$Documentation \ for \ \textbf{Hawaii} \ \ \textbf{Hybrid} \ v.0.1$

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1 Problems addressed by library

- 1. Compute statistical averages over ensembles of classical trajectories restricted by chosen phase-space domains (e.g., bound or unbound states), enabling the calculation of dipole autocorrelation functions.
- 2. Calculate static phase-space averages using rejection sampling and adaptive Monte Carlo algorithms to estimate zeroth and second spectral moments.
- 3. Transform correlation functions into spectral functions, apply smoothing, and convert them into spectral profiles.

To ensure clarity, we will outline the key objects and their corresponding units of measurement. In the context of this document, the correlation function of the dipole moment, C(t), is defined as:

$$C(t) = V\langle \mu(0)\mu(t)\rangle. \tag{1}$$

Note that this definition differs from others, such as the one in Ref. [1], as it does not include the $1/(4\pi\varepsilon_0)$ factor. Instead, this factor is incorporated into the definition of the spectral function. The correlation function values are produced by calculate_correlation_and_save and calculate_correlation_array_and_save in units of (m³· atomic unit of dipole). The time is treated internally in atomic time units, and the produced CFnc object stores the time in the atomic time units. However, when the correlation function is to be processed in order to be converted into spectral function (by fit_baseline function), time axis needs to be in seconds.

The spectral function is defined as follows:

$$G(\nu) = \frac{1}{2\pi} \frac{1}{4\pi\varepsilon_0} \int_{-\infty}^{+\infty} C(t)e^{-2\pi ic\nu t} dt,$$
 (2)

where ν is the wavenumber (cm⁻¹). The values of spectral function are $J \cdot m^6 \cdot s = kg \cdot m^8 \cdot s$. The binary absorption coefficient is then related to the spectral function according to:

$$\alpha(\nu) = \frac{\tau(\nu)}{\rho_1 \rho_2} = \frac{(2\pi)^4 N_L^2}{3h} \nu \left[1 - \exp\left(-\frac{hc\nu}{k_B T}\right) \right] G(\nu), \tag{3}$$

where the absorption coefficient is

$$\tau(\nu) = L^{-1} \ln(I_0/I),\tag{4}$$

and ρ_1 and ρ_2 are gas densities in mixture. We will express the binary absorption coefficient as per convention in cm⁻¹Amagat⁻².

2 Input file format

This section provides an overview of the input file format, including its structures, features, and supported data types. The format uses &SECTION_NAME and &END delimiters to define sections containing case-insensitive key-value pairs (assigned with =) or function calls. Comments are indicated with exclamation marks! throughout the input file. The input file parser enforces type safety, requiring each key to have a value of the expected type; any type mismatch triggers an error message and terminates program execution.

Supported data types include integers, floating-point values, boolean values, strings and arrays. Underscores may be used in integers for readability (e.g., 1_000_000). For floating point values the scientific notation may be used (e.g., 1e2 instead of 100). The arrays are denoted with curly braces {}.

&INPUT section provides overall control of the calculation, for example, calculation type, pair state, reduced mass, paths to dynamic libraries containing potential energy and induced dipole functions, and output files. The available input parameters are described in Section 2.1.

&MONOMER section specifies the types of interacting monomers through the MONOMER_TYPE field along with their molecular constants such as inertia tensor values and centrifugal constant. The available parameters are

listed in Section 2.2.

&PROCESSING section configures the data pipeline using a stack machine model, where each command manipulates the implicit data stack. Unlike the &INPUT and &MONOMER sections, the &PROCESSING section also contains an order-sensitive sequence of commands that execute sequentially, consuming inputs from the top of the stack and pushing the results back for subsequent commands. A typical pipeline begins with loading data using read_cf, which pushes a correlation function onto the stack. Then cf_to_sf consumes the correlation function and pushes corresponding spectral function. Commands like compute_m0_classical_detailed_balance and compute_m2_classical_detailed_balance consume the spectral function, requiring the user to first call dup() to preserve the original spectral function if further commands will be operating on it. By the end of the processing script, the stack must be empty, ensuring all data has been processed or saved. The available functions are described in Section 2.3.

2.1 INPUT section

Here we provide a detailed description of keywords handled in the INPUT block.

- #1 CALCULATION_TYPE
- #2 PAIR_STATE
- #3 PAIR_REDUCED_MASS
- #4 SO_POTENTIAL
- #5a SO_DIPOLE
- #5b SO_DIPOLE.1
- #5c SO_DIPOLE.2
- #6 TEMPERATURE
- **#7** TEMPERATURES
- #8 SATELLITE_TEMPERATURES
- #9 TOTAL_TRAJECTORIES
- #10 CVODE_TOLERANCE
- #11 SAMPLING_TIME
- #12 MAXTRAJECTORYLENGTH
- #13 SAMPLER_RMIN
- #14 SAMPLER_RMAX
- #15 PESMIN
- #16 INITIALMO_NPOINTS
- #17 INITIALM2_NPOINTS
- #18 HEP_MO_NPOINTS
- #19 HEP_MO_NITERATIONS
- #20 HEP_M2_NPOINTS
- #21 HEP_M2_NITERATIONS
- #22 HEP_PPF_NITERATIONS
- #23 HEP_PPF_NPOINTS
- #24 SF_FILENAME
- #25 CF_FILENAME
- #26 CF_FILENAMES
- #27 R0
- #28 RCUT
- #29 PARTIAL_PARTITION_FUNCTION_RATIO
- #30 PARTIAL_PARTITION_FUNCTION_RATIOS
- #31 APPROXIMATEFREQUENCYMAX
- #32 ODD_J_SPIN_WEIGHT
- #33 EVEN_J_SPIN_WEIGHT
- #34 USE_ZIMMERMANN_TRICK
- #35 AVERAGE_TIME_BETWEEN_COLLISIONS
- #1. CALCULATION_TYPE. The program performs calculations based on the specified CALCULATION_TYPE. This

field is required and expected to be a plain string (no quotation marks). Note that there is no default value - the field must be explicitly provided. Supported calculation types:

- 1. CALCULATION_PR_MU
- 2. CALCULATION_CORRELATION_SINGLE
- 3. CALCULATION_CORRELATION_ARRAY
- 4. CALCULATION_PROCESSING
- 5. CALCULATION_PHASE_SPACE_MO
- 6. CALCULATION_PHASE_SPACE_M2
- #2. PAIR_STATE. The pair state is specified using the keyword PAIR_STATE as a plain string (no quotation marks). This field is required, no default value is implied. This parameter is applicable for all calculation types except for CALCULATION_PROCESSING, where it's ignored. Available pair states:
 - 1. FREE_AND_METASTABLE
 - 2. BOUND
 - 3. ALL
- #3. PAIR_REDUCED_MASS. The reduced mass for a pair is specified as a floating-point number. This parameter is applicable for all calculation types except for CALCULATION_PROCESSING.
- #4. SO_POTENTIAL specifies a double-quoted string path to a dynamic library containing the required symbols double pes_lab(double*) and void dpes_lab(double*, double*). Missing symbols trigger immediate termination. Functions pes_lab and dpes_lab operate on laboratory-frame coordinates and compute the potential energy and its derivative (outputted through the second pointer), respectively. If present, a symbol void pes_init(void) will be executed before any calls pes_lab and dpes_lab will be issued, enabling library initialization (e.g., loading the parameters from file).
- #5a. SO_DIPOLE specifies a double-quoted string path to a dynamic library implementing the required symbol void dipole_lab(double *, double [3]), which computes laboratory-frame dipole moment components from input coordinates in the laboratory frame of reference. Missing symbol trigger immediate termination. The loaded function is assigned to dipole_1 and dipole_2, enabling calculation of autocorrelation functions. if present, a symbol void dipole_init(void) will be executed before any calls dipole_lab will be issued, enabling library initialization. This setting is equivalent to assigning the same path to both SO_DIPOLE.1 and SO_DIPOLE.2.
- #5b, c. SO_DIPOLE.1, SO_DIPOLE.2 specify independent paths to dipole function implementations as dynamic libraries. The required and optional symbols match those mentioned in #5.a. Setting these paths to different dynamic libraries enables the calculation of mixed correlation correlation.
- #6. TEMPERATURE

2.2 MONOMER section

Here we provide a detailed description of keywords handled in the MONOMER block.

```
MONOMER_TYPE
#36
#37
     DJ
#38
     II
#39
     INITIAL_J
#40
     TORQUE_CACHE_LEN
#41
     TORQUE_LIMIT
#42
     NSWITCH_HISTOGRAM_BINS
#43
     NSWITCH_HISTOGRAM_MAX
#44
     NSWITCH_HISTOGRAM_FILENAME
#45
     JINI_HISTOGRAM_BINS
#46
     JINI_HISTOGRAM_MAX
#47
     JINI_HISTOGRAM_FILENAME
#48
     JFIN_HISTOGRAM_BINS
#49
     JFIN_HISTOGRAM_MAX
#50
     JFIN_HISTOGRAM_FILENAME
```

2.3 PROCESSING section

Here we provide a detailed description of keywords

#51 SPECTRUM_FREQUENCY_MAX

and internal functions handled in the MONOMER block.

```
READ_CF(\1)
#52
     DUP()
#53
#54
     INT3()
#55
    AVERAGE_CFS()
#56
     COMPUTE_MO_CLASSICAL_DETAILED_BALANCE()
#57
     COMPUTE_M2_CLASSICAL_DETAILED_BALANCE()
#58 FIT_BASELINE()
#59
     CF_TO_SF()
#60
     ALPHA()
#61
     D3()
#62
     WRITE_CF(\1)
#63
     WRITE_SF(\1)
     WRITE_SPECTRUM(\1)
#64
```

3 Examples

3.1 Propagating trajectory for CO_2 -Ar

As an initial illustration of the use of Hawaii Hybrid package, we give a sample program called trajectory_co2_ar.cpp to propagate a trajectory of collisional dynamics of CO_2 -Ar system. The problem consists of solving the following Hamilton dynamical equations:

	t_{P} d	Phase-Space moments	single correlation function	^{an} ay of correlation functions
He-Ar	✓	√	✓	×
CO-Ar	×	×	\checkmark	×
CO_2 -Ar	\checkmark	\checkmark	\checkmark	\checkmark
N_2 -Ar	×	\checkmark	\checkmark	\checkmark
H_2 -Ar	\checkmark	×	×	×
CH_4-CO_2	✓	✓	✓	✓

$$\begin{split} \dot{R} &= \frac{p_R}{m}, \\ \dot{p}_R &= \frac{p_\Theta^2}{mR^3} + \frac{p_\Phi^2}{mR^3 \sin^2 \Theta} - \frac{\partial U}{\partial R}, \\ \dot{\Phi} &= \frac{p_\Phi}{mR^2 \sin^2 \Theta}, \\ \dot{p}_\Phi &= -\frac{\partial U}{\partial \Phi}, \\ \dot{P}_\Phi &= -\frac{\partial W}{\partial \Phi}, \\ \dot{P}_\Theta &= \frac{p_\Theta}{mR^2}, \\ \dot{p}_\Theta &= \frac{p_\Phi^2 \cos \Theta}{mR^2 \sin^3 \Theta} - \frac{\partial U}{\partial \Theta}, \\ \dot{p}_1 &= \frac{p_1^\varphi}{I_1 \sin^2 \vartheta_1}, \\ \dot{p}_1^\varphi &= -\frac{\partial U}{\partial \varphi_1}, \\ \dot{p}_1^\varphi &= \frac{p_1^\vartheta}{I_1}, \\ \dot{p}_1^\theta &= \frac{(p_1^\varphi)^2 \cos \theta_1}{I_1 \sin^3 \theta_1} - \frac{\partial U}{\partial \theta_1}. \end{split}$$

The hawaii.h header files provides definition of the struct MoleculeSystem that is used to evaluate the right-hand side of the dynamical equations. The trajectory.h header provides struct Trajectory that provides interface with CVode library that performs the integration of dynamical equations (see Section ??). Next, we provide implementations for functions that calculate potential energy pes and its derivatives with respect to coordinates dpes. Each user program that instantiates the MoleculeSystem is expected to implement these functions. To implement these functions hawaii provides functions to transform angles between different reference frames (see Section 4.9).

- 3.2 Propagating trajectory for H_2 -Ar while requantizing the angular momentum of H_2
- 3.3 Calculating the zeroth and second spectral moments of CO_2 -Ar as phase-space averages using rejection-based sampler

This example demonstrates how to use interface with hep to calculate spectral moments. First, the zeroth moment over unbound states is calculated as follows:

$$M_0 = \frac{(2\pi)^4 N_L^2}{3(Q/V)h} \frac{1}{4\pi\epsilon_0} \frac{1}{2\pi c} \int_{\Omega: H > 0} \mu^2 \exp\left(-\frac{H}{kT}\right) d\mathbf{q} d\mathbf{p}.$$

The integral is evaluated using mpi_perform_integration, whereas the partition can be calculated using analytic_full_partition_function_by_V.

- 3.4 Calculating a single correlation function for CO₂-Ar
- 3.5 Calculating a spectral function using p_r/μ -representation for CO_2 -Ar
- 3.6 Calculating an array of correlation functions for CO₂-Ar
- 3.7 Processing correlation function for CO₂-Ar

4 Complete list of function and code organization

4.1 Setting up a molecule pair

enum MonomerType

Values

ATOM = O

LINEAR_MOLECULE = 4

LINEAR_MOLECULE_REQ_INTEGER = MODULO_BASE + 4

LINEAR_MOLECULE_REQ_HALFINTEGER = 2*MODULO_BASE + 4

/* LINEAR_VIBRATING_MOLECULE = MODULO_BASE + 6 */

ROTOR = 6

ROTOR_REQUANTIZED_ROTATION = 2*MODULO_BASE + 6

Description

This enum is used to distinguish between systems of different types and store the size of the phase point: size(phase_point) = MonomerType % MODULO_BASE, where MODULO_BASE is #defined to 100 by default.

struct Monomer

Fields

MonomerType t

double I[3] – values of tensor of inertia

 ${\tt double}\ {\tt DJ-centrifugal\ distortion\ constant}$

double *qp - dynamic variables (currently, Euler angles and conjugated momenta) at the current step of simulation

double *dVdq - the derivatives of potential energy with respect to coordinates pertaining to this monomer (the order of coordinates is the same as for qp)

bool apply_requantization

Description

The apply_requantization will be set to true in rhs to signal that the requantization of the monomer's angular momentum is required during trajectory propagation. The order of variables in the qp array is specified by the following indices:

#define IPHI 0

#define IPPHI 1

#define ITHETA 2

#define IPTHETA 3

#define IPSI 4
#define IPPSI 5

enum PairState

Values

 $FREE_AND_METASTABLE = 0$

BOUND = 1

Description

struct MoleculeSystem

Fields

double intermolecular_qp[6] – Coordinates and conjugated momenta that correspond to the intermolecular motion: $(\Phi, p_{\Phi}, \Theta, p_{\Theta}, R, p_R)$.

 ${\tt Monomer}\ {\tt m1}$

Monomer m2

double mu - reduced mass of molecule pair

size_t Q_SIZE - total number of coordinates for molecule pair

size_t QP_SIZE - total number of coordinates and momenta for molecule pair (a.k.a. size(phase_point))

double *intermediate_q - contiguous vector of coordinates

double *dVdq - contiguous vector of potential energy derivatives

time_t init_rawtime - the initialization time, represented as the number of seconds since the Unix Epoch; set in init_ms

time_t temp_rawtime - a temporary time field for the time at which the previous iteration was completed, used for tracking elapsed time between iterations

Description

Keep in mind that angular variables and momenta are stored in the same order as for qp in Monomer. These variables' locations are #defined as follows:

#define IPHI 0

#define IPPHI 1

#define ITHETA 2

#define IPTHETA 3

#define IR 4

#define IPR 5

Keep in mind that intermolecular coordinates and monomer's coordinates are not stored contiguously. The contiguous vector of coordinates can be assembled by calling <code>extract_q_and_write_into_ms</code> function, which stores the coordinates in memory pointed at by <code>intermediate_q</code>. These coordinates are passed to external functions that compute the values of intermolecular energy, its derivatives with respect to coordinates and induced dipole (see section ??). There is no guarantee that coordinates stored in <code>Monomer</code>'s and coordinates in memory at <code>intermediate_q</code> are always in sync. The function <code>extract_q_and_write_into_ms</code> must be invoked if the contiguous vector of coordinates is desired at a certain point of the program execution.

Function init_ms

Call

MoleculeSystem *ms = MoleculeSystem init_ms(mu, t1, t2, I1, I2, seed)

Arguments

double mu - the reduced mass of the molecule pair

 ${\tt MonomerType\ t1\ specifies\ the\ type\ of\ first\ monomer}$

MonomerType t2 specifies the type of second monomer

double* I1 contains inertia tensor values for first monomer. If the monomer is atom, no values will be read from the pointer, so NULL can be passed. Two and three values are expected for the rotor and linear molecule, respectively.

double* I2 contains inertia tensor values for second monomer.

size_t seed is the seed for random number generator. A unique seed will be produced if 0 is passed.

Description

The function prepares the MoleculeSystem struct based on the specified monomer types, allocates the memory using malloc (probably should be avoided due to cache locality) and initializes the random number generator.

Function kinetic_energy

Call double kinetic_energy(*ms)

Arguments MoleculeSystem* ms

Description The kinetic energy function is calculated at the phase-point stored in MoleculeSystem. Currently,

implemented for intermolecular degrees of freedom, linear molecule (tested) and rotor (tested).

Parts of the kinetic energy corresponding to intermolecular degrees of freedom:

$$T_{\rm int} = \frac{p_R^2}{2\mu} + \frac{p_{\Theta}^2}{2\mu R^2} + \frac{p_{\Phi}^2}{2\mu R^2 \sin^2 \Theta};$$

to linear molecule:

$$T_{\rm lin} = \frac{p_{\theta}^2}{2I} + \frac{p_{\varphi}^2}{2I\sin^2\theta};$$

to rotor:

$$\begin{split} T_{\text{rotor}} &= \frac{1}{2I_1 \sin^2 \theta_2} \Big[\left(p_\varphi - p_\psi \cos \theta \right) \sin \psi + p_\theta \sin \theta \cos \psi \Big]^2 + \\ &+ \frac{1}{2I_2 \sin^2 \theta} \Big[\left(p_\varphi - p_\psi \cos \theta \right) \cos \psi - p_\theta \sin \theta \sin \psi \Big]^2 + \frac{1}{2I_3} \left(p_\psi \right)^2. \end{split}$$

Function Hamiltonian

Call double Hamiltonian(*ms)

Arguments MoleculeSystem* ms

Description Calls to kinetic energy, assembles a contiguous vector of coordinates via extract_q_and_write_into_ms and passes it to external pes.

The potential energy and its derivatives are expected to be provided as functions with the following signatures:

Function pes

Call double pes(*q);

Arguments double *q

Description An array of Q_SIZE variables is passed to this function; MoleculeSystem describes their order.

Intermolecular distance measured in Bohrs, while angles are measured in radians. The function is

expected to return energy in Hartree.

Function dpes

Call void dpes(*q, *dpesdq);

Arguments double *q

double *dpesdq

Description An array of Q_SIZE variables is passed to this function; MoleculeSystem describes their order.

Intermolecular distance measured in Bohrs, while angles are measured in radians. The function is expected to return an array of derivatives of energy in units of Hartree/Bohr of length Q_SIZE.

Function dipole

Call void dpes(*q, *dpesdq);

Arguments double *q

double *dpesdq

Description

pes and dpes are defined as external functions in the header hawaii.h, so they have to be implemented in the user program, whereas the dipole function needs to be set through function pointer named dipole.

4.2 Sampling the phase-space point

Call void q_generator(*ms, *params)

Arguments MoleculeSystem* ms

CalcParams *params

Description Generates R with density $\rho \sim R^2$ in the range [params.sampler_Rmin, params.sampler_Rmax].

The distributions of φ, ψ are $\varphi, \psi \sim U[0, 2\pi]$ and for θ is $\cos \theta \sim U[0, 1]$. Implemented for inter-

molecular degrees of freedom, linear molecule and rotor.

Function p_generator

Call void p_generator(*ms, T)
Arguments MoleculeSystem* ms

double T

Description Samples momenta p from distribution $\rho \sim e^{-K/kT}$ at given temperature. Switches between

p_generator_linear_molecule and p_generator_rotor to sample momenta for monomers.

Function reject

Call bool reject(*ms, Temperature, pesmin)

Arguments MoleculeSystem* ms

double Temperature

double pesmin - the minimum value of PES

Description Applies the rejection step to the phase-point that is stored in the MoleculeSystem. It presupposes

that the provided phase-point is sampled from $\rho \sim e^{-K/kT}$ using q-generator and p-generator functions. The random variable $u \sim U[0,1]$ is chosen, to determine whether the current phase-point

is to be accepted with probability $\rho \sim \exp(-H/kT)$.

4.3 Calculating a trajectory: interfacing with CVode library

struct Trajectory

Fields size_t DIM

size_t mxsteps
N_Vector y
N_Vector abstol
N_Vector reltol
SUNMatrix A

SUNLinearSolver LS
void *cvode_mem

Description

Function init_trajectory

Call Trajectory init_trajectory(*ms, *transformed, reltol);

Arguments MoleculeSystem *ms

double reltol

Description

Function free_trajectory

Call void free_trajectory(*traj);

Arguments Trajectory *traj

Description

Function | reinit_trajectory Call void reinit_trajectory(*traj, t); Trajectory *traj Arguments double t Description Function | make_step Call int make_step(*traj, tout, *t); Arguments Trajectory *traj double tout double *t Description Invokes the CVode main function and, if enabled, applies requantization to the monomers' angular momenta. When the apply_requantization flag is set, the momenta in qp are rescaled to adjust the angular momentum to the nearest half-integer or integer, according to the requantization rule which is represented with the Monomer type. Subsequently, the function synchronizes the internal trajectory phase-space vector with the MoleculeSystem phase-space vector, and reinitializes the state of CVode to reset its state since a sudden change of phase-point has occured. Function set_initial_condition Call void set_initial_condition(*traj, qp); Arguments Trajectory *traj Array qp Description Function | make_vector Call N_Vector make_vector(size); Arguments int size Description Function | set_tolerance Call void set_tolerance(*traj, tolerance); Arguments Trajectory *traj * double tolerance Description Function | rhs Call void rhs(t, y, ydot, *user_data); Arguments UNUSED(realtype t) N_Vector y stores coordinates and conjugated momenta N_Vector ydot is filled by function with numerical values of right-hand side of Hamilton's equations of motion at provided phase point void *user_data is employed to pass MoleculeSystem* inside the function (see section ??) Description

This function is passed to CVode library to propagate the trajectory (see section ??). First, the phase-point coordinates are stored into MoleculeSystem struct. A contiguous vector of coordinates is assembled via extract_q_and_write_into_ms. Next, by calling the external function dpes, the derivatives of potential energy are computed and stored into MoleculeSystem.dVdq. The components of derivative vector are then copied into the field Monomer.dVdq of the corresponding monomer via the call to extract_dVdq_and_write_into_monomers. The right-hand side of Hamilton's equations with respect to intermolecular degrees of freedom are readily obtained and filled into ydot, while the derivatives with respect to monomer's coordinates are handled by rhsMonomer function.

Function rhsMonomer

Call void rhsMonomer(*m, *deriv);

Arguments Monomer *m

double *deriv stores the right-hand side of Hamilton's equations of motion with respect to coor-

dinates and momenta that correspond to the passed-in monomer

Description In addition to differentiating the kinetic energy, the derivatives of potential energy, which are

taken from Monomer.dVdq, are also added to compute the right-hand side. Implemented for cases

of LINEAR_MOLECULE (tested) and ROTOR (tested).

Function | compute_numerical_rhs

Call Array compute_numerical_rhs(*ms, order);

Arguments MoleculeSystem *ms

size_t order - the order of the central finite-difference formula (implemented for 2, 4 or 6)

Description Computes numerically the right-hand side of the Hamiltonian equations of motion. The order

corresponds to the order of variables employed in MoleculeSystem struct.

4.4 Conducting angular momentum requantization

Function j_monomer

Call void j_monomer(m, j);

Arguments Monomer m

double j[3]

Description Computes the magnitude of angular momentum of passed-in monomer. Currently, implemented

only for linear molecules.

[linear molecule] :
$$j = \begin{bmatrix} -p_{\theta} \sin \varphi - p_{\varphi} \cos \varphi / \tan \theta \\ p_{\theta} \cos \varphi - p_{\varphi} \sin \varphi / \tan \theta \\ p_{\varphi} \end{bmatrix}$$

Function torque_monomer

Call double torque_monomer(m);

Arguments Monomer m

Description Computes the magnitude of torque of passed-in monomer. Needs the derivative of potential m. dVdq

to be set within the monomer. Currently, implemented only for linear molecules.

$$[\text{linear molecule}]: \tau = \begin{bmatrix} \sin\varphi \frac{\mathrm{d}V}{\mathrm{d}\theta} + \cos\varphi/\tan\theta \frac{\mathrm{d}V}{\mathrm{d}\varphi} \\ -\cos\varphi \frac{\mathrm{d}V}{\mathrm{d}\theta} + \sin\varphi/\tan\theta \frac{\mathrm{d}V}{\mathrm{d}\varphi} \\ -\frac{\mathrm{d}V}{\mathrm{d}\varphi} \end{bmatrix}$$

Function | find_closest_integer

Call double find_closest_integer(j);

Arguments double j

Description Requantizes a given value of the angular momentum to the nearest integer, effectively rounding

to the closest whole number (0, 1, 2, 3 ...). Note that the value returned as double-precision value. Using the floor function as the implementation results in artifact trajectories with angular

momentum being brought to zero.

Function | find_closest_half_integer

Call double find_closest_half_integer(j);

Arguments double j

Description Requantizes a given value of the angular momentum to the nearest half-integer, effectively rounding

to the closest value of the form $(0,\,1.5,\,2.5,\,3.5...)$. Note that 0.5 is skipped to represent the S-band

rotational structure of the homonuclear diatomics (see Sokolov coursework).

4.5 Computing averages over phase-space

 $Function \ \middle| \ analytic_full_partition_function_by_V$

Call double analytic_full_partition_function_by_V(*ms, *params, Temperature)

Arguments MoleculeSystem *ms

double Temperature

Description Returns analytically calculated value for the ratio of the partition function for the totality of states

to the volume. Currently, implemented only for atom-atom, linear molecule-atom and rotor-linear

molecule.

$$[\text{linear molecule - atom}]: Q = 4\pi \times (2\pi\mu kT)^{3/2} \times (2\pi IkT)$$

$$[\text{rotor - linear molecule}]: Q = 32\pi^3 \times (2\pi\mu kT)^{3/2} \times (2\pi kT)^{3/2} \sqrt{I_1^{\text{rotor}}I_2^{\text{rotor}}I_3^{\text{rotor}}} \times (2\pi I_1^{\text{lin}}kT)$$

Function calculate_MO

Call void calculate_MO(*ms, *params, Temperature, *m, *q);

Arguments MoleculeSystem *ms

CalcParams *params double Temperature

double *m - the estimate of M_0 double *q - the error of the estimate

Description

By sampling from $\rho \sim e^{-K/kT}$ and rejecting some of the points using the reject function, params.initialMO_npoints phase-points are produced to estimate M_0 and its error. The average over the sampled phase-points is multiplied by the params.partial_partition_function_ratio, which is supposed to be a ratio of the partition function over the part of the phase space that is pertinent to the select pair state to the total partition function and multiplied by volume. This ratio can be calculated using mpi_perform_integration function that invokes hep library.

Function | calculate_M2

Call void calculate_MO(*ms, *params, Temperature, *m, *q);

Arguments MoleculeSystem *ms

CalcParams *params double Temperature

double *m - the estimate of M_2 double *q - the error of the estimate

Description

By sampling from $\rho \sim e^{-K/kT}$ and rejecting some of the points using the reject function, params.initialM2_npoints phase-points are produced to estimate M_2 and its error. The derivative of dipole with respect to coordinates is done using cental 2-point finite-difference formula. The energy is differentiated with respect to momenta using compute_dHdp. Here we use GSL's wrappers over BLAS to conduct matrix-by-vector multiplication. The average over the sampled phase-points is multiplied by the params.partial_partition_function_ratio, which is supposed to be a ratio of the partition function over the part of the phase space that is pertinent to the select pair state to the total partition function and multiplied by volume. This ratio can be calculated using mpi_perform_integration function that invokes hep library.

MPI Function mpi_calculate_MO

Call void calculate_MO(*ms, *params, Temperature, *m, *q);

Arguments MoleculeSystem *ms

CalcParams *params
double Temperature

double *m
double *q

Description The task of iterating params.initialMO_npoints points is split equally between processes of

communicator. The behavior is the same as in calculate_MO.

MPI Function | mpi_calculate_M2

Call void calculate_M2(*ms, *params, Temperature, *m, *q);

Arguments MoleculeSystem *ms

CalcParams *params double Temperature

double *m
double *q

Description The task of iterating params.initialM2_npoints points is split equally between processes of

communicator. The behavior is the same as in calculate_M2.

Function compute_dHdp

Call void compute_dHdp(*ms, *dHdp);

Arguments MoleculeSystem *ms

gsl_matrix *dHdp

Description Calls to rhsMonomer to fill in derivatives of Hamiltonian with respect to momenta pertaining to

monomers.

Function | transform_variables

Call void transform_variables(x, *transformed, *Jac);

Arguments hep::mc_point<double> const& x

double *transformed

double *Jac

Description Transforms a point x from n-dimensional hypercube $[0,1]^n$ to a point in phase-space of the

MoleculeSystem (which is passed in using global pointer). The following transformations are

implemented

$$R \leftarrow 1/x,$$

$$\theta \leftarrow \pi x,$$

$$\phi \leftarrow 2\pi x,$$

$$p \leftarrow \tan(\pi(x - 1/2)).$$

The jacobian of the transformation is accumulated along the steps. Implemented for linear molecule-atom and rotor-linear molecule pairs.

Function integrand_pf

Call double integrand_pf(x);

Arguments hep::mc_point<double> const& x

Description Based on the point in hypercube returns the value of the following integrand:

$$\operatorname{jac} \cdot \exp\left(-\frac{E}{kT}\right)$$
.

(The energy is computed for MoleculeSystem that is passed in using global pointer.)

Function integrand_MO

Call double integrand_MO(x);

Arguments hep::mc_point<double> const& x

Description Based on the point in hypercube returns the value of the following integrand:

$$jac \cdot \mu^2 \cdot \exp\left(-\frac{E}{kT}\right).$$

(The energy is computed for MoleculeSystem that is passed in using global pointer. The pointer to dipole function is also assumed to be set globally.)

Function integrand_M2

Call double integrand_M2(x);

Arguments hep::mc_point<double> const& x

Description Based on the point in hypercube returns the value of the following integrand:

$$\operatorname{jac} \cdot \left[\mu, H\right] \cdot \exp\left(-\frac{E}{kT}\right).$$

The derivative of dipole with respect to coordinates is computed using 2-point central finite-difference formula. (The energy is computed for MoleculeSystem that is passed in using global pointer. The pointer to dipole function is also assumed to be set globally.)

Function mpi_perform_integration

Call double mpi_perform_integration(*ms, integrand, *params, Temperature, niterations,

npoints, *m, *q);

Arguments MoleculeSystem *ms

Integrand integrand - typedef double (*Integrand)(hep::mc_point<double> const&)

CalcParams *params double Temperature size_t niterations size_t npoints double *m

double *q

Description Evaluates the integral of requested integrand

4.6 Performing averaging over trajectory ensembles

struct CalcParams

Fields PairState ps /* sampling */ double sampler_Rmin // a.u. double sampler_Rmax // a.u. double pesmin // Hartree /* initial spectral moments check */ size_t initialMO_npoints size_t initialM2_npoints double partial_partition_function_ratio /* requantization */ size_t torque_cache_len double torque_bound /* trajectory */ double sampling_time size_t MaxTrajectoryLength

```
bool allow_truncating_trajectories_at_length_limit
double cvode_tolerance
/* applicable to both correlation function AND spectral function calculations */
size_t niterations
size_t total_trajectories
/*correlation function calculation ONLY */
const char *cf_filename
double Rcut // distance at which the trajectory is forcefully stopped, a.u.
/* pr/mu calculation ONLY */
const char *sf_filename
double ApproximateFrequencyMax // cm-1
double RO // initial distance, a.u.
double average_time_between_collisions // a.t.u.
/*correlation function array ONLY */
double* partial_partition_function_ratios
double *satellite_temperatures
size_t num_satellite_temperatures
const char **cf_filenames
```

struct CFnc

```
Fields double *t
double *data
size_t len - number of samples in *t and *data
size_t capacity - capacity of *t and *data
size_t ntraj - number of trajectories used for averaging
double Temperature
bool normalized - flag that indicates whether the data samples are normalized by # of trajectories
```

struct CFncArray

```
Fields

double *t

double **data

size_t ntemp

size_t len - number of samples in *t and elements of *data

double* nstar - number of effective trajectories at each temperature

size_t ntraj - number of calculated trajectories at base temperature

Description

An array of correlation functions for a fixed base temperature (sampling temperature) and a

satellite temperature (target temperature for re-weighting).
```

4.6.1 Calculating correlation functions

Function correlation_eval

```
Call int correlation_eval(*ms, *traj, *params, *crln, *tps);

Arguments MoleculeSystem *ms

Trajectory *traj
CalcParams *params
double *crln
int *tps

Description Evaluates the mean between forward and backward correlation functions from the initial condition set in the *ms. It also tracks the number of turning points and returns it using the output parameter
```

MPI Function calculate_correlation_and_save

*tps.

Call CFnc calculate_correlation_and_save(*ms, *params, Temperature);

Arguments MoleculeSystem *ms

CalcParams *params
double Temperature

Description First, esti

First, estimates of the zeroth and second special moments are obtained using params.initialMo_npoints and params.initialMo_npoints points, respectively. The accumulation of params.total_trajectories individual correlation functions is divided into params.niterations iterations. The individual correlation functions are obtained using correlation_eval function. The current aggregate estimate of the correlation function is saved to params.cf_filename at the end of each iteration. The zeroth moment based on the current estimate of the correlation function is made and compared to the value obtained during static phase-space sampling. NEED to correct the calculation of the estimate of the second moment based on the current correlation function. If FREE_AND_METASTABLE pair state is requested, then we SHOULD divide the contributions depending on the number of turning points provided by correlation_eval. The tracking of the number of turning points is already implemented. The communication between processes is realized using MPI_Allreduce function.

MPI Function | calculate_correlation_array_and_save

Call CFncArray calculate_correlation_array_and_save(*ms, *params, Temperature);

Arguments MoleculeSystem *ms

CalcParams *params double base_temperature

Description

Function save_correlation_function

Call void save_correlation_function(*fp, cf, *params);

Arguments FILE *fp

CFnc cf

CalcParams *params

Description The correlation function values cf.data assumed to be unnormalized by the number of trajectories

cf.ntraj. The file stream fp is truncated using ftruncate before writing into it.

Function | invert_momenta

Call void invert_momenta(*ms);

Arguments MoleculeSystem *ms

Description Inverts the momenta stored inside MoleculeSystem.

4.6.2 Calculating spectral functions

struct SFnc

```
Fields double *nu
double *data
size_t len - number of samples in *nu and *data
size_t capacity - capacity of *nu and *data
size_t ntraj - number of trajectories used for averaging
double Temperature
```

MPI Function | calculate_spectral_function_using_prmu_representation

Call SFnc calculate_spectral_function_using_prmu_representation_and_save(*ms, *params,

Temperature)

Arguments MoleculeSystem *ms

CalcParams *params

double Temperature

Description

First, we check that params.ApproximateFrequencyMax is less than Nyquist frequency of the signal that will be sampled with requested params.sampling_time. Then, based on the calculated frequency step, a length of the frequency array is calculated. Since the frequency step depends on the params.sampling_time and params.MaxTrajectoryLength, the maximum frequency of the calculated spectral function will be somewhat close to the requested params.ApproximateFrequencyMax (as close as possible using the frequency step). The estimates of the zeroth and second spectral moments are obtained using params.initialMo_npoints and params.initialMo_npoints, respectively. The accumulation of params.total_trajectories is divided into params.niterations. For each trajectory the intermolecular distance is set to params.Ro. The trajectory is cut at the same distance params.Ro. Connes apodization is applied to time dependencies of Cartesian components of dipole throughout collisional trajectory before applying Fourier transform to them. The length of the dipole array which is equal to params.MaxTrajectoryLength needs to be a power of 2. The current estimate of the spectral function is saved to params.sf_filename at the end of each iteration. The zeroth and second spectral moments are obtained from the current estimate of the spectral function and compared to the values obtained through static phase-space sampling.

When requantization is enabled, the inertia tensor value is adjusted to account for the effect of centrifugal distortion. The line position of rotational transition is given by

$$\tilde{\nu}_{J' \leftarrow J''} = 2B(J'' + 1) - 4D(J'' + 1)^3 = 2B_{\text{eff}}(J'' + 1). \tag{5}$$

The line position can be expressed in terms of the effective rotational constant, B_{eff} , which depends on the rotational quantum number:

$$B_{\text{eff}} = B - 2D(J+1)^2. \tag{6}$$

To enable the adjustment of the inertia tensor, the DJ field must be set, and the MonomerType must be either LINEAR_MOLECULE_REQ_INTEGER or LINEAR_MOLECULE_REQ_HALFINTEGER.

Function save_spectral_function

Call void save_spectral_function(*fp, sf, *params);

Arguments FILE *fp

SFnc sf

CalcParams *params

Description The spectral function values sf.data assumed to be unnormalized by the number of trajectories

sf.ntraj. The file stream fp is truncated using ftruncate before writing into it.

Function connes_apodization

Call void connes_apodization(a, sampling_time);

Arguments Array a

double sampling_time

Description Multiplies the provided array by Connes apodization with time-normalization factor (a) set to

sampling_time: $A(t) = (1 - t^2/a^2)^2$. See link.

4.7 Processing results

Function read_correlation_function

Call bool read_correlation_function(*filename, *sb, *cf);

Arguments const char *filename

String_Builder *sb

CFnc *cf

Reads the correlation function from the file filename expecting a header containing metainformation (the lines should begin with '#') and data presented in the two-column format. The header is read into String_Builder without parsing. The numerical values are parsed using fscanf function and checked to be non-NaN. Returns false when either the file could no be correctly opened to a parser error was encountered.

Function read_spectral_function

Call bool read_spectral_function(*filename, *sb, *sf);

Arguments const char *filename

String_Builder *sb

SFnc *sf

Description

Reads the spectral function from the file filename expecting a header containing metainformation (the lines should begin with '#') and data presented in the two-column format. The header is read into String_Builder without parsing. The numerical values are parsed using fscanf function and checked to be non-NaN. Returns false when either the file could no be correctly opened to a parser error was encountered.

Function | writetxt

```
Call bool writetxt(*filename, *x, *y, len, *header);
Arguments const char *filename
double *x
double *y
size_t len
const char *header
```

Description

Function integrate_composite_simpson

```
Call double integrate_composite_simpson(*x, *y, len);
```

Arguments double *x

double *y
size_t len

Description Performs numerical integration using composite Simpson's 3/8 rule. See link.

$Function | \verb| compute_Mn_from_sf_using_classical_detailed_balance| \\$

```
Call double compute_Mn_from_sf_using_classical_detailed_balance(sf, n);
```

Arguments SFnc sf

 $size_t n$

Description

Calculates the n-th moment of the spectral function via numerical integration using the composite Simpson's rule <code>integrate_composite_simpson</code>. The expressions for classical detailed balance are employed. The implementation employs a straightforward application of the Simpson's rule, without invoking any special handling of lower- or higher-frequency ranges. It is assumed that the spectral function is provided solely for positive frequencies. The dimensions of the frequency and spectral function values are expected to be $\rm cm^{-1}$ and $\rm J\cdot m^6\cdot s^{-1}$, respectively. The resulting spectral moment is obtained in units of $\rm cm^{-n-1}\cdot Amagat^{-2}$.

```
Function | compute_Mn_from_sf_using_quantum_detailed_balance
```

```
Call double compute_Mn_from_sf_using_quantum_detailed_balance(sf, n);
Arguments SFnc sf
size_t n
```

Calculates the n-th moment of the spectral function via numerical integration using the composite Simpson's rule <code>integrate_composite_simpson</code>. The expressions for quantum detailed balance are employed. The implementation employs a straightforward application of the Simpson's rule, without invoking any special handling of lower- or higher-frequency ranges. It is assumed that the spectral function is provided solely for positive frequencies. The dimensions of the frequency and spectral function values are expected to be cm⁻¹ and $J \cdot m^6 \cdot s^{-1}$, respectively. The resulting spectral moment is obtained in units of cm⁻ⁿ⁻¹ · Amagat⁻².

struct WingParams

Fields double A

double B

double C

Description Stores the parameters for the Lorentzian function shifted vertically by constant value:

$$y = C + A/(1 + B^2x^2).$$

struct WingData

Fields size_t n

double *t

double *y

 ${\bf Description}$

Function wingmodel

Call double wingmodel(*wp, t);

Arguments WingParams *wp

double t

Description

Graphical interface is needed to visualize this step...

Function | fit_baseline

Call WingParams fit_baseline(*cf, EXT_RANGE_MIN);

Arguments CFnc *cf

size_t EXT_RANGE_MIN

Description

Function dct

Call double* dct(*v, len);

Arguments double *v

 $\mathtt{size_t\ len}$

Description

Implements the Fast Discrete Cosine Transform using the trick of Makhoul (2N + half-sample shift) and FFT, also known as DCT-II. For details, see original paper. A new vector with result is allocated and returned. Note that the output vector is not scaled by any time/ frequency units, making the implementation unit-agnostic.

Function | idct

Call double* idct(*v, len);

Arguments double *v

 $size_t len$

Implements the Inverse Fast Discrete Cosine Transform using the trick of Makhoul (2N + half-sample shift) and IFFT. For details, see original paper. A new vector with result is allocated and returned. Note that the output vector is not scaled by any time/ frequency units, making the implementation unit-agnostic.

```
Function | dct_sf_to_cf(SFnc sf)
 Call
              CFnc dct_sf_to_cf(cf);
 Arguments
              SFnc sf
 Description
Function | dct_cf_to_sf(CFnc cf)
 Call
              CFnc dct_cf_to_sf(cf);
 Arguments
              CFnc cf
 Description
Function dct_numeric_sf
 Call
              SFnc dct_numeric_sf(cf, *wp);
 Arguments
              CFnc cf
              WingParams *wp
 Description
Function | desymmetrize_d2
 Call
              SFnc desymmetrize_d2(sf);
 Arguments
              SFnc sf
 Description
              needs at least visual check
Function | desymmetrize_sch
 Call
              SFnc desymmetrize_sch(sf);
 Arguments
              SFnc sf
 Description
Function | desymmetrize_egf
 Call
              SFnc desymmetrize_egf(sf);
 Arguments
              SFnc sf
 Description
Function | desymmetrize_frm
 Call
              SFnc desymmetrize_frm(sf);
 Arguments
              SFnc sf
 Description
struct Spectrum
 Fields
        double *nu
         double *data
         size_t len - number of samples in *nu and *data
         size_t capacity - capacity of *nu and *data
Function | compute_alpha
```

Call Spectrum compute_alpha(sf);
Arguments SFnc sf
Description

4.8 LOESS: locally weighted polynomial regression

LOESS blends the simplicity of linear least squares regression with the adaptability of nonlinear regression. It achieves this by fitting simple model to localized subset of the data, gradually constructing a function that captures the deterministic pattern of the variation in the data – effectively filtering out the random component that follows some probability distribution.

Degree of local polynomials. The local polynomials fitted to each subset of the data are typically of either first or second degree. Employing a zero-degree polynomial reduces LOESS to a weighted moving average. While higher-degree polynomials could theoretically be used, they are not aligned with the spirit of LOESS. Such polynomials are prone to overfitting within each subset and thus often lead to numerical instability.

Weight function. The weight function, gives the most weight to the data points nearest the point of estimation and the least weight to the data points that are furthest away. The use of the weights is based on the idea that points near each other are more likely to be related to each other in a simple way than points that are further apart. The traditional weight function used for LOESS is the tricube weight function: $w(x) = (1 - |x|^3)^3$ for |x| < 1 and 0 otherwise. The main criteria for the weight function are the following (Cleveland, 1979):

- w(x) > 0 for |x| < 1 since negative weights do not make sense
- w(-x) = w(x): there is no reason to treat points to the left of x differently from those to the right
- w(x) is a nonincreasing function for $x \ge 0$: it seems unreasonable to allow a point that is closer to x to have less weight than the one that is further away
- w(x) = 0 for $|x| \geqslant 1$

In addition it seems desirable that w(x) decrease smoothly to 0 as x goes from 0 to 1. Such a weight function is more likely to produce a smoothed result. The tricube has been chosen since it enhances a χ^2 -distributional approximation of an estimate of the error variance. So it should provide an adequate smooth in many situations. The weight for a specific point in any localized subset of data is obtained by evaluating the weight function at the distance between that point and the point of estimation, after scaling the distance so that the maximum absolute distance over all of the points in the subset of data is exactly one.

Let us consider the case of local second-degree polynomials:

$$\hat{y} = \beta_0 + \beta_1 x + \beta_2 x^2$$

In this model, β_0 , β_1 , β_2 are coefficients needed to be estimated using the data pairs (x_i, y_i) within a specified window. The predicted value \hat{y} is derived from the this local polynomial model.

The matrix X is built using the x-values within the window. Each row of X corresponds to a separate x-value, while the columns represent the constant term (β_0) , the linear term (β_1) , and the quadratic term (β_2) .

$$X = \begin{bmatrix} 1 & x_1 & x_1^2 \\ 1 & x_2 & x_2^2 \\ \dots & \dots & \dots \\ 1 & x_n & x_n^2 \end{bmatrix}$$

In weighted regression, each observation is assigned a weight w_i . These weights are organized into a digonal weight matrix W, defined as:

$$W = \begin{bmatrix} w_1 & 0 & 0 & \dots \\ 0 & w_2 & 0 & \dots \\ \dots & \dots & \dots & \dots \\ 0 & 0 & 0 & w_n \end{bmatrix}$$

The normal equations for weighted regression problem are expressed as:

$$(X^T W X)\beta = X^T W y$$

To solve these equations we explore the following methods:

- complete orthogonal decomposition of X^TWX to compute its pseudo-inverse
- QR decomposition of X^TWX (no pivoting) to compute its pseudo-inverse
- robust Cholesky decomposition-based solver

For smoothing CH₄-CO₂ spectra we hit windows consisting 10,000 points in the tail.

enum WEIGHT_FUNC

Values WEIGHT_TRICUBE – defined as $(1 - |x|^3)^3$ for |x| < 1

WEIGHT_BISQUARE – defined as $(1 - |x|^2)^2$ for |x| < 1

Description Enumeration defines the types of weight functions available for assigning weights to data points.

The choice of weight function determines how influence is assigned to observations based on their

proximity to the central point at which smoothing is performed.

enum LS_METHOD

Values LS_COMPLETE_ORTHOGONAL_DECOMPOSITION – uses complete orthogonal decomposition to compute

the pseudo-inverse of the matrix (X^TWX)

LS_QR_NO_PIVOTING - fastest out of variants of QR decompositions, but maybe unstable if the

matrix is not rull rank

LS_CHOLESKY_SOLVER – is usually the fastest. However, if the matrix is even mildly ill-conditioned, this is not a good method. It loses roughly twice as many digits of accuracy using the normal

equation, compared to the more stable methods mentioned above.

Description Enumeration of numerical methods for solving linear least squares problem: $(X^TWX)beta =$

 X^TWY

Function loess_init

Call void loess_init(*x, *y, len);

Arguments double *x

double *y
size_t len

Description This function prepares the input data for LOESS by storing the predictor 'x' and response 'y'

values, and setting up the necessary values for subsequent LOESS calculations. The function assumes that the input arrays \dot{x} and \dot{y} are of equal length and contain valid numerical data.

struct Smoothing_Config

```
Fields size_t degree - degree of local polynomial [recommended: 2-3]
```

size_t ws_min - minimum window size

size_t ws_step - window size step

size_t ws_delay - optional: the index at which the window is starting to increase

size_t ws_cap - optional: cap on window size

Function | loess_estimate

Call double loess_estimate(x, window_size, degree);

Arguments double x

size_t window_size

size_t degree

Description This function computes a local polynomial fit of provided degree within a specified window around

the predictor value 'x' and uses it to estimate the response value 'y'.

```
Function loess_create_grid
```

Call double *loess_create_grid(xmin, xmax, npoints);

Arguments double xmin

double xmax
size_t npoints

Description Generates a uniformly spaced grid of predictor values within a specified range. This function creates

a grid of equally spaced 'grid_npoints' points between 'grid_xmin' and 'grid_xmax' (inclusive). The function checks the validity of the provided grid bounds with respect to the range of the previously

provided input data for smoothing.

Function loess_apply_smoothing

Call double* loess_apply_smoothing(*config);

Arguments Smoothing_Config *config

Description This function performs LOESS smoothing on the input data by fitting local polynomials to subsets

of the data and combining the results to produce a smoothed curve. The behavior of the smoothing process is controlled by the Smoothing_Config struct, which specifies parameters such as the

polynomial degree, window size, and parameters of window extension

Function loess_free

Call void loess_free();

Description Releases the memory allocated for smoothing.

4.9 Functions to transform angles between frames of reference

```
Function linear_molecule_atom_lab_to_mol
```

Call void linear_molecule_atom_lab_to_mol(*qlab, *qmol);

Arguments double *qlab

double *qmol

Description

Put here a description from dissertation

Function | linear_molecule_atom_Jacobi_mol_by_lab

Call void linear_molecule_atom_Jacobi_mol_by_lab(jac, *qlab, *qmol);

double *qlab
double *qmol

Description

Put here a description from dissertation

Function CH4_linear_molecule_lab_to_kal

Call void CH4_linear_molecule_lab_to_kal(*qlab, *qkal);

Arguments double *qlab

double *qkal

Description

Put here a description from dissertation

Function CH4_linear_molecule_Jacobi_kal_by_lab

Call void CH4_linear_molecule_Jacobi_kal_by_lab(jac, *qlab, *qkal);

Arguments Eigen::Ref<Eigen::MatrixXd> jac

double *qlab

Put here a description from dissertation

4.10 Utility structs & functions

```
struct Array
```

Fields double *data

 $size_t n$

Description Sized array in dynamic memory.

Function | create_array

Call Array create_array(n);

Arguments size_t n

Description

Function | init_array

Call void init_array(n);

Arguments Array *a

double *data

size_t n

Description

Function | free_array

Call void free_array(n);

Arguments Array *a

Description

struct String_Builder

Fields char *items - dynamically allocated buffer holding the string data

size_t count - the current number of characters in the buffer

size_t capacity - the total allocated capacity of the buffer in bytes

Description This struct represents a resizable buffer designed to build strings dynamically. It stores characters

in a contiguous block of memory, allowing for manipulation of strings.

Function sb_append

Call void sb_append(*sb, *line, n);

Arguments String_Builder *sb

const char *line

size_t n

Description Appends a sequence of characters to the String_Builder. This function appends the first 'n'

characters from the provided 'line' to the String_Builder. If 'n' exceeds the length of 'line', the behavior is undefined. The String_Builder automatically resizes its buffer if necessary to

accommodate the new characters.

Function sb_append_cstring

Call void sb_append(*sb, *line, n);

Arguments String_Builder *sb

const char *line

Description This function appends a C-style (null-terminated) string line to the String Builder's buffer.

If the String_Builder does not have sufficient capacity, its storage is automatically extended to accommodate the new content. If the String_Builder's capacity is zero, it is first resized to INIT_SB_CAPACITY bytes before any extension occurs.

Function | sb_append_format

Call void sb_append_format(*sb, *format, ...);

Arguments String_Builder *sb

const char *format

varargs(...)

Description This function takes a format string and a variable number of arguments, formats them according

to the specified format, and appends the resulting string to the provided <code>String_Builder</code>. If the <code>String_Builder</code> lacks sufficient capacity, its storage is automatically extended to accommodate the new content. If the <code>String_Builder</code>'s capacity is zero, it is first resized to <code>INIT_SB_CAPACITY</code>

bytes before any extension occurs.

Function | sb_append_seconds_as_datetime_string

Call void sb_append_seconds_as_datetime_string(*sb, s);

Arguments String_Builder *sb

int s

Description Appends a string representation of a time duration in seconds to a String Builder. The string is

formatted as a human-readable datetime string (e.g., "1h 2m 3s").

Function | sb_reset

Call void sb_reset(*sb);
Arguments String_Builder *sb

Description This function effectively clears the content of the String_Builder by setting its length to zero.

The underlying buffer is not deallocated, allowing it to be reused for subsequent operations.

Function | sb_free

Call void sb_free(*sb);
Arguments String_Builder *sb

Description This function releases the memory held by the internal buffer of the String_Builder and resets

the fields.

Supplied routines:

- 1. spherical decomposition for ab initio PES for CO₂-Ar (Kalugina/Lokshtanov)
- 2. spherical decomposition for ab initio IDS for CO₂-Ar (Kalugina/Lokshtanov)
- 3. spherical decomposition for ab initio PES for CH₄-CO₂ (Finenko)
- 4. spherical decomposition for ab initio IDS for $\mathrm{CH_4-CO_2}$ (Finenko)
- 5. spherical decomposition for ab initio PES for H₂-Ar (LeRoy/Chistikov)
- 6. spherical decomposition for ab initio PES for CO-Ar (Pederson)
- 7. spherical decomposition for ab initio IDS for CO-Ar (Rizzo)
- 8. PIP-NN representation for ab initio PES surface for N2-Ar (Finenko)
- 9. PIP-NN representation for ab initio IDS surface for N₂-Ar (Finenko)
- 10. PIP-NN representation for ab initio IDS for H₂-Ar (Meyer/Finenko)

Routines to be added in the future:

- 1. spherical decomposition for full-dimensional ab initio PES for N₂-Ar (Finenko)
- 2. spherical decomposition for long-range IDS for N₂-Ar (Wang)
- 3. spherical decomposition for long-range $d\mu/dr$ surface for N₂-Ar (Wang)
- 4. spherical decomposition for long-range IDS for H₂-Ar (Kalugina)
- 5. spherical decomposition for ab initio induced dipole for H₂-Ar (Meyer)
- 6. spherical decomposition for ab initio PES for CO₂-CO₂ (Kalugina/Lokshtanov)
- 7. spherical decomposition for ab initio IDS for CO₂-CO₂ (Kalugina/Lokshtanov)
- 8. spherical decomposition for ab initio PES for N₂-N₂ (Karman/Chistikov)
- 9. spherical decomposition for ab initio IDS for N₂-N₂ (Karman/Chistikov)
- 10. spherical decomposition for ab initio PES for N₂-H₂ (Kalugina)
- 11. spherical decomposition for long-range IDS for N_2-H_2 (Kalugina)
- 12. spherical decomposition for ab initio PES for CH₄-N₂ (Finenko)
- 13. spherical decomposition for ab initio IDS for CH₄-N₂ (Finenko)
- 14. PIP-NN representation for full-dimensional ab initio PES for CH₄-N₂ (Finenko)

5 A skeleton of the user's program

For now, let us assume that the user is supposed to use Hawaii Hybrid as a library, not via the configuration file to a driver program (which could be arranged in the future). Then the following structure is expected:

1. Initialize parallel environment

Call MPI_Init to initialize MPI if desired.

2. Initialize MoleculeSystem

Call init_ms specifying types of monomers, their tensors of inertia, the reduced mass of the molecule pair and the generator seed.

3. ...

References

- [1] D. N. Chistikov et al. "Simulation of collision-induced absorption spectra based on classical trajectories and ab initio potential and induced dipole surfaces. II. CO₂-Ar rototranslational band including true dimer contribution". In: *Journal of Chemical Physics* 155.6, 064301 (2021), p. 064301. DOI: 10.1063/5.0060779.
- [2] Piotr M. Kowalski. "Infrared absorption of dense helium and its importance in the atmospheres of cool white dwarfs". In: *Astronomy & Astrophysics* 566, L8 (June 2014), p. L8. DOI: 10.1051/0004-6361/201424242. arXiv: 1406.4591 [astro-ph.SR].

6 Changelog

- 24.12.2024 rhsMonomer: accepts pointer so that monomer's qp can be changed if apply_requantization flag is set.
- 06.01.2025 Makefile: switched to Makefile from build script.
- 08.01.2025 added MPI library which is hidden by the guard macro, implemented mpi_calculate_MO. Now the MPI and non-MPI versions of hawaii.c are compiled into two object files.
- 09.01.2025 implemented calculate_correlation_and_save.
- 09.01.2025 Porting some of hep-functionality in hep_hawaii.cpp. Zeroth moment can be calculated using adaptive Monte Carlo integration over phase space.
- 10.01.2025 Tracking the number of turning points is added to correlation_eval.
- 10.01.2025 PES and its derivatives for CH_4-CO_2 are adapted.
- 11.01.2025 fixing error inverting momenta in correlation_eval. Calculating correlation function for CO₂-Ar seems to be working correctly. Need to run a calculation with larger number of trajectories.
- 12.01.2025 added higher-order finite-difference formulas for differentiating energy: compute_numerical_rhs
- 12.01.2025 trajectory for CH₄-CO₂ using kinetic energy only works correctly. Angles transformation and its jacobian needs to be adapted from FUNCHAL to account for different order of coordinates.
- 15.01.2025 Run calculation of correlation function for $\rm CO_2-Ar$ for 10.000.000 trajectories. The correlation function is in close agreement with the previous results. Now moving on to adapting code for processing correlation functions in hawaii.
- 17.01.2025 Adapted functions related to processing correlation function from FUNCHAL. Code implementing loess algorithm is left as-is because of the use of Eigen3.
- 19.01.2025 Added checks for NaN values of dipole in correlation_eval to avoid the corruption of the correlation function estimate. At least one corrupted value of correlation function for CO₂—Ar has occurred during the calculation of 10.000.000 trajectories. The reason for the occurence of the NaN is unknown. Should investigate when initial condition leading to NaN values is found. Could this NaN be the consequence of overflow in generate_normal?
- 19.01.2025 Implemented calculate_spectral_function_using_prmu_representation_and_save function and tested that it produces correct result for CO₂—Ar at 300 K.
- 20.01.2025 Enable switch-enum option for compiler to invoke warning in the switch-cases where one of the cases of the enum is not explicitly handled even though default case is present.
- 20.01.2025 Caught a possible (but really rare) overflow error in the implementation of Box-Mueller algorithm for sampling normally distributed variable: generate_normal.
- 20.01.2025 Estimate M₀ based on the spectral function obtained during each iteration of the pr-mu calculation
- 20.01.2025 Estimate M₂ using rejection sampling and using hep. Works for CO₂-Ar
- 22.01.2025 Trying out calculating correlation function for CO-Ar. Why is the error in M_2 is 13% for 20.000.000 points? $M_0 = 1.847 \cdot 10^{-4}$, $M_2 = 4.148$. There was error in passing the arguments to dipole function (fixed). The correlation function "breaks" at some point during the calculation. Extremely large values (1e194) occur at approximately the same time intervals in the correlation function rendering it useless. Maybe it happens because of the PES, not because of IDS?
- 25.01.2025 jacobian is working for CH₄-CO₂. hawaii is extended with function to compute numerical jacobian; analytical and numerical jacobians are in close agreement.
- 25.01.2025 fixed a bug in extract_q_and_write_into_ms where coordinates for second monomer actually were overwriting the coordinates for the first monomer

- 26.01.2025 adapted dipole function for CH₄-CO₂
- 26.01.2025 examples/mpi_phase_space_integration_ch4_co2 produces M_0 and M_2 spectral moments at 300 K which are consistent with previous estimations: $M_0 = 8.29 \cdot 10^{-4}$, $M_2 = 5.37$.
- 26.01.2025 differentiate between debug/release build in Makefile: apply separate compilation flags
- 27.01.2025 examples/correlation_ch4_co2 produces what seems to be a correct spectrum, the spectral moments are in agreement with their phase-space counterparts.
- 27.01.2025 bug fix: desymmetrization procedure didn't propagate temperature to the output structure
- 31.01.2025 Implementing calculate_correlation_array_and_save that employs individual trajectory reweighting to produce results at satellite temperatures: testing on CO₂—Ar in examples/correlation_array_co2_ar
- 01.02.2025 During the correlation and pr-mu calculations for CO₂-Ar on cluster, the file with temporary result is not written (at least for several iterations at the end of which the file is supposed to be written). Turns out that if stream is flushed using fflush and filesystem caches for a given file descriptor are forced to be committed to disk using syncfs the problem is resolved.
- 01.02.2025 calculate_correlation_array_and_save produces spectral profiles for CO₂-Ar (unbound states) that are in close agreement with the results obtained in 2021. Ensemble of approximately 8 million trajectories was used. Need to check for bound states as well.
- 06.02.2025 CO-Ar 300K: $M_0 = 1.850 \cdot 10^{-4}$, $M_2 = 1.884$.
- 08.02.2025 the factor by ALU³ is moved from save_correlation_function to correlation_eval.
- 09.02.2025 added normalized flag to CFnc, changed reading/saving correlation function from/to file (not documented)
- 09.02.2025 added parsing Temperature and ntraj from header of the file using regex and added averaging of correlation functions
- 09.02.2025 rewrite of LOESS code to better understand what's going on there: prepare for OpenMP parallelization, check out different approaches to solve the weighted linear squares problem
- 09.02.2025 loess: added OpenMP parallelization of the main loop
- 11.02.2025 loess: refactored creating window & added stubs for running outside OpenMP environment
- 12.02.2025 added checking for energy conservation inside trajectory. maybe need to check this info in correlation_eval and discard the outlying trajectories?
- 12.02.2025 implemented average_correlation_functions using vararg
- 12.03.2025 calculated correlation functions for He-Ar at 50K and 300K using 500mln and 300mln trajectories, respectively. At 300K we achieved a decent looking spectrum, however at 50K it still looks not converged at large frequencies. Moreover, the correlation function at 50K exhibits pulsing (beating) oscillations which result in some spectral feature at 3.5 cm-1. What is it? Is it reproducible in quantum calculation? It is somewhat reproduced in quantum calculation. More investigation is needed. Probably should be discussed in separate paper on He-Ar calculations with spectra provided in wide temperature range.
- 16.03.2025 hawaii: use appropriate partial partition function for each temperature when calculating M_0 before proceeding to propagate trajectories
- 18.03.2025 Caught this mysterious error during bound states calculation for CH₄-CO₂: [n01p012:2488401:0:2488401] Caught signal 7 (Bus error: nonexistent physical address). No stack trace, location of error unknown.
- 23.03.2025 Added functions to perform D4 and D4a desymmetrization. Some fixes included as of 26.06 (d4b file).
- $02.05.2025 \text{ N}_2 \text{Ar PIP-NN}$ potential and dipole are added. $N_2 \text{Ar } 300\text{K}$: $M_0 = 3.833 \cdot 10^{-5}$, $M_2 = 0.2321$.

- 03.05.2025 When performing calculation of the correlation function, garbage-like values were found in the correlation accumulator. Fixed: forgot to memset before use. Also call to memset is added in calculate_correlation_array_ar
- 03.05.2025 Call to MPI_Reduce is changed to the MPI_Send/MPI_Recv pair in calculate_correlation_and_save.
- 14.05.2025 Applying requantization to H₂-Ar trajectory.
- 18.05.2025 Added some timestamps for HAWAII output and when saving spectral function to file in prmurepresentation. Added the same timestamps for correlation function and correlation function arrays.
- 19.05.2025 Added PIP-NN representation of the H_2 -Ar *ab initio* induced dipole. Checked that it produces the same values as in FUNCHAL.
- 19.05.2025 experimenting with the Poisson distribution for free path in prmu representation
- 19.05.2025 reviewed molecular constants for CO. Be careful with modifying molecular constants, since potential or dipole function may heavily rely on the very specific values. It's not the case for PES for CO-Ar but may easily be in other cases.
- 19.05.2025 MPI_AllReduce → MPI_Send/Recv for calculation using prmu representation. Need to check that after this change procedure produces correctly normalized results
- 22.05.2025 Analytic consideration of the part of the trajectory when the linear molecule rotates freely
- 24.05.2025 Receive packets out-of-order from slave processes in calculate_spectral_function_using_prmu_representation_and_
- 26.05.2025 Fixed gsl_histogram_extend_right
- 26.05.2025 Introduced Arena allocator and handled the extension of the histogram when received on the master process
- 26.05.2025 Introduced histograms of initial and final angular momenta in calculate_spectral_function_using_prmu_representation for now, they are printed in the output file (stdout stream). We need to redirect them in the corresponding files and allow to customize the path throw some variable in CalcParams.
- 27.05.2025 Fixed a bug in the logic of turning on/off requantization in calculate_spectral_function_using_prmu_representation. It was noticed when histogram of the number of requantization switches has become available. After the fix the intensity distribution of requantized lines in the rotational band of CO-Ar became more logical and resembling the HITRAN one.
- 28.05.2025 Account for the effect of centrifugal distortion when requantization is applied. The line positions of CO became in excellent agreement with those from HITRAN spectra
- 13.07.2025 fixed parsing error in negative floating-point numbers in driver
- 13.07.2025 fixed printing accumulated results in mpi_calculate_MO
- 13.07.2025 mixed spectral moments for CO-Ar at 300 K are $M_0(\text{mixed}) = 5.44 \cdot 10^{-5}$, $M_2(\text{mixed}) = 1.46 \cdot 10^{-1}$.

7 Todo's

- Investigate whether two-peak structure appears for He—Ne CIA in trajectory-based calculation using Karman PES & IDS surfaces
- Investigate three-body He-He-He CIA [2]
- When processing correlation function, add the Q-branch to the spectral function. I expect that part of the Q-branch is seen for CH₄-CO₂ after numerical FFT, but it's analytical part is missing.
- Support N₂ vibration for N₂-Ar calculation
- mpi_calculate_M0/mpi_calculate_M2: the temporary result is not accumulated over communicator, it is printed only for values accumulated only for zeroth process.

- log more information in the output file
- get rid of OpenMP magic in LOESS and instead use explicitly pthread library
- Allow providing the value (values) of zeroth and second spectral moment to calculate_correlation_and_save and calculate_correlation_array_and_save. For CH₄-CO₂ the calculation of second moments is time-consuming, so the values should probably be cached somehow. Honestly, calculating zeroth moments for 10+ temperatures is also no joke. It should be possible to cache these values as well.
- CO₂-CO₂ in 200-300K range: communicate with Wishnow about the possibility to conduct an experiment. And write a small article about trajectory-based results?
- How to organize and store the calculated results for examples? The results include CFs, SFs, spectra, spectral moments and partial partition functions.
- We should store the results of individual iterations of calculate_correlation_and_save and calculate_correlation_a
 Maybe use SQL database for this?
- Arena Allocator for storing some temporary strings and small arrays. At the moment, there are waaaay too many mallocs/frees.
- Saving the contributions of free and metastable states in correlation function separately. How can we
 pass in the information about the desired filenames? How to organize the storage for these contributions?
- The information about the number of turning points could be saved into gsl_histogram, and then partially displayed in the output file and saved to file system
- We are using MPI_Allreduce mechanism for broadcasting. This should be changed to direct MPI_Send / MPI_Recv calls. It is changed in calculate_correlation_and_save.
- Maybe we need a check that processes in the communicator get different seeds each?
- In case when correlation function is calculated for bound states, we can separate the contributions for bound-bound and bound-free transitions (Fakhardji trick)
- Adapt transforming angles between frames of reference for linear molecule-linear molecule
- How to organize Makefiles for use in different environments (several clusters)? If we push the changes to remote git repo on the remote machine, the local changes get overriden, thus removing the changes made to local Makefile. Annoying...
- I would like to make a graphical shell in which it will be possible to set parameters for calculation (for example, in the form of a list with available options for each parameter). The program is given paths to files (or drag-and-dropped) with potential and dipole functions and it assembles an executable file. Here I make an assumption that it is better to have the potential to be statically compiled with the rest of the code rather than dynamically loading it from library. It would probably be nice to be able to run the trajectory program from graphical shell locally and display the calculation result. If the correlation function is calculated, then there is no need to add its processing within the same "stage". Let the correlation function be calculated separately, then the result should be loaded and converted into a spectrum. In the case of calculating the spectral function in the pr-mu representation, smoothing is also required, it is also carried out in a separate "stage". This would enable us to have a starting point for a graphical shell that can demonstrate some of the features of the library without overcomplicating from the very beginning. Later on, we could think about establishing TCP connection between shell and the main program running remotely enabling the user to setup the calculation with the desired parameters and monitor the calculation.

8 Notes

Usage of GNU profiler (gprof):

- 1. Compile the program with the -pg flag: g++ -pg -o test test.cpp
- 2. Run the program: ./test
- 3. Generate the profiling report: gprof test gmon.out > analysis.txt