	500 500 500 500	number of epochs for stepwise learning rate reduction	

node_features	number of features for node embedding. It needs to be larger than the largest atomic number in the training data.
n_features	number of trainable node feature for graph convolution.
n_edges	maximum number of neighboring atoms within the radius cutoff.
n_filters	number of trainable Gaussian filters to extract the edge features.
use_filter_bias	add bias to the Gaussian filters.
cfc_activ	sets the activation function for Gaussian filters. shifted_softplus is only option.

n\_blocks number of interaction blocks. Larger number will increase the training time.

3-5 usually works well.

maxradius sets a radius in Angstrom to cut a spheric atomic environment.

offset apply an offset to the center of the Gaussiann filters.

sigma sets the widtch of the Gaussian filters. Narrower Gaussian filter requires a

greater number of filter

mlp specifies the neurons per hidden layers in the output MLP, e.g., 64 64 64

will build three hidden layers and each contains 64 neurons.

use\_mlp\_bias add bias to the output MLP layers.

mlp\_activ sets the activation function for the output MLP layers. shifted softplus is

only option.

use\_step\_callback turn on the stepwise learning rate schedular. It is turned on in default.

use\_output\_bias add bias to the last output layer.

use\_step\_callback turn on the stepwise learning rate schedular. It is turned on in default.

loss weights sets the weights of energy and gradient loss in the total loss function. It

reads two values, e.g., 11

**learning\_rate** sets the initial learning rate.

epo sets the number of epochs.

**epostep** sets the number of epochs to validate the model.

learning rate step sets the stepwise reduced learning rates for each portion of epochs.

epoch\_step\_reduction sets the number of epochs for each portion of learning rates reduction.

## 5.15. SCH\_SOC

&SCH_SOC			
node_features	128	number of node-embedding feature	
n_features	64	number of trainable node features	
n_edges	10	maximum number of neighbors	
n_filters	64	number of Gaussian filters	
use_filter_bias	True	add filter bias	
cfc_activ	shifted_softplus	activation function for the filters	
n_blocks	3	number of interaction blocks	
maxradius	4 maximum radius cutoff		
offset	0.0 offset of Gaussian filter center		
sigma	0.4 width of Gaussian filters		
mlp	64	neurons per layer in the output MLP	
use_mlp_bias	True	add bias to the output MLP	
mlp_activ	shifted_softplus	activation function for the MLP	
use_output_bias	True	add bias to the output layer	
use_step_callback	True	turn on stepwise learning rate schedular	
еро	2000	number of epochs	
epostep	10	number of epochs for validation	
learning_rate	1e-3	initial learning rate	
learning_rate_step	1e-3 1e-4 1e-5 1e-6	stepwise learning rates	
epoch_step_reduction	500 500 500 500	number of epochs for stepwise learning rate reduction	

node_features	number of features for node embedding. It needs to be larger than the largest atomic number in the training data.
n_features	number of trainable node feature for graph convolution.
n_edges	maximum number of neighboring atoms within the radius cutoff.
n_filters	number of trainable Gaussian filters to extract the edge features.

use filter bias add bias to the Gaussian filters.

cfc activ sets the activation function for Gaussian filters, shifted softplus is only

option.

n blocks number of interaction blocks. Larger number will increase the training time.

3-5 usually works well.

maxradius sets a radius in Angstrom to cut a spheric atomic environment.

offset apply an offset to the center of the Gaussiann filters.

sigma sets the widtch of the Gaussian filters. Narrower Gaussian filter requires a

greater number of filter

mlp specifies the neurons per hidden layers in the output MLP, e.g., 64 64 64

will build three hidden layers and each contains 64 neurons.

use\_mlp\_bias add bias to the output MLP layers.

mlp\_activ sets the activation function for the output MLP layers. shifted softplus is

only option.

use\_step\_callback turn on the stepwise learning rate schedular. It is turned on in default.

**use output bias** add bias to the last output layer.

use\_step\_callback turn on the stepwise learning rate schedular. It is turned on in default.

**learning rate** sets the initial learning rate.

epo sets the number of epochs.

**epostep** sets the number of epochs to validate the model.

learning rate step sets the stepwise reduced learning rates for each portion of epochs.

epoch\_step\_reduction sets the number of epochs for each portion of learning rates reduction.

## 5.16. E2N2 EG

&E2N2_EG		
n_edges	10	maximum number of neighbors
maxradius	4	maximum radius cutoff
n_features	64	number of trainable node features
n_blocks	3	number of interaction blocks
I_max	1	rotation order
parity	True	Use tensor parity
n_rbf	20	number of radial basis functions
trainable_rbf	True	trainable rbf weights
rbf_cutoff	6	exponential of the rbf cutoff function
rbf_layer	2	number of radial net hidden layer
rbf_neurons	64	number of radial net neurons/layer
rbf_act	silu	activation function for the radial net
normalization_y	component	spheric harmonic normalization scheme
normalize_y	True	Normalize spheric harmonic vectors
self_connection	True	add self-connection contribution
gate	True	use gated activation
loss_weights	11	weights of energy and gradient loss
еро	400	number of epochs
epostep	10	number of epochs for validation
subset	0 use part of training data	
batch_size	64	batch size
val_batch_size	0	validation batch size
nbatch	0	number of batch
learning_rate	1e-3	initial learning rate
learning_rate_step	1e-3 1e-4 1e-5 1e-6	stepwise learning rates
epoch_step_reduction	100 100 100 100	number of epochs for stepwise learning rate reduction

**n\_edges** maximum number of neighboring atoms within the radius cutoff.

maxradius sets a radius in Angstrom to cut a spheric atomic environment.

n\_features number of trainable node feature for graph convolution.

n blocks number of interaction blocks. Larger number will increase the training time.

3-5 usually works well.

I\_max the largest rotation order that will be kept in tensor products

parity include the parity of tensors

n\_rfb number of Bessel radial basis function

trainable\_rbf update the weights of the radial basis function during training.

**rbf\_cutoff** exponential of the cutoff function used in DimeNet.

rbf layers number of hidden layers in the radial basis network

rbf\_neurons number of neurons per hidden layer in the radial basis network

rbf act activation function in the radial basis network. SiLU is recommended.

Another option is shifted softplus.

**normalization\_y** chooses the normalization scheme for spheric harmonic vectors.

**normalize\_y** normalizes the spheric harmonic vectors.

self\_connection include self-connection when updating the node feature

gate use gated activation for tensor convolution. The activation functions are

silu for even scaler and even tensor and tanh for old scaler and old tensor.

The keywords are:
act\_scalars\_e silu
act\_scalars\_o tanh
act\_gates\_e silu
act\_gates\_e tanh

loss weights sets the weights of energy and gradient loss in the total loss function. It

reads two values, e.g., 11

epo sets the number of epochs.

**epostep** sets the number of epochs to validate the model.

**subset** use a portion of the training data for training.

batch size specify the batch size of training data. It is recommended to use 5.

val batch size specify the batch size of validation data. It is recommended to use 5.

nbatch specify the number of batches in training data. A value greater than 0 will

overwrite the **batch\_size**, the batch size will be automatically determined by the number of training data. The default value is 0. It could be useful to keep the same number of batches during adaptive sampling as the total

number of training data is increasing.

**learning\_rate** sets the initial learning rate.

learning\_rate\_step sets the stepwise reduced learning rates for each portion of epochs.

epoch\_step\_reduction sets the number of epochs for each portion of learning rates reduction.

#### 5.17. E2N2 NAC

(not available yet)

# 5.18. E2N2\_SOC

&E2N2_SOC		
n_edges	10	maximum number of neighbors
maxradius	4	maximum radius cutoff
n_features	64	number of trainable node features
n_blocks	3	number of interaction blocks
I_max	1	rotation order
parity	True	Use tensor parity
n_rbf	20	number of radial basis functions
trainable_rbf	True	trainable rbf weights
rbf_cutoff	6	exponential of the rbf cutoff function

rbf_layer	number of radial net hidden lag		
rbf_neurons	64	number of radial net neurons/layer	
rbf_act	silu	activation function for the radial net	
normalization_y	component	spheric harmonic normalization scheme	
normalize_y	True	Normalize spheric harmonic vectors	
self_connection	True	add self-connection contribution	
gate	True	use gated activation	
еро	400	number of epochs	
epostep	10	number of epochs for validation	
subset	0	use part of training data	
batch_size	64	batch size	
nbatch	0	number of batch	
learning_rate	1e-3 initial learning rate		
learning_rate_step	1e-3 1e-4 1e-5 1e-6 stepwise learning rates		
epoch_step_reduction	100 100 100 100	number of epochs for stepwise learning rate reduction	

n_edges	maximum number of neighboring atoms within the radius cutoff.
maxradius	sets a radius in Angstrom to cut a spheric atomic environment.
n_features	number of trainable node feature for graph convolution.
n_blocks	number of interaction blocks. Larger number will increase the training time. 3–5 usually works well.
I_max	the largest rotation order that will be kept in tensor products
parity	include the parity of tensors
n_rfb	number of Bessel radial basis function
trainable_rbf	update the weights of the radial basis function during training.
rbf_cutoff	exponential of the cutoff function used in DimeNet.

rbf layers number of hidden layers in the radial basis network

rbf\_neurons number of neurons per hidden layer in the radial basis network

rbf act activation function in the radial basis network. SiLU is recommended.

Another option is shifted softplus.

**normalization\_y** chooses the normalization scheme for spheric harmonic vectors.

**normalize\_y** normalizes the spheric harmonic vectors.

self connection include self-connection when updating the node feature

gate use gated activation for tensor convolution. The activation functions are

silu for even scaler and even tensor and tanh for old scaler and old tensor.

The keywords are:
act\_scalars\_e silu
act\_scalars\_o tanh
act\_gates\_e silu
act\_gates e tanh

epo sets the number of epochs.

**epostep** sets the number of epochs to validate the model.

subset use a portion of the training data for training.

batch size specify the batch size of training data.

nbatch specify the number of batches in training data. A value greater than 0 will

overwrite the **batch\_size**, the batch size will be automatically determined by the number of training data. The default value is 0. It is useful to keep the same number of batches during adaptive sampling as the total number

of training data is increasing.

**learning\_rate** sets the initial learning rate.

learning rate step sets the stepwise reduced learning rates for each portion of epochs.

epoch\_step\_reduction sets the number of epochs for each portion of learning rates reduction.

# 5.19. DIME\_NAC

The keywords, default values, and short descriptions are listed below.

&DIME_NAC		
model_type	None	Choose DimeNet model
batch_size	64	batch size
val_size	64	validation size
hidden_channels		
blocks	3	number of interaction blocks
bilinear	True	Use tensor parity
spherical	20	number of radial basis functions
radial	True	trainable rbf weights
Ir	1e-3	initial learning rate
еро	400	number of epochs

Full descriptions for all available keywords are summarized below.

None DimeNet model. This is default.

pp DimeNet++ model.

batch size specify the batch size of training data.

val\_size specify the size of validation data.

**hidden\_channels** specify the number of hidden channels.

blocks specify the number of interaction blocks.

bilinear specify the number of bilinear functions.

**spherical** specify the number of spherical functions.

radial specify the number of radial functions.

Ir specify the starting learning rate.

epo specify the number of training epochs

#### 5.20. FILE

The keywords, default values, and short descriptions are listed below.

&FILE		
natom	0	number of atoms
file	None	path to a list file to read QC calculation results

Full descriptions for all available keywords are summarized below.

natom sets the number of atoms for reading the coordinates from the QC

calculation logfiles.

file read the path to a list file for extracting the QC data from the calculation

logfiles. If a file name is provided, it assumes that the list file is in the current folder. In the list file, each line should contain a path to a QC calculation

folder.

# 6. Nonadiabatic molecular dynamics

# 5.1. Fewest switches surface hopping

## 5.2. Zhu-Nakamura surface hopping

# 7. Machine learning models

## 6.1. Preparing training data

# 6.2. Creating a neural network

First, we create a model to predict energies, forces, and non-adiabatic couplings (if requested). The input example below shows the frequently used keywords for creating modes.

**jobtype** determines the type of calculation. It takes 'train' for training neural networks, 'prediction' for predicting e

nergies, forces, and non-adiabatic couplings, 'adaptive' for adaptive sampling of conformational space using molecular dynamics trajectories, and 'md' for molecular dynamics simulation.

PyRAI<sup>2</sup>MD has a flexible training scheme depending on the available computing resources. When **ml\_ncpu = 1**, all models will be trained sequentially. When **ml\_ncpu <=4**, all models will be trained in subprocess so they can use all given numbers of CPUs. If **ml\_ncpu > 4**, the extra CPU resources will be used to parallelize the training, which is automatically managed by TensorFlow.

- 6.3. Training a neural network
- 6.5. Adaptive sampling

- 8. External quantum chemical program
- 7.1. Molcas
- 7.2. BAGEL
- 7.3. ORCA

- 7.4. OpenQP
- 7.5. GFN-xTB
- 7.6. MNDO