

MixPI: User Guide

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II. DOWNLOADING AND COMPILING MIXPI

A. Download

1. *MixPI*

The most recent version of MixPI is v1.0. Subsequent versions will be developed and updated to this page. The following subdirectories are located within the MixPI download:

- `src/` : Contains the source codes for MixPI.
- `examples/` : Example input files for a MTS bulk water simulation and a $\text{Co}^{3+} + e^-$ simulation
- `updated-cp2k-files/` : The cp2k source files that were updated to interface with MixPI

All files in the `src/` folder should be downloaded. Files that contain the changes to CP2K are also located in the `updated-cp2k-files/` and need to be downloaded as well.

2. *CP2K*

MixPI requires the installation of the molecular dynamics package CP2K v2023.2¹ which can be found on the CP2K website. CP2K must be installed as both an executable and library. The instructions for each are also located on the CP2K website. A few changes to

the CP2K source code were made in the development of MixPI. These changes were made to the following CP2K files:

- `constraint.F`
- `constraint_vsite.F`
- `start/libcp2k.F`
- `subsys/molecule_kind_types.F`
- `topology_connectivity_util.F`
- `topology_coordinate_util.F`
- `topology_generate_util.F`
- `topology_psf.F`
- `topology_types.F`
- `topology_util.F`

Unless indicated, these files are located in the `src/` directory. The updated versions of these files are located in the MixPI directory `updated-cp2k-files/`. These changes do not interfere with running CP2K as normal.

B. Compile

1. *CP2K*

After replacing the updated files in the `src/` directory; the authors suggestion using the toolchain build to generate arch files and compile CP2K. The instructions below will assume the toolchain was used. Instructions for the toolchain build are found in the CP2K documentation. To build the CP2K version used by the authors, the following steps were used.

- From the `cp2k/` directory, access the toolchain and run the install command. `cd /tools/toolchain/`

`./install_cp2k_toolchain.sh`

For additional features to be linked to CP2K (like PLUMED), include those in the install line. (For example `./install_cp2k_toolchain.sh - -with-plumed`)

- After build, copy the arch files generated in `install/arch` to the `cp2k arch` directory
`cp install/arch/local* ../../arch/`

- Source the locations of the libraries that are linked to CP2K
`source install/setup`

- Return to the `cp2k` directory and compile.

`cd ../../`

`make -j 12 ARCH=local VERSION='ssmp sdbg ssmp pdbg'`

- After successful compilation, compile `cp2k` as library.

`make -j 12 ARCH=local VERSION='ssmp sdbg ssmp pdbg' libcp2k`

2. *MixPI*

After compiling CP2K as an executable and library, the paths in the makefile for MixPI must be updated to find the correct libraries. This is done with three separate updates:

- Copy all libraries flags from the CP2K arch file (The arch file is `arch/local.psm` and the `-l***` flags are located following `"LIBS="`.) onto the `"LIB ="` line in the makefile located in the `src/` directory of MixPI. (To find these libraries, updated the `-I` and `-L` paths within the makefile or, if compiled with the toolchain build, source the same setup file as used when building `cp2k`.)

- Update the locations of the `cp2k/dbcsr` libraries and object files. They are located at:

`CP2K_LIB = /**cp2k-directory**/lib/local/psmp`

`DBSCR_LIB = /**cp2k-directory**/lib/local/psmp/objs/dbcsr`

```
CP2K_INC = /**cp2k-directory**/obj/local/psmp
DBSCR_INC = /**cp2k-directory**/obj/local/psmp/exts/dbcsr
```

This should generate an executable (cp2k_pimd.out) which will run MixPI.

III. INPUT

In this section, we introduce the specific input files needed to run the MixPI software program. Small sections of input from a MTS water example (box of water with 4 bead H atoms and 2 bead O atoms; an aqueous Co^{3+} ion with an explicit electron treated as 1024 bead ring polymer) are shown for clarity. A full set of example input files are located in the example directory on the MixPI GitHub page.

A. MixPI Input Files: Path-Integral Variables

A file, called `rp_molecule_info.inp`, contains information about which molecules contain atoms treated as path integrals and the number of beads for each of those atoms. The structure is as follows:

```
The number of total molecules in the simulation that contain at least one PI atom
***The following block is repeated for as many molecule types as are present ***
Number of atoms in the molecule number of molecules of this type PSF file that describes
this molecule type
    Number of beads for atom 1 in molecule
    Number of beads for atom 2 in molecule
    ...
```

In Fig. 1, we show the `rp_molecule_info.inp` file for a bulk water simulation with 90 water molecules where the O atoms are treated as 4 bead RPs and the H atoms are treated as 8 bead RPs. This file will be used by the `sample make.input.F` program (which will convert classical input files to the RP input files) and the MixPI program itself.

```

90      90      SPCFw.psf      !total number of molecules with PI
3      2      !number of atoms in molecule type 1      number of molecules of type 1      the PSF file
      4      !number of beads on atom 1, molecule 1
      4      !number of beads on atom 2, molecule 1
      4      !number of beads on atom 3, molecule 1 ...

```

FIG. 1: The `rp_molecule_info.inp` file for a simulation of 90 water molecules described by the `SPCFw.psf` file with 4 bead RPs for the oxygen atoms and 8 bead RPs for the hydrogen atoms.

B. CP2K Input Files

A set of CP2K input files, similar to those needed for a traditional CP2K MD simulation, are required. While the input structures are close to those used by classical MD, the MixPI CP2K input does need to be modified to reflect the new force and structure information for the path-integral set-ups. A simple program (called `make_input.F` which is located within the `examples/generate-input` directory) will take the classical inputs and convert them into MixPI inputs. Each of the input files is outlined in a section below.

1. *cp2k.inp*

The CP2K input only needs those input parameters dedicated to the force evaluations; any parameters concerning the molecular dynamics steps (like temperature, time step, etc.) are read in from a different file. The force definitions used in `cp2k.inp` must describe the interactions between the classical atoms and RP beads, the RP beads on separate atoms, and the classical atom interactions. For example, a Lennard Jones interaction of the form

$$V_{\text{LJ}} = 4\epsilon_{\text{OO}} \left[\left(\frac{\sigma_{\text{OO}}}{r_{\text{OO}}} \right)^{12} - \left(\frac{\sigma_{\text{OO}}}{r_{\text{OO}}} \right)^6 \right] \quad (1)$$

which describes the interactions between the oxygen atoms of two water molecules has the form shown in Fig. 2. As indicated, the overall strength of the interaction (ϵ_{OO}) is divided by the number of beads ($N = 2$) to account for the ring polymer bead interactions.

A similar force scaling is also done for the bonded interactions as well as the electrostatics.

```

&LENNARD-JONES
  ATOMS OW      OW
  EPSILON [K_e]      78.265400000000000
  SIGMA [angstrom]   3.1654920000000000
  RCUT [angstrom]    7.000000000000000
&END LENNARD-JONES

```

(a) The Lennard Jones parameters between two oxygen atoms in a classical simulation using the qSPC/E force-field.

```

&LENNARD-JONES
  ATOMS 2OW      2OW
  EPSILON [K_e]      39.132700000000000
  SIGMA [angstrom]   3.1654920000000000
  RCUT [angstrom]    7.000000000000000
&END LENNARD-JONES

```

(b) The Lennard Jones parameters between the beads of two oxygen atoms that are represented by 2 bead ring polymers using the qSPC/E force-field.

FIG. 2: The CP2K nonbonded force section between the oxygen atoms in a water simulation for the (a) classical and (b) ring polymer simulations.

2. *structure.psf*

A PSF file (protein structure file) is used to indicate the atom types and bonded information for the system. A psf file for "classical" qSPC/E water is shown in Fig. 3. The first block contains the atom information. The order of the columns is: atom ID, segment name, residue ID, residue name, atom name, atom type, charge, mass, and an unused 0. As we will see below, the important columns are segment name, atom name, atom type (which, when you compare to Fig. 2, matches the force-field information), charge, and mass.

IV. THEORY

In this section, we review the theory behind PIMD and MTS-PIMD; similar derivations have been presented previously^{2,3} and are also shown in the code.

```

PSF EXT

1 !SPCFw
CP2K PSF File for qSPC/E water molecule

3 !NATOM
1 MOL1      1      SOL      OW      OW      -0.840000      15.999      0
2 MOL1      1      SOL      HW      HW      0.420000      1.008      0
3 MOL1      1      SOL      HW      HW      0.420000      1.008      0

2 !NBOND
1          2          1          3

1 !NTHETA
2          1          3

0 !NPHI

0 !NIMPHI

0 !NDON

0 !NACC

0 !NNB

```

FIG. 3: The PSF file for qSPC/E water.

1. All Replica Ring Polymer Hamiltonian

The quantum mechanical canonical partition function can be written as

$$\begin{aligned}
Z &= \text{Tr}[e^{-\beta\hat{H}}] \\
&= \int dq \langle q | e^{-\beta\hat{H}} | q \rangle,
\end{aligned} \tag{2}$$

where $|q\rangle$ is a position eigenstate of the one-dimensional Hamiltonian

$$\hat{H} = T(\hat{p}) + V(\hat{q}). \tag{3}$$

We use the asymmetric Trotter approximation to split the Hamiltonian into a potential and kinetic components to rewrite the partition function as

$$Z = \lim_{N \rightarrow \infty} \int dq \langle q | \left(e^{-\frac{\beta}{N}V(\{\hat{q}\})} e^{-\frac{\beta}{N}T(\{\hat{p}\})} \right)^N | q \rangle, \tag{4}$$

and insert N-1 copies of the identity in the position basis ($\hat{I} = \int dq |q\rangle \langle q|$) between the two components. We evaluate the potential

$$Z = \lim_{N \rightarrow \infty} \int d\{\mathbf{q}\} \prod_{\alpha=1}^N \langle q_{\alpha} | \left(e^{-\frac{\beta}{N}V(\{\hat{q}\})} e^{-\frac{\beta}{N}T(\{\hat{p}\})} \right) | q_{\alpha+1} \rangle, \tag{5}$$

where $q_{N+1} = q_1$ to maintain the trace.

The potential and kinetic terms are evaluated with respect to the position basis.

$$Z = \lim_{N \rightarrow \infty} \left(\frac{mN}{2\pi\beta\hbar^2} \right)^{\frac{N}{2}} \int d\{\mathbf{q}\} e^{-\beta \sum_{\alpha=1}^N \left(\frac{1}{N} V(q_\alpha) - \frac{Nm}{2\beta^2\hbar^2} (q_\alpha - q_{\alpha+1})^2 \right)} \quad (6)$$

By introducing identity via a Gaussian integral over momentum divided by itself, we can rewrite the expression above as

$$Z = \lim_{N \rightarrow \infty} \left(\frac{\beta}{2\pi m_f} \right)^{1/2} \left(\frac{mN}{2\pi\beta\hbar^2} \right)^{\frac{N}{2}} \int d\{\mathbf{q}\} e^{-\beta \sum_{\alpha=1}^N \left(\frac{p_\alpha^2}{2m_f} + \frac{1}{N} V(q_\alpha) - \frac{Nm}{2\beta^2\hbar^2} (q_\alpha - q_{\alpha+1})^2 \right)} \quad (7)$$

We now have an expression in which the exponential looks similar to a classical Hamiltonian of the form

$$H_{\text{RP}} = \sum_{\alpha=1}^N \frac{p_\alpha^2}{2m_f} + \frac{1}{N} V(q_\alpha) + \frac{Nm}{2\beta^2\hbar^2} (q_\alpha - q_{\alpha+1})^2 \quad (8)$$

which is referred to as the one-dimensional PIMD Hamiltonian. For equilibrium properties, one can choose any quantity as m_f , the fictitious mass. As discussed later, this choice of m_f can improve the sampling efficiency of PIMD. This Hamiltonian can be easily expanded to multiple degrees of freedom.

A. MTS-Expansion in Three-Dimensions

Using a similar methodology, we derive the MTS-PIMD Hamiltonian below. We choose to derive the Hamiltonian for three particles in one-dimension to highlight the form of the Hamiltonian used with multiple quantization levels that the MTS-PIMD Hamiltonian can accommodate. This Hamiltonian can easily be extended to handle an arbitrary number of degrees. The quantum Hamiltonian operator for three particles is written

$$\begin{aligned} \hat{H} &= \sum_{i=1}^3 \frac{\hat{p}_i}{2m_i} + \sum_{i=1}^3 V_1(\hat{q}_i) + \sum_{i=1}^3 \sum_{j<i} V_2(\hat{q}_i, \hat{q}_j) + \sum_{i=1}^3 \sum_{j<i} \sum_{k<j} V_3(\hat{q}_i, \hat{q}_j, \hat{q}_k) \\ &= H_1(\hat{q}_1, \hat{q}_2, \hat{q}_3) + H_2(\hat{q}_2, \hat{q}_3) + H_3(\hat{q}_3) \end{aligned} \quad (9)$$

where \hat{p}_i , \hat{q}_j , and m_i are the momentum, position, and mass of the i^{th} quantum particle. $V_1(\hat{q}_i)$ is the uncoupled portion of the potential energy for the i^{th} degree while $V_2(\hat{q}_i, \hat{q}_j)$ and $V_3(\hat{q}_i, \hat{q}_j, \hat{q}_k)$ are the two-body and three-body coupled potential energy contributions, respectively. We assume that the q_1 degree of freedom experiences the largest quantum mechanical effect, followed by q_2 and then q_3 ; therefore, q_1 requires more time slices, and a larger N

value, than q_2 . We define N_1 , N_2 , and N_3 as the number of time slices along each degree with $N_1 \geq N_2 \geq N_3$. We split the Hamiltonian by dependence on particle number where any two- or three-body terms are included in the Hamiltonian corresponding to the highest quantized variable. For example, $H_1(\hat{q}_1, \hat{q}_2, \hat{q}_3) = T_1(\hat{p}_1) + V(\hat{q}_1) + V(\hat{q}_1, \hat{q}_2) + V(\hat{q}_1, \hat{q}_3) + V(\hat{q}_1, \hat{q}_2, \hat{q}_3)$. As describe in Sec. IV 1, we use the asymmetric Trotter approximation on this partitioned Hamiltonian to first split H_3 from H_{12}

$$Z = \lim_{N_3 \rightarrow \infty} \int d\{q\} \langle q_1 q_2 q_3 | \left(e^{-\frac{\beta H_3}{N_3}} e^{-\frac{\beta H_{12}}{N_3}} \right)^{N_3} | q_1 q_2 q_3 \rangle \quad (10)$$

and then apply the Trotter approximation twice more to $H_{12} = H_1 + H_2$ and, finally, to H_1 to obtain

$$Z = \lim_{N_3, N_2, N_1 \rightarrow \infty} \int d\{q\} \langle q_1 q_2 q_3 | \left(\Lambda_3 (\Lambda_{21})^{\frac{N_2}{N_3}} \right)^{N_3} | q_1 q_2 q_3 \rangle. \quad (11)$$

where we define $\Lambda_i = e^{-\frac{\beta H_i}{N_i}}$ and $\Lambda_{ij} = \Lambda_i \Lambda_j^{\frac{N_j}{N_i}}$. Following a similar series of steps as employed in the all-replica derivation in the previous section, we insert N_3 copies of identity with respect to the position basis q_3 and evaluate Λ_{12} since it is diagonal with respect to q_3 .

$$Z = \lim_{N_3, N_2, N_1 \rightarrow \infty} \int d\{q\} \langle q_1 q_2 | \prod_{\alpha=1}^{N_3} \langle q_{3,\alpha} | \Lambda_3 | q_{3,\alpha+1} \rangle \Lambda_{12}(q_{3,\alpha+1})^{\frac{N_2}{N_3}} | q_1 q_2 \rangle. \quad (12)$$

We now have the one-particle results with H_3 and can use the result of the previous derivation to obtain

$$\begin{aligned} Z = \lim_{N_3, N_2, N_1 \rightarrow \infty} & \left(\frac{\beta}{2\pi m_{3,f}} \right)^{1/2} \left(\frac{m_3 N_3}{2\pi \beta \hbar^2} \right)^{\frac{N_3}{2}} \\ & \int d\{q\} e^{-\beta \sum_{\alpha=1}^{N_3} \left(\frac{p_{3,\alpha}^2}{2m_{3,f}} + \frac{1}{N_3} V_1(q_{3,\alpha}) - \frac{N_3 m_3}{2\beta^2 \hbar^2} (q_{3,\alpha} - q_{3,\alpha+1})^2 \right)} \\ & \langle q_1 q_2 | \prod_{\alpha=1}^{N_3} \Lambda_{12}(q_{3,\alpha+1})^{\frac{N_2}{N_3}} | q_1 q_2 \rangle. \end{aligned} \quad (13)$$

Next, we introduce N_3 copies of identity with respect to q_2 between each of the $\Lambda(q_{3,\alpha+1})^{N_2/N_3}$ terms

$$\begin{aligned} Z = \lim_{N_3, N_2, N_1 \rightarrow \infty} & A_3 \int d\{q\} e^{-\beta H_{3,RP}(q_3)} \\ & \langle q_1 | \prod_{\alpha=1}^{N_3} \langle q_{2,(\alpha-1)*N_2/N_3+1} | \Lambda_{12}(q_{3,\alpha+1})^{\frac{N_2}{N_3}} | q_{2,\alpha*N_2/N_3+1} \rangle | q_1 \rangle. \end{aligned} \quad (14)$$

Since q_2 is more quantized than q_3 , and needs a greater number of slices than q_3 , we introduce additional N_2/N_3 copies of identity between each $\Lambda_{12}(q_{3,\alpha+1})$

$$Z = \lim_{N_3, N_2, N_1 \rightarrow \infty} A_3 \int d\{q\} e^{-\beta H_{3,RP}(q_3)} \langle q_1 | \prod_{\alpha=1}^{N_3} \prod_{\gamma=1}^{N_2/N_3} \langle q_{2,(\alpha-1)*N_2/N_3+\gamma} | \Lambda_2(q_{3,\alpha+1}) \Lambda_1(q_{3,\alpha+1})^{\frac{N_1}{N_2}} | q_{2,(\alpha-1)*N_2/N_3+\gamma+1} \rangle | q_1 \rangle . \quad (15)$$

$\Lambda_1(q_{3,\alpha+1})$ is diagonal with respect to q_2 and we have obtained the one-dimensional ring-polymer solution for q_2 .

$$Z = \lim_{N_3, N_2, N_1 \rightarrow \infty} A_3 A_2 \int d\{q\} e^{-\beta H_{3,RP}(\{q_3\})} e^{-\beta H_{2,RP}(\{q_2\}, \{q_3\})} \prod_{\alpha=1}^{N_3} \prod_{\gamma=1}^{N_2/N_3} \langle q_1 | \Lambda_1(q_{3,\alpha+1}, q_{2,(\alpha-1)*N_2/N_3+\gamma+1})^{\frac{N_1}{N_2}} | q_1 \rangle . \quad (16)$$

We follow the same procedure to introduce N_2 copies of identity between each $\Lambda_1(q_{3,\alpha+1}, q_{2,(\alpha-1)*N_2/N_3+\gamma+1})^{\frac{N_1}{N_2}}$ and N_1/N_2 copies of identity between each $\Lambda_1(q_{3,\alpha+1}, q_{2,(\alpha-1)*N_2/N_3+\gamma+1})$ to obtain the all-replica Hamiltonian for q_1 . Once again, we introduce N_2 copies between each of the copies of $\Lambda_1(q_{3,\alpha+1}, q_{2,(\alpha-1)*N_2/N_3+\gamma+1})^{\frac{N_1}{N_2}}$

$$Z = \lim_{N_3, N_2, N_1 \rightarrow \infty} A_3 A_2 \int d\{q\} e^{-\beta H_{3,RP}(\{q_3\})} e^{-\beta H_{2,RP}(\{q_2\}, \{q_3\})} \prod_{\alpha=1}^{N_3} \prod_{\gamma=1}^{N_2/N_3} \langle q_1, ((\alpha-1)*N_2/N_3+\gamma-1)*N_1/N_2+1 | \Lambda_1(q_{3,\alpha+1}, q_{2,(\alpha-1)*N_2/N_3+\gamma+1})^{\frac{N_1}{N_2}} | q_1, ((\alpha-1)*N_2/N_3+\gamma)*N_1/N_2+1 \rangle . \quad (17)$$

as well as N_1/N_2 copies of identity in q_1 between each $\Lambda_1(q_{3,\alpha+1}, q_{2,(\alpha-1)*N_2/N_3+\gamma+1})$

$$Z = \lim_{N_3, N_2, N_1 \rightarrow \infty} A_3 A_2 \int d\{q\} e^{-\beta H_{3,RP}(\{q_3\})} e^{-\beta H_{2,RP}(\{q_2\}, \{q_3\})} \prod_{\alpha=1}^{N_3} \prod_{\gamma=1}^{N_2/N_3} \prod_{\lambda=1}^{N_1/N_2} \langle q_1, ((\alpha-1)*N_2/N_3+\gamma-1)*N_1/N_2+\lambda | \Lambda_1(q_{3,\alpha+1}, q_{2,(\alpha-1)*N_2/N_3+\gamma+1}) | q_1, ((\alpha-1)*N_2/N_3+\gamma-1)*N_1/N_2+\lambda+1 \rangle . \quad (18)$$

We obtain an expression for the three-level mixed-time slicing RPMD partition function and Hamiltonian

$$Z = \lim_{N_3, N_2, N_1 \rightarrow \infty} A_3 A_2 A_1 \int d\{q\} e^{-\beta H_{3,RP}(\{q_3\})} e^{-\beta H_{2,RP}(\{q_2\}, \{q_3\})} e^{-\beta H_{1,RP}(\{q_1\}, \{q_2\}, \{q_3\})} . \quad (19)$$

where

$$\begin{aligned}
H_{3,RP} &= \sum_{\alpha=1}^{N_3} \frac{p_{3,\alpha}^2}{2m_{3,f}} + \frac{1}{N_3} V_1(q_{3,\alpha}) - \frac{N_3 m_3}{2\beta^2 \hbar^2} (q_{3,\alpha} - q_{3,\alpha+1})^2 \\
H_{2,RP} &= \sum_{\alpha=1}^{N_3} \sum_{\gamma=1}^{N_2/N_3} \frac{p_{2,(\alpha-1)*N_2/N_3+\gamma}^2}{2m_2} + \frac{1}{N_2} V_1(q_{2,(\alpha-1)*N_2/N_3+\gamma}) + \frac{1}{N_2} V_2(q_{2,(\alpha-1)*N_2/N_3+\gamma}, q_{3,\alpha}) \\
&\quad + \frac{N_2 m_2}{2\beta^2 \hbar^2} (q_{2,(\alpha-1)*N_2/N_3+\gamma} - q_{2,(\alpha-1)*N_2/N_3+\gamma+1})^2 \\
H_{1,RP} &= \sum_{\alpha=1}^{N_3} \sum_{\gamma=1}^{N_2/N_3} \sum_{\lambda=1}^{N_1/N_2} \frac{p_{1,((\alpha-1)*N_2/N_3+\gamma-1)*N_1/N_2+\lambda}^2}{2m_1} + \frac{1}{N_1} V_1(q_{1,((\alpha-1)*N_2/N_3+\gamma-1)*N_1/N_2+\lambda}) \\
&\quad + \frac{1}{N_1} V_2(q_{1,((\alpha-1)*N_2/N_3+\gamma-1)*N_1/N_2+\lambda}, q_{2,(\alpha-1)*N_2/N_3+\gamma}) \\
&\quad + \frac{1}{N_1} V_2(q_{1,((\alpha-1)*N_2/N_3+\gamma-1)*N_1/N_2+\lambda}, q_{3,\alpha}) \\
&\quad + \frac{1}{N_1} V_3(q_{1,((\alpha-1)*N_2/N_3+\gamma-1)*N_1/N_2+\lambda}, q_{2,(\alpha-1)*N_2/N_3+\gamma}, q_{3,\alpha}) \\
&\quad + \frac{N_3 m_3}{2\beta^2 \hbar^2} (q_{1,((\alpha-1)*N_2/N_3+\gamma-1)*N_1/N_2+\lambda} - q_{1,((\alpha-1)*N_2/N_3+\gamma-1)*N_1/N_2+\lambda+1})^2
\end{aligned} \tag{20}$$

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