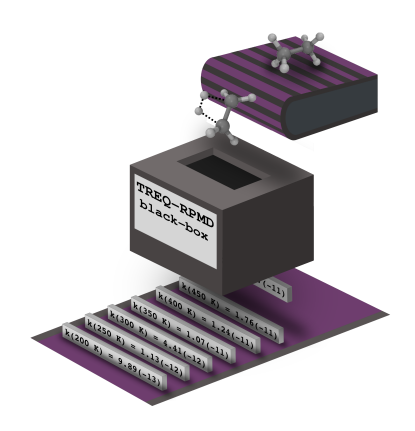
EVB-QMDFF

RPMD and Rate Constant Calculations on Black-Box Potential Energy Surfaces



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EVB-QMDFF Manual

Contents

1	General Information Installation				
2					
	2.1	Prerec	quisites	5	
	2.2	Settin	gs	5	
	2.3	Comp	ilation	6	
3	Prog	Program handling			
	3.1	qmdffg	${ m gen}$	7	
		3.1.1	Introduction	7	
		3.1.2	Example 1: Octane with Orca	8	
		3.1.3	Example 2: Guaicol with Gaussian	9	
	3.2	evbop	vt	10	
		3.2.1	Introduction	10	
		3.2.2	dE-EVB	10	

1 General Information

EVB-QMDFF is a program package for molecular dynamics simulations. It can both perform classical MD and ring polymer molecular dynamics (RPMD) on potential energy surfaces built upon the quantum mechanically derived force field (QMDFF) by GRIMME and the empirical valence bond method by WARSHEL and others.

QMDFF is able to generate a fully-fledged potential energy description for an arbitrary chemical system, leading to a reasonable description of the accessible energy landscape as well as of bond breakings. With this ability, one can e.g. simulate mechanochemical reaction with a single QMDFF. Two or three QMDFF can be coupled via EVB in order to simulate arbitrary chemical elementary reactions. Based on this simple idea, a list of EVB coupling methods, including the newly-developed transition path corrected reaction path EVB-QMDFF (TREQ) was implemented into the program package. Based upon umbrella samplings of the reaction path, a protocol for black-box calculations of chemical reaction rate constants was implemented, serving as a simple and powerful tool for the calculation of those values.

A second branch is the simulation of liquid phases via QMDFF. It is possible to generate a force field for a liquid of arbitrary composition from QMDFFs of the single components. The targeted optimization of noncovalent QMDFF parameters is now possible as well, leading to force fields of high quality for the liquid of interest, which can be sampled via NVT or NpT dynamics classically or with RPMD.

EVB-QMDFF consists of a number of independent programs, serving to fulfill a number of different tasks:

- qmdffgen: Generates a QMDFF from given output of a QM calculation containing the bond orders, charges and frequencies of the reference structure.
- evbopt: Optimizes the parameters of a chosen EVB coupling term (dE, dQ or DG coupling) for a chemical reaction based on two QMDFFs (reactand and product) and reference energies of the reaction path.
- egrad: Calculates energies and gradients for a given trajectory of structures with one of the given PES descriptions.
- evb_qmdff: Performs geometry optimizations, gradient and frequency calculations of chemical systems with one of the given PES descriptions.

- irc: Optimizes the transition state and, starting from it, a reaction path with the IRC method.
- dynamic: Molecular dynamics simulations, both classical and RPMD, on QMDFF and EVB-QMDFF potential energy surfaces. Forces can be applied on different collective variables, for sampling of certain chemical processes.
- rpmd: Calculation of chemical reaction rate constants based on RPMD on EVB-QMDFF potential energy surfaces.
- evb_kt_driver: With a reaction path as input, reference calculations, PES setup with TREQ and RPMD samplings are done in black-box fashion, resulting in temperature-dependent reaction rate constants and Arrhenius parameters.
- mult_qmdff: The QMDFF of a solvent box is built from given QMDFFs of single solvent molecules.
- qmdffopt: After automatic builtup of a suited trainset, noncovalent QMDFF parameters can be optimized for better solvent properties.

2 Installation

EVB-QMDFF consists of pure Fortran code and thus is quite easy to install.

2.1 Prerequisites

All you need is:

- A Fortran compiler (gfortran or ifort)
- BLAS and LAPACK linear algebra libraries
- FFTW3 Fourier transform libraries
- Some of the programs (**evbopt** and **rpmd**) can be run in parallel using MPI. If you want to compile and run them in parallel, MPI must be installed.

2.2 Settings

After downloading EVB-QMDFF, go into the EVB-QMDFF main folder and open the Makefile located there. In its header, the Fortran compilers (GNU or Intel) can be chosen:

```
# Fortran compiler for serial version:
FC = gfortran # GNU Fortran
# FC = ifort # Intel Fortran compiler
# Fortran compiler for MPI version
# FC = mpif90 # GNU Fortran
# FC = mpifort # Intel Fortran compiler
```

The serial GNU compiler is chosen as default, please uncomment your preferred version. By default, the single program binaries are located both into the bin subdirectory of the program folder and the bin directory in your home (/bin). Plase edit this part right below the compiler settings to your preferred location:

```
# Location of the compiled and linked executables
BINDIR = ~/bin
```

Now save the Makefile and copy it to the src subdirectory.

2.3 Compilation

In the src subdirectory, execute:

make

and the objects and executables will be compiled. Now, EVB-QMDFF is ready to use.

3 Program handling

In this chapter, the handling of each program will be explained. For each of them, at least one example is located into the examples folder of the EVB-QMDFF directory. Switch to the subfolders named after the respective programs in order to calculate the examples prepared for you if you want to get more experienced in using EVB-QMDFF.

Besides the descriptions provided below, each program has a help function, that gives general information and a complete list of available keywords for handling. It can be invoked with:

```
[program.x] -h
or
```

[program.x] -help

For almost all programs, the main input file is a [name].key file, where all needed commands to execute your calculation are listed.

3.1 qmdffgen

3.1.1 Introduction

qmdffgen uses the output of an QM calculation for a molecule to generate a QMDFF for it. This QMDFF is able to simulate the full configurational landscape of this molecule and further to describe bond breakings correctly. A QMDFF needs a predefine set of information provided by the QM program:

- Optimized geometry
- Wiberg-Mayer bond orders
- Hirshfeld charges
- Hessian matrix

Provided the correct settings, all those can be obtained from a single QM calculation (see below). After doing the QM calculation, qmdffgen can be invoked:

```
qmdffgen.x
```

The program is interative and asks you for the Software used for the reference calculation (currently available: orca, Gaussian, Turbomole and CP2K), the number of QMDFFs to generate (one to three), and the prefix of the QM input files (name without file ending). Then, the QMDFF is generated in a black-box procedure and directly useable for other calculations!

3.1.2 Example 1: Octane with Orca

Folder: qmdff/ocane_orca/

The input as well as the output of the QM reference calculation with orca is placed here. Basically, the keywords for the reference calculation with orca are always the same. Open the file octane.inp:

```
! PBEO D3 def2-SVP opt freq
* xyz 0 1
C
          -8.36972
                            0.54078
                                           -0.36595
C
          -6.84957
                            0.56234
                                           -0.35222
Η
          -0.30022
                            5.11409
                                            2.54651
Η
          -0.26113
                            4.29744
                                            0.97442
%output
        Print[ P_Hirshfeld ] 1
end
```

A geometry optimization of the octane molecule was done with the PBEO DFT functional (with Grimme D3 dispersion correction) and the cheap def2-SVP basis set. For useful output, a geometry optimization opt followed by a frequency calculation freq need to be done, further, the section

```
%output
Print[P_Hirshfeld] 1
end
```

must be added for the calculation of Hirshfeld charges. The **qmdffgen** program needs the octane.hess and octane.out files as reference.

Since these are present, simply type

qmdffgen.x

and give the respective commands during the interactive input. Then, the QMDFF octane.qmdff will be generated.

During the calculation, the central quality parameters are printed into the terminal:

Zero point vibr. energy comparison (FF/true, kcal) : 150.432 154.650 Mean absolute deviation (MAD) of frequencies (cm-1) : 54.560

The MAD value in particular is a good indicator of the QMDFF-quality. It is calculated by performing frequency calculations for the QM reference and the QMDFF hessian and comparing them. Good QMDFFs achieve values between 20 and 50, if the value is above 100, something might went wrong during the QM reference calculation or the optimization. If the error cannot be located, please let me know. The QMDFF was written into octane.qmdff. This file is the entrypoint for all the other programs included into EVB-QMDFF.

A number of additional output files was produced during the QMDFF optimization. these are:

- octane_qmdff.log: Detailed information about the QMDFF generation (QMDFF internal coordinates, fragments, Levenberg-Marquardt fit of the force constants and QMDFF quality). See the QMDFF paper [?] for more details about the whole procedure.
- octane_molden.out: This file can be opened with the molden molecule viewer in order to visualize the normal modes of the different frequencies of the QMDFF hessian.
- coord_def.inp: List of internal coordinates used to construct the QMDFF (bonds, angles, dihedrals) in the formate also used for several other programs inside QMDFF (irc, rpmd, ...).
- coord_analysis.dat: Values of all internal coordinates, of the Wilson matrix for transformation of cartesian gradients into intenal coordinates and the Wilson matrix derivative for the transformation of cartesian Hessians into internal coordinates. These matrices are used for setup of the DG-EVB coupling term in evbopt.

3.1.3 Example 2: Guaicol with Gaussian

Folder: qmdff/guaicol_gaussian/

For Gaussian, geometry optimization and frequency calculation must be done separately, since a combined geopt+freq calculation writes out the Hessian in an incomplete formate. Therefore, the file guaicol_opt.gjf performs the geometry optimization and the file guaicol_freq.gjf performs the calculation of the Hessian.

The QMDFF can again be calculated by generated by typing:

```
qmdffgen.x
```

Note, that the prefix of the QM reference must be that of the frequency calculation, i.e., guaicol_freq.

The quality of the generated QMDFF can be seen from the lines:

```
Zero point vibr. energy comparison (FF/true, kcal) : 85.684 86.016
Mean absolute deviation (MAD) of frequencies (cm-1) : 26.815
```

The MAD value of 26 suggests a good description of the low-energy part of the potential energy surface of Guaicol with the generated QMDFF (which was written to guaicol_freq.qmdff. The other output files are the same as explained in the first example.

3.2 evbopt

3.2.1 Introduction

This program generates and optimizes the parameters of EVB coupling terms. The coupling smoothly connects two QMDFFs, therefore these need to be present (generated with qmdffgen) when an EVB coupling term is optimized. In addition, some reference data describing the reaction must be present, serving as the training set for the optimization routine. The xyz-coordinates of the structures and the respective reference energies in a linewise format must be present. Usually, a reaction path, generated from a NEB or IRC calculation, would be used for this. For the RP-EVB and the related TREQ method, a reaction path is mandatory.

3.2.2 dE-EVB

Folder: evbopt/dE-EVB/

The simplest EVB coupling version is the energy-gap (dE)-EVB coupling. Here, the offdiagonal coupling term depends only on the energetic distances of both QMDFFs at the current coordinates $(C(\Delta E))$. A bunch of different functions depending on ΔE can be used, the easiest are the constant coupling C = a with no ΔE -dependence at all and the 1g coupling $C(\Delta E) = a \exp(-b(\Delta E)^2)$ with a Gaussian shape.

The main input file is the evbopt_de_1g.key file, where all essentially all control parameters are listed:

```
FFNAME min1.qmdff min2.qmdff

2EVB -272.22659664700754 -272.16798367875066

path_energies ref.dat

path_structure struc.xyz

coupling sd2

shift_manual
```