

EREORG User Manual

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About PCET

PCET (**P**roton-**C**oupled **E**lectron **T**ransfer) is a program package originally developed in the Sharon Hammes-Schiffer research group at the Department of Chemistry and Biochemistry, University of Notre Dame by A. Soudackov, H. Decornez and I. Rostov in the framework of the project "Theoretical Study of Proton-Coupled Electron Transfer Reactions in Solution" funded by the National Science Foundation. The package is the property of the Sharon Hammes-Schiffer research group.

Disclaimer

The developers and maintainers of the PCET package and its offsprings *do not* guarantee that the package is free from errors. They *do not* accept any responsibility for any loss or damage that may result from its use. Users are not entitled to redistribute the program to third parties without consent of the owners.

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Chapter 1

Introduction

EREORG is a utility package (part of PCET) designed to calculate the reorganization energy matrices using the Ellipsoidal Model (ELCM) or Frequency Resolved Cavity Model (FRCM) for advanced solvation calculations in the framework of the dielectric continuum model.

1.1 Functionality

1.1.1 General features

- Solvent models
 - continuum electrostatic model with ellipsoidal cavity [6]
 - advanced continuum Frequency Resolved Cavity Model (FRCM) with cavities of molecular shape [7]
- Free energy surfaces as functions of scalar solvent coordinates (on the grid or along a given path)
 - diabatic and adiabatic free energy curves for single ET reaction between any two EVB states
- Rates
 - nonadiabatic rates for a single ET reaction for any pair of EVB states

1.1.2 Limitations

There are two main limitations in the current version of PCET program:

- the four-state MS-EVB model is implemented. It means that maximum two coupled charge transfer processes can be described. Accordingly, four different charge distributions in the reacting system should be defined.

1.2 Programming Language

All the routines in the current version of PCET are written mostly in FORTRAN 90. Memory allocation is done using FORTRAN 90 rules and syntax. FORTRAN 90 features used in FRCM related routines were removed to maintain the backward compatibility with the original FRCM code written in FORTRAN 77.

1.3 Target Computers

Currently, PCET package can be built on the following platforms:

- Linux (Intel ifort and PGI pgf90 compilers with optional use of Intel MKL libraries)
- OS X 10.7+ (GNU gfortran compiler)

1.4 Required External Libraries

BLAS and LAPACK libraries (MKL libraries for Intel compiler)

1.5 Arithmetic Precision

- All real variables and parameters are specified as `real(kind=8)`
- All integer variables are specified as `integer(kind=4)`

1.6 Units

The program uses kcal/mol for energy, Angströms for length, and picoseconds for time. All the charges and masses are in atomic units.

1.7 The REORG directory structure

subdirectory	contents
bin	executable and submission scripts
docs	documentation (in TeX and PDF formats)
source	FORTRAN source files, Makefile etc.
tests	input and output files for testing

Chapter 2

Theoretical Background

2.1 Solvation models

2.1.1 Ellipsoidal Model

In this model the solute is represented by a set of point charges located on one axis. Different charge distributions corresponding to VB basis states are modelled by different magnitudes of point charges. The point charges representing the solute are placed on the main axis of an axially symmetrical ellipsoidal cavity embedded in a dielectric continuum solvent characterized by the inertial (ϵ_0) and electronic (optical) (ϵ_∞) dielectric constants (see Fig. 2.1).

The total polarization potential $\phi(\mu, \lambda)$ at the point (μ, λ) (elliptical coordinates) inside the cavity can be expressed as[6]

$$\phi(\mu, \lambda) = \sum_{n=0}^{\infty} B_n(\epsilon_0) P_n(\mu) P_n(\lambda) \quad (2.1)$$

where B_n are constants and P_n are the Legendre functions of the first kind. The expression for the coefficients B_n is readily obtained from traditional electrostatics:

$$B_n = \frac{2(2n+1)\beta_n}{R} \left(\frac{1}{\epsilon_0} - 1 \right) \frac{Q_n(\lambda_0)}{P_n(\lambda_0)} \left[1 - \frac{1}{\epsilon_0} \frac{\lambda_0 - P_{n-1}(\lambda_0)/P_n(\lambda_0)}{\lambda_0 - Q_{n-1}(\lambda_0)/Q_n(\lambda_0)} \right]^{-1} \quad (2.2)$$

Here R is the interfocal distance, λ_0 defines the ellipsoid boundary, Q_n are the Legendre functions of the second kind, and β_n is given by the sum over

$$\beta_n = \sum_{k=1}^N q_k P_n(\mu_k) \quad (2.3)$$
$$\phi_{\text{in}} = \phi - \phi_{\infty} \quad (2.4)$$

The reorganization energy matrix elements are calculated then on the grid along the proton coordinate by using standard definitions[1]:

$$t_{ij} = - \sum_k q_k^{(i)} (\phi^{(j)} - \phi_\infty^{(j)}) \quad (2.5)$$

$$t_{ij}^{(\infty)} = -\sum_k q_k^{(i)} \phi_\infty^{(j)} \quad (2.6)$$

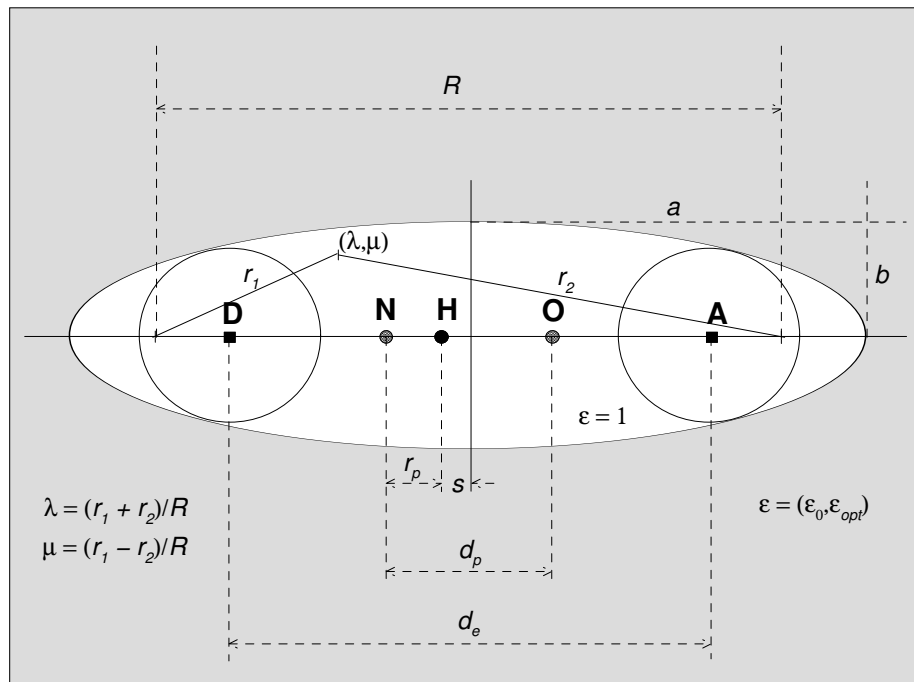


Figure 2.1: Ellipsoidal model cavity and its geometrical parameters.

where indices i and j label charge distributions corresponding to the VB basis states.

2.1.2 FRCM

Relevant references:

- Chudinov, Napolov, and Basilevsky, Chemical Physics **160**, 41-54 (1992).
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- Soudackov and Hammes-Schiffer, J. Chem. Phys. **111**, 4672 (1999).

Description of algorithm

Reorganization energy matrix elements

The multistate continuum theory requires the calculation of the solvent reorganization energy matrix elements defined as

$$t_{ij} = - \int d\mathbf{r} \rho_i \hat{\mathcal{K}}_{\text{in}} \rho_j = - \int d\mathbf{r} \rho_i \phi_j^{(\text{in})}, \quad \phi_j^{(\text{in})} = \hat{\mathcal{K}}_{\text{in}} \rho_j \quad (2.7)$$

and

$$t_{ij}^{(\infty)} = - \int d\mathbf{r} \rho_i \hat{\mathcal{K}}_{\infty} \rho_j = - \int d\mathbf{r} \rho_i \phi_j^{(\infty)}, \quad \phi_j^{(\infty)} = \hat{\mathcal{K}}_{\infty} \rho_j. \quad (2.8)$$

Here the operator $\hat{\mathcal{K}}_{\infty} = \hat{\mathcal{K}}(\epsilon_{\infty})$ and $\hat{\mathcal{K}}_{\text{in}} = \hat{\mathcal{K}}(\epsilon_o) - \hat{\mathcal{K}}(\epsilon_{\infty})$, where $\hat{\mathcal{K}}(\epsilon)$ is a dielectric Green function for the medium with dielectric constant ϵ and ϵ_o and ϵ_{∞} are the static and electronic dielectric constants, respectively. The calculation of these reorganization energies requires the calculation of the potentials $\phi_j^{(\infty)} = \hat{\mathcal{K}}(\epsilon_{\infty})\rho_j$ and $\phi_j^{(\text{tot})} = \hat{\mathcal{K}}(\epsilon_o)\rho_j$ caused by the charge density ρ_j in a dielectric continuum with dielectric constant ϵ_{∞} and ϵ_o , respectively. The inertial potential may be calculated from these two potentials:

$$\phi_j^{(\text{in})} = \phi_j^{(\text{tot})} - \phi_j^{(\infty)}. \quad (2.9)$$

The FRCM method is used to calculate these potentials.

Calculation of the electrostatic potential due to solvent response

The standard PCM method is used to calculate the electronic solvent potential $\phi_j^{(\infty)}$. In this method, the entire charge density ρ_j is placed in a cavity of arbitrary shape with surface S . The dielectric constant is equal to unity inside the cavity and is equal to ϵ_∞ outside the cavity. The electronic solvent potential $\phi_j^{(\infty)}$ is calculated by solving the Poisson equation with the appropriate boundary conditions at the surface of the cavity. (Note that the total potential used in the Poisson equation is the sum of the potential due to the charge density ρ_j in a vacuum and the solvent potential $\phi_j^{(\infty)}$ describing the potential due to the solvent response to the density ρ_j .) The solvent potential may be expressed in terms of a surface charge density σ_j on the surface S . Thus, the problem is reduced to the calculation of this surface charge density σ_j corresponding to the charge density ρ_j . The surface charge density σ_j may be defined in terms of a standard integral equation, which must be solved iteratively.

The FRCM method is used to calculate the total solvent potential $\phi_i^{(\text{tot})}$. In this method, two cavities are formed around the charge density ρ_j . This leads to an inner surface S_1 and an outer surface S_2 . The entire charge density is assumed to be contained inside the inner cavity. The dielectric constant is equal to unity inside the inner cavity, to ϵ_∞ in the region between the two surfaces S_1 and S_2 , and to ϵ_o in the region outside the outer cavity. The solvent potential $\phi_j^{(\text{tot})}$ is calculated by solving the Poisson equation with the appropriate boundary conditions at the two surfaces S_1 and S_2 . (Again, note that the total potential used in the Poisson equation is the sum of the potential due to the charge density ρ_j in a vacuum and the solvent potential $\phi_j^{(\text{tot})}$ describing the potential due to the solvent response to the density ρ_j .) The solvent potential may be expressed in terms of two surface charge distributions $\sigma_j^{(1)}$ and $\sigma_j^{(2)}$ on the surfaces S_1 and S_2 , respectively. In this case, the problem requires the calculation of both surface charge distributions corresponding to the charge density ρ_j . These surface charge densities may be defined in terms of integral equations which must be solved iteratively.

In order to numerically solve the integral equations for both PCM and FRCM, the surface integrals are computed by dividing the surface into small pieces. These surface elements are denoted “tesserae” or “sectors” (where the terms are used interchangeably, although a sector actually refers to a three-dimensional segment and a tesserae actually refers to only the surface

element). The convergence of the iterative solution of the Poisson equation is determined by the dimensionless parameter SELFCR, where the convergence criterion is defined to be $\text{SELFCR} \times 10^{-4}$ for the outer cavity and $\text{SELFCR} \times 10^{-5}$ for the inner cavity. The default is $\text{SELFCR}=2.0$. In past versions, the keyword PRECISE decreases this to $\text{SELFCR}=0.2$, but this word should not be used in the current version. After the surface charge density has converged, it is renormalized according to the total charge of the solute. If the total surface charge computed from the numerical surface charge density differs from the value determined by the total solute charge by more than CHDIFF directly prior to the renormalization of the surface charge density, an error message is printed. The default value is $\text{CHDIFF}=0.1$. The maximum number of iterations is ITSE, after which an error message is printed and the program is stopped. The default value is $\text{ITSE}=15$.

Thus, the calculation of the electronic and inertial potentials requires the following steps:

1. Generate the surfaces of the two cavities.
2. Generate the tesserae independently for the two cavities.
3. Numerically solve the Poisson equation using PCM for the inner cavity surrounded by solvent with dielectric constant ϵ_∞ to obtain the electronic potential $\phi_j^{(\infty)}$.
4. Numerically solve the Poisson equation using FRCM for the two cavities with solvent of dielectric constant ϵ_∞ in between the two surfaces and ϵ_o outside both cavities to obtain the total potential $\phi_j^{(\text{tot})}$.
5. Calculate the inertial potential $\phi_j^{(\text{in})} = \phi_j^{(\text{tot})} - \phi_j^{(\infty)}$.
6. Use these potentials to calculate the electronic and inertial reorganization energy matrix elements.

Generation of surfaces

In FRCM, the inner cavity is generated by placing a sphere centered on each atom with radius κr_{vdw} , where r_{vdw} is the van der Waals radius for the specific atom and κ is independent of the solvent with a default value of 0.9. In the FRCM program, the surface of the inner cavity may be smoothed by including additional spheres. The key words SMOOTH and NOSMOOTH denote cavity smoothing or the absence of cavity smoothing. The default is

to include smoothing. The parameter SOLRD defines the effective radius (in Angstroms) of the solvent molecules and is used to calculate the excluded volume at the seam of a pair of overlapping spheres. The default value is SOLRD=1 Å. The parameter EXVOL defines the minimum excluded volume (in cubic Angstroms) at such a seam for which an auxiliary sphere is added during smoothing. The default value is EXVOL=1 Å³. If EXVOL (or SOLRD) is large enough, the effect is the same as using the key word NOSMOOTH. The outer sphere cavity is generated by adding δ to the radius of each sphere used to create the inner cavity (including the spheres added for smoothing). In general, δ depends on the solvent.

Generation of tesserae/sectors

The premise of the method for generation of the tesserae (or sectors) is that the surface charges change more rapidly closer to the seam of the intersection between two spheres. Thus, the surface elements should be chosen to be smaller in the regions of close contact of the spheres. The following three steps are used to generate the tesserae.

1. The first step is to create a network of A-sectors for each pair of spheres. These sectors are smaller in contact regions (i.e., where the spheres overlap) and larger in regions near the poles. The tesserae are created through a series of rings moving from a contact region to a pole. The tesserae (or sectors) are defined in terms of the standard polar angles θ and ϕ , where $0 < \theta < 180^\circ$ and $0 < \phi < 360^\circ$, and θ is measured relative to the axis connecting the centers of the pair of spheres. The parameters that determine the sizes of the A-sectors and the basic procedure used to create the A-sectors are as follows.
 - The parameters that determine the sizes of the A-sectors are MODFE1 and MODFE2 for the inner and outer surfaces, respectively. This discussion will use MODFE1 as the example. The maximum angular step size $\Delta\theta$ for θ is defined as

$$N = 400/3/\text{MODFE1}. \quad (2.10)$$

For the default value of MODFE1=6, $N = 22.2^\circ$. The first two rings near the contact region have $\Delta\theta = 0.6N$ and $\Delta\theta = 0.9N$, respectively, and the remaining rings have $\Delta\theta = N$. The values for these angular step sizes are stored in the array DDTET, where the first element is $0.6N$, the second element is $0.9N$, and the

remaining elements are N . The number of elements is such that the sum of all angles is less than 180° . Thus, for MODFE1=6, DDTET=(13.3, 20.0, 22.2, 22.2, 22.2, 22.2, 22.2, 22.2). The angle ϕ has the same angular step size $\Delta\phi$ as θ except for the first two rings, where $\Delta\phi = 1.33\Delta\theta$ and $\Delta\phi = 1.28\Delta\theta$, respectively.

- The A-sectors are created for each pair in the following manner. First, the angle in θ for the overlap between the spheres is determined to exclude it from consideration. Second, the sectors are created through a series of rings, using the angular step sizes from the array DDTET, starting at the contact region and moving toward the pole. The sectors are renormalized to render the total sum of angular steps for θ and ϕ 180° and 360° , respectively. Since this procedure is done for each pair of spheres, for each sphere there is a combination of overlapping networks of A-sectors due to all of its neighbors.
- The relation between MODFE1 and MODFE2 is critical to the numerical accuracy. Typically, for MODFE1=MODFE2, the number of tesserae is significantly greater for the inner surface than for the outer surface. This results from the greater overlap between the larger spheres for the outer surface, leading to a larger value of the excluded angle θ and thus to a smaller number of tesserae for each pair of spheres. In addition, the spheres added for smoothing of the inner surface do not contribute as much to the surface area of the outer cavity. Furthermore, for MODFE1=MODFE2, the tesserae are larger in surface area for the outer surface than for the inner surface. This inconsistency between the size and number of tesserae on the two surfaces leads to numerical instabilities. One approach is to choose MODFE2 such that the maximum area of the tesserae for the inner and outer surfaces is approximately equal. This can be achieved using the equation

$$\text{MODFE2} = \text{MODFE1} * (\kappa r_{vdw} + \delta) / (\kappa r_{vdw}) \quad (2.11)$$

for a typical value of r_{vdw} for the system of interest. Another approach is to choose MODFE2 such that the number of tesserae on the inner and outer surfaces is nearly equal.

2. The second step to create a network of small elements, called B-sectors. These are also defined in terms of the angles θ and ϕ . The angular step sizes are defined in terms of NTETFI1 and NTETFI2 for the inner and

outer cavities, respectively. This discussion will use NTETFI1 as the example. The angular step sizes are defined as

$$\Delta\phi = 360/(\text{NTETFI1} * 120) \quad (2.12)$$

$$\Delta\theta = 180/(\text{NTETFI1} * 60). \quad (2.13)$$

The default is NTETFI1=NTETFI2=1, which leads to step sizes of 3° . This leads to 7200 B-sectors per sphere.

3. In the third step of the generation of the tesserae, the A-sectors and the B-sectors are used to construct a unique set of sectors completely covering the whole surface, called the C-sectors. In this procedure, each B-sector is related to the A-sector derived from the neighboring sphere closest to it. Thus, each B-sector is uniquely related to an A-sector. The B-sectors related to the same A-sector are combined as a cluster called a C-sector. The network of C-sectors completely and uniquely covers the cavity surface. In order to decrease the total number of C-sectors, some of the C-sectors are combined with neighboring smaller ones.

Chapter 3

Compiling and Execution

3.1 Compiling EREORG

To compile EREORG program for a specific platform you need change to the source directory, then copy or link `Makefile.xxx` to `Makefile.machine`, for example

```
ln -s Makefile.xxx Makefile.machine
```

```
make all
```

where `<xxx>` must be substituted by the platform-specific suffix. With the present version only one makefile is supplied:

`Makefile.gfortran` - for GFORTRAN compiler

`Makefile.lnx` - for LINUX platforms with Portland Group
FORTRAN compiler `pgf77`

This will initiate compiling and linking of the program. First, the makefile will build an object library containing all the object files except the object file for the main program. Then the makefile will combine the main object file with the library and produce the executable `reorg_frm-<x.x>-gfortran.x` which will be placed in the `bin` directory.

When you modify the include files with parameters and array dimensions, you **must** recompile all the source codes and build a new library. To remove the old object library run the command

```
make clean
```


3.2 Running EREORG

You can run EREORG program from any directory you wish provided that the path to the executable is listed in your `PATH` variable or you have a link to the executable in your current directory.

You will need two mandatory (and maybe more optional) input files in your current directory. The main input file (control file) with arbitrary name contains the control information (keywords) about the jobs you want to run. It contains also the names of other optional input files you may want to use. The other mandatory file is the file with geometry specification for solvation calculations. The name of this file is specified in the keyword section of the control input file. The detailed information about the structure of the input files is given in the next chapter.

To initiate execution you have to run the command

```
reorg_frm-<x.x>_gfortran.x <name-of-the-control-input-file>
```

The program will generate a screen output (you might want to redirect it to disk using standard redirection rules) and also other optional output files which will be placed in the new subdirectory with the name specified in the title section of the control input file.

Chapter 4

Structure of the DATA Files

4.1 Control Input File

4.1.1 General structure

The control input file consists of blocks separated by empty lines. Each block contains the control information for a particular job. The program reads blocks sequentially and terminates when it finds the word `tw END` in the beginning (first three positions) of the block. All the lines in the control input file must be not longer than 80 symbols. The symbols after first 80 are ignored.

Each block consists of three sections:

1. TITLE SECTION (one line) - the title of the job and optionally the keyword specifying the name of the job. This keyword has the following format: `JOBNAME=<name-of-the-job>`. If this keyword is specified the program will create a subdirectory `name-of-the-job` for optional output files. Otherwise a subdirectory with the standard name (consecutive number of the job) will be created.
2. COMMENT SECTION (one line) - any text specifying the current job
3. KEYWORDS SECTION (as many lines as you want but the total number of symbols should not exceed 480) - the keywords controlling the job. The detailed description of all the keywords and related options is given in the next subsection.

4.1.2 Keywords

There are two types of keywords in the PCET 2.0 program. The keywords of the first type set some global controls for the job and are mandatory. The keywords of the second type provide the control information about specific tasks which should be performed. Usually these keywords require mandatory set of options given in brackets. The format for the first type of keywords is (no spaces before or after "="!)

KEYWORD=<value>

The format for the second type is (no spaces before the opening bracket!)

KEYWORD(Option1,Option2,Option3,...)

The keywords are case-insensitive and should be separated by spaces. The options are separated by commas and are case-insensitive as well.

Below is the list of keywords with detailed descriptions.

SOLV(*Options*)

Defines the model and parameters for solvation calculations. The available options are:

WATER - the solvent is water

H2O - the solvent is water

CH2CL2 - the solvent is methylene chloride (dichloromethane)

MEOH - the solvent is methanol

ETOH - the solvent is ethanol

CH3CN - the solvent is CH₃CN

DCE - the solvent is DCE

THF - the solvent is THF (tetrahydrofuran)

NBZ - the solvent is nitrobenzene

DMF - the solvent is DMF

EPS0= ϵ_0 - static dielectric constant

EPS8= ϵ_{∞} - optical (electronic) dielectric constant

ELLIPSE - simple electrostatic ellipsoidal model will be used

A= a - major semiaxis of the ellipsoidal cavity

B= b - minor semiaxis of the ellipsoidal cavity

FRCM - advanced FRCM model will be used

KAPPA= κ - factor for VdW radii

DELTA= δ - thickness of the inner layer in FRCM

GEOM=<file> - the geometry and EVB charges are given in the external file <file> (the format of the file is described in the next section).

TREAD=<file> - the reorganization energy matrices on the grid are read from external binary file <file> saved in previous calculation.

TWRITE=<file> - the reorganization energy matrices on the grid are saved in the external binary file <file>.

XYZOUT=<file> - the cartesian coordinates are written to the external file <file> in XYZ format.

NOSYMD - the reorganization energy matrices are not about the diagonal from top left to bottom right. This keyword should only be used to check how good the approach is working. By default the reorganization energy matrices will be symmetrized. (See solint.f and tmat.f)

SYMT - the solvent reorganization energy matrices are symmetrized so that the properties for a symmetric PCET system are correctly reproduced.

SYMET - the solvent reorganization energy matrices are symmetrized so that the properties for a symmetric ET system are correctly reproduced. (NOTE, this should only be used with the ET2 keyword.

SYMPT - the solvent reorganization energy matrices are symmetrized so that the properties for a symmetric PT system are correctly reproduced. (NOTE, this should only be used with the PT2 keyword.

REDDENS - All the elements of the solvent reorganization energy matrices which depend on 2b are obtained from all the other elements depending only on 1a,1b,2a. This should always be true anyway. A system for

which this is not true has some major problems and should be used with caution. This keyword can also be used when the charges for the 2b state for solvation are not obtainable. (simply put in any charges that are not all zero and such that the column has at least one different element from the other columns:1a,1b,and 2a.)

EREORG(*Options*)

Calculates [4x4] reorganization energy matrices for four given charge distributions. Also calculates one-dimensional free energy curves and nonadiabatic rate for a single ET reaction between specified EVB states. The available options are:

REAC=<N> - the EVB state N (1/2/3/4) will be considered as a reactant state in the ET reaction. If this option is not specified then the state 1 (1a) is assumed to be the reactant state.

PROD=<M> - the EVB state M (1/2/3/4) will be considered as a product state in the ET reaction. If this option is not specified then the state 3 (2a) is assumed to be the product state.

ZE1=<ZE1> - the left limit on the grid along the solvent coordinate z_e (kcal/mol). The default is $-3\lambda_{ET}$, where λ_{ET} is the ET reorganization energy.

ZE2=<ZE2> - the right limit on the grid along the solvent coordinate z_e (kcal/mol). The default is $+3\lambda_{ET}$, where λ_{ET} is the reorganization energy.

NZE=<N> - number of points along the grid. The default is 100.

EBIAS=<E> - the gas phase bias (kcal/mol) for the ET pair. The default is 0.

VEL=<E> - electronic coupling (kcal/mol) for the ET pair. The default is 0.

PLOT=<filename> - name of the output file with the free energy profiles (the default filename is `et.dat`).

RATE=<T> - the nonadiabatic rate at $T = <T>$ K will be calculated. The default temperature is 298.15 K.

4.2 Additional INPUT Files

4.2.1 Geometry and charges specification

This input file specifies the geometry and basis EVB point charge distributions for solvation calculations. The examples are stored in the `tests` subdirectory as files with `.sol` extensions.

Line 1: Keywords

One keyword is mandatory and defines the format for the geometry specification. You should specify either `XYZ` (cartesian coordinates) or `tw INT` (internal coordinates). If the file is intended for FRCM solvation calculations, you can also specify any FRCM keywords described in the FRCM manual.

Line 2 - Title

Any title specifying the geometry.

Line 3 - Comment

Any text related to geometry (can be empty).

Lines 4:N - Geometry specification

The geometry can be specified either in cartesian coordinates (`<element_symbol> <x> <y> <z>`) or in internal coordinates (MOPAC Z-matrix format). The allowed element symbols are H-No and also symbols 'De', 'Ae', 'Dp', 'Ap', 'Ps' for general electron donor, electron acceptor, proton donor, proton acceptor, and pseudoatom, respectively. Format is free and the section is terminated by an empty line.

Line N+2 - Title for charge distribution section

Any title specifying the EVB charges (must contain `CHARGES` keyword).

Line N+3:M - Charges for EVB states

Four columns in free format, the columns specify charges on atoms for EVB states 1a, 1b, 2a, and 2b, respectively. Note that the order of point charges should exactly correspond to the order of atoms in the geometry section.

The input of charges is also terminated by an empty line. **IMPORTANT: the first line in this section should contain TOTAL charges corresponding to each EVB state.**

4.3 OUTPUT Files

4.3.1 Standard output

Standard unit 6 is used for standard output throughout the program. You can redirect the standard output to the disk file using standard redirection rules for your operating system.

4.3.2 Geometry output

The program can produce the geometry files if the XYZOUT option is specified with the HGAS or SOLV keywords. The files are written in standard "xyz" format, so you can use any visualization software to check the geometry (for example, MOLDEN).

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