ETO-HEOM Tutorial

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Quick Start: Workflow Summary

This section provides a high-level summary of the full workflow. Follow these steps to run a full 2DES simulation using ETO-HEOM:

1. Build the Solver Compile the CPU and/or GPU solvers:

```
\begin{array}{ll} \mathbf{cd} & /\operatorname{path}/\operatorname{to}/\operatorname{ETO-\!HEOM}/\\ \mathrm{make} & \end{array}
```

2. Initialize Working Directory Use the setup script to create a simulation workspace:

```
./setup_cpu_job.sh JOBNAME BATHTYPE
# or
./setup_gpu_job.sh JOBNAME BATHTYPE
```

- 3. Edit Template Input File Modify key.key-tmpl to set up:
 - System size, HEOM level, Hamiltonian, Disorder
 - Dipole directions, pulse settings, and time domain
- 4. Generate Bath Parameters Automatically fill in the bath section by executing:

```
\begin{array}{ll} {\rm python 3~BATHTYPE.ETOM.\,py} \\ \# \ or \ run \ BATHTYPE\_ETOM\_example.\,ipynb \end{array}
```

5. Generate Input Files Configure and run the input generation script:

```
./gen_2d_data.sh
```

6. Submit PBS Jobs Submit the simulation jobs to the cluster:

```
./submit_jobs.sh
```

- Wait Until Completion Monitor jobs with qstat or log files. Ensure all output files are generated in ./2d-output.
- 8. Analyze Results and Generate 2D Spectrum After all jobs finish, analyze the results:

```
python3 gen_2d_spectrum.py
# or run gen_2d_spectrum_example.ipynb
```

For detailed instructions, continue reading the full tutorial below.

1 Build Instructions

Before setting up any jobs, compile the solvers by running make in the project root:

```
\begin{array}{ll} \mathbf{cd} & /\operatorname{path}/\operatorname{to}/\operatorname{ETO-\!HEOM}/\\ \mathrm{make} & \end{array}
```

This builds the following binaries:

- CPU_2DES CPU-based HEOM 2DES simulator
- GPU_2DES GPU-accelerated HEOM 2DES simulator (if applicable)

Ensure the binaries are placed in their respective src/ directories. For example:

- etoheom-cpu/src/
- etoheom-gpu/src/

2 Job Setup Scripts

Before using any setup scripts, ensure that the HOME_PATH variable in the script files is set to the current path of your ETO-HEOM project.

2.1 CPU Job Setup

Initializes a simulation folder for CPU-based 2DES simulations.

Usage

```
./setup_cpu_job.sh JOBNAME BATHTYPE
```

Arguments

- JOBNAME: Custom identifier for the simulation (e.g., dimer)
- BATHTYPE: Type of spectral density must be one of:
 - $\ {\tt debye_lorentz}$
 - ohmic
 - superohmic

Resulting Structure

```
JOBNAME_BATHTYPE/
|-- 2d-input/
                                 # Input .key files
|-- 2d-output/
                                 # Output .out files
|-- pbs-script/
                                 # PBS scripts
   |-- pbserr/
                                 # PBS error logs
   |-- pbslog/
                                 # PBS output logs
|-- key.key-tmpl
                                 # Input template file
|-- gen_2d_data.sh
                                 # Script to generate .key files
|-- clean_2d_data.sh
                                 # Script to clean .key and .out files
|-- gen_2d_spectrum.py
                                 # Script to gen 2d spectrum
|-- gen_2d_spectrum_example.py
                                 # gen 2d spectrum example
|-- BATHTYPE_ETOM.py
                                 # ETO model for the specified bath type
|-- BATHTYPE_ETOM_example.ipynb  # ETO model example
|-- submit_jobs.sh
                                 # Script to generate & submit PBS jobs (CPU)
|-- README.md
                                 # Usage of working file
```

2.2 GPU Job Setup

Initializes a simulation folder for GPU-based 2DES simulations.

Usage

```
./setup_gpu_job.sh JOBNAME BATHTYPE
```

Arguments

- JOBNAME: Custom identifier for the simulation (e.g., dimer)
- BATHTYPE: Type of spectral density must be one of:
 - debye_lorentz
 - ohmic
 - superohmic

Resulting Structure

```
JOBNAME_BATHTYPE/
|-- 2d-input/
                                # Input .key files
|-- 2d-output/
                                # Output .out files
|-- pbs-script/
                                # PBS scripts
  |-- pbserr/
                               # PBS error logs
   |-- pbslog/
                               # PBS output logs
|-- key.key-tmpl
                               # Input template file
|-- gen_2d_data.sh
                               # Script to generate .key files
|-- clean_2d_data.sh
                               # Script to clean .key and .out files
|-- gen_2d_spectrum.py
                                # Script to gen 2d spectrum
|-- gen_2d_spectrum_example.py # gen 2d spectrum example
|-- BATHTYPE_ETOM.py
                                # ETO model for the specified bath type
|-- BATHTYPE_ETOM_example.ipynb  # ETO model example
|-- submit_jobs.sh
                                # Script to generate & submit PBS jobs (GPU)
|-- README.md
                                # Usage of working file
```

3 Template Input Setup

After setting up your job folder, you must modify the key.key-tmpl file located in the working directory. This is the template input file that defines all system-specific parameters for the simulation.

Update the following sections to reflect your system configuration:

SIZE

Define the system size including the ground and first excited states.

HEOM

Define the HEOM configuration:

• Format: (SITE_NUMBER) (TRUNCATION_LEVEL)

HAMILTONIAN

Define the system Hamiltonian, including the ground and excited states. All values are in cm⁻¹.

DISORDER

Static disorder matrix of the Hamiltonian, also in cm⁻¹. First row: number of samples and random seed.

BATH

Bath information, automatically generated by {BATHTYPE}_ETOM.py. You typically do not need to modify this section.

DIPOLE

- First row: Number of transition dipole vectors, usually same as (SITE_NUMBER)
- Following rows: Dipole direction and amplitude in XYZ components

POLARIZATION

Define the polarization angles for the four pulses. You typically do not need to modify this section.

PULSE

- First row: Number of pulses (currently only supports 3)
- Next 3 rows: Amplitude ($< 10 \text{ cm}^{-1}$), central time (fs), width (fs), and frequency (cm⁻¹)
- Use placeholders TAU1, TAU2, TAU3 that will be replaced automatically by gen_2d_data.sh

TIME

Simulation time settings:

- Format: start time, end time (T_END), time step, and sampling interval
- T_END will be replaced dynamically during gen_2d_data.sh execution

4 Input File Generation Script

Modify gen_2d_data.sh

This shell script controls how the 2DES input files are generated. You may adjust the following key parameters at the top of the script:

- t0=210 Initial central time for the first pulse (in fs)
- propagate_time=600 Duration to propagate after the last pulse (in fs)
- tau_step=10 Step size for τ_2 (in fs)
- tau_bound=600 Upper and lower bounds for τ_2 offset (in fs)
- T=(0) Array of population times T (in fs). You can add more values.
- input_file="key.key-tmpl" Path to the input template file

Script Functionality

The script will:

- Loop over values of τ_2 from -tau_bound to +tau_bound
- Automatically compute TAU1, TAU2, TAU3, and T_END based on t0, tau_step, and each value in T
- Replace placeholders in the template file with computed values
- Output input files as: ./2d-input/key_{tau2}_{T}.key

You can modify this script to include more population times or to adjust the pulse timing logic as needed.

Run gen_2d_data.sh

Once the script is configured, execute it to generate input files:

```
./gen_2d_data.sh
```

The generated input files will be saved in the ./2d-input directory.

Clean Input Files: clean_2d_data.sh

To remove all generated input files and reset the input directory, use:

```
./ clean_2d_data.sh
```

This will delete all files inside the ./2d-input/ folder.

5 Generate Bath Parameters

After the working directory is created using the setup script, the bath type (BATHTYPE) is already specified and embedded into the directory structure. You do not need to manually modify or set the bath type again.

Execute BATHTYPE_ETOM.py

To generate bath parameters for your simulation, run the corresponding Python script inside your working directory. This script will automatically insert bath information into key.key-tmpl:

```
python3 debye_lorentz_ETOM.py
```

This command calculates the ETO model parameters based on the specified spectral density type and updates the BATH section in the template file accordingly.

Use Jupyter Notebook: BATHTYPE_ETOM_example.ipynb

You may also use the accompanying Jupyter notebook to interactively generate bath parameters:

• Launch the notebook with:

```
jupyter notebook debye_lorentz_ETOM_example.ipynb
```

• Execute the notebook cells step-by-step to calculate and visualize the bath parameters.

This method is recommended if you wish to explore or fine-tune the bath model before applying it.

6 Submit PBS Jobs

After generating all input files, execute the submission script to generate and submit PBS job scripts.

$Run submit_jobs.sh$

From your working directory, run:

```
./submit_jobs.sh
```

This script will automatically:

- Create PBS job scripts for the CPU or GPU version based on your setup
- Divide the τ_2 range into manageable chunks (CPU only)
- Submit the generated PBS scripts using qsub

Important Parameters to Check

Before execution, ensure the following variables in submit_jobs.sh are properly set:

- HOME_PATH Automatically set by the setup script
- JOBNAME Automatically set by the setup script
- CPU_2DES or GPU_2DES Path to your binary executable
- \bullet INPUT_DIR Should be ./2d-input
- OUTPUT_DIR Should be ./2d-output
- ERR_DIR, LOG_DIR Should point to subfolders inside ./pbs-script/
- TLIST Array of population times T (e.g., 0 or (0 100 200))
- START_TAU, END_TAU, STEP_TAU Define the au_2 scan range

PBS Script Generation: CPU Version

For CPU-based simulations, the script:

- Splits the τ_2 scan range into equal parts (NUM_SCRIPTS)
- Generates one PBS script per chunk under pbs-script/
- Submits them using qsub

Each PBS job:

- Executes up to 11 parallel CPU processes (ppn=11)
- Launches CPU_2DES for each key_{TAU}_{T}.key input
- Outputs result to 2d-output/out_{TAU}_{T}.out

PBS Script Generation: GPU Version

For GPU-based simulations, the script:

- Generates a single PBS job script under pbs-script/
- Sets GPU-specific resources (e.g., nodes=gpu02, CUDA environment variables)

The job:

- Executes GPU_2DES for each input file
- Logs results and errors in pbs-script/pbslog and pbs-script/pbserr

Output

Submitted PBS jobs will automatically:

- Read input files from ./2d-input
- Write output files to ./2d-output
- Write logs to ./pbs-script/pbslog/
- Write errors to ./pbs-script/pbserr/

Use qstat, tail -f, or PBS web portal to monitor job progress.

7 Post-Processing: Generate 2D Spectrum

After all PBS jobs have completed and the output files are available in the ./2d-output directory, you can generate the 2D spectrum by running the analysis script.

Execute gen_2d_spectrum.py

This script will parse the output files in 2d-output/, perform Fourier transforms over delay times, and assemble the final 2D spectrum data.

To run the script:

```
python3 gen_2d_spectrum.py
```

The resulting 2D spectrum will be saved in a format specified within the script (e.g., NumPy arrays, figures, or HDF5 files).

Use Jupyter Notebook: gen_2d_spectrum_example.ipynb

Alternatively, if you prefer interactive analysis or visualization, you can use the accompanying Jupyter notebook:

• Launch the notebook with:

```
jupyter notebook gen_2d_spectrum_example.ipynb
```

- Follow the instructions and code cells to:
 - Load output data
 - Apply any windowing or filtering
 - $-\,$ Compute the 2D Fourier transform
 - Plot the 2D spectrum using Matplotlib

This notebook is useful for adjusting analysis parameters and visually verifying results.