Manual for SS-QRRK utility code

SS-QRRK: A Program for System-Specific Quantum Rice-Ramsperger-Kassel Theory

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1. Introduction and recommended citations

This is a manual for the *SS-QRRK* utility code, which is included in the *Polyrate* program (in the directory polyrate2016–2A/util/SS-QRRK/). The *SS-QRRK* utility code is a program for calculating pressure-dependent rate constants using variational transition state theory. To execute the code, put input files with the .exe file in a working directory, and then type " ./FALLOFF.exe &".

Recommend citations:

(1) J. Zheng, J. L. Bao, R. Meana-Pañeda, S. Zhang, B. J. Lynch, J. C. Corchado, Y.-Y. Chuang, P. L. Fast, W.-P. Hu, Y.-P. Liu, G. C. Lynch, K. A. Nguyen, C. F. Jackels,

A. Fernandez Ramos, B. A. Ellingson, V. S. Melissas, J. Villà, I. Rossi, E. L. Coitiño, J. Pu, T. V. Albu, A. Ratkiewicz, R. Steckler, B. C. Garrett, A. D. Isaacson, and

D. G. Truhlar, POLYRATE–version 2016-2A, University of Minnesota, Minneapolis, 2016 (2) "Kinetics of Hydrogen Radical Reactions with Toluene by Chemical Activation Theory Employing System-Specific Quantum RRK Theory Calibrated by Variational Transition State Theory," J. L. Bao, J. Zheng, and D. G. Truhlar, Journal of the American Chemical Society **138**, 2690-2704 (2016). dx.doi.org/10.1021/jacs.5b11938

(3) "Predicting Pressure-Dependent Unimolecular Rate Constants Using Variational Transition State Theory with Multidimensional Tunneling Combined with System-Specific Quantum RRK Theory: A Definitive Test for Fluoroform Dissociation,"
J. L. Bao, X. Zhang, and D. G. Truhlar, Physical Chemistry Chemical Physics 18, 16659-16670 (2016). dx.doi.org/10.1039/C6CP02765B

See also:

(4) "Barrierless association of CF_2 and dissociation of C_2F_4 by variational transition state theory and system-specific quantum Rice-Ramsperger-Kassel theory," J. L. Bao, X. Zhang, and D. G. Truhlar, Proceedings of the National Academy of Sciences, U.S.A., 113, 13606–13611 (2016). dx.doi.org/10.1073/pnas.1616208113

2. Theoretical background of SS-QRRK

System-specific quantum RRK (SS-QRRK) theory is an efficient way to incorporate variational effects, multidimensional quantum mechanical tunneling, and multi-structural torsional anharmonicity into the calculation of pressure-dependent rate constants in the falloff region of chemical reactions. Recall that unimolecular reactions may have pressure-dependent rate constants because only in the high-pressure limit can the usual assumption of thermally equilibrated states in the reactant be justified. Furthermore, association reactions (whose reverse reactions are unimolecular) may also be pressure dependent because the newly formed states of the associated species (which can be an intermediate or a product) may redissociate before they are equilibrated. (Nonequilibrium effects can also be present in bimolecular reactions that produce two or more products, but in such cases they are expected to be much smaller, and we do not include them in those cases.)

SS-QRRK theory uses information obtained from the high-pressure-limit canonical rate constants (i.e., the thermal equilibrium rate constants) to compute the microcanonical rate constants using quantum RRK theory fitted to these high-pressure results; the resulting microcanonical rate constants are then used in the computation of the pressure-dependent rate constants. Current implementations of SS-QRRK are for three types of reactions: unimolecular reactions (both isomerization, $A \rightarrow P$, and dissociation, $A \rightarrow P1 + P2$), bimolecular association reactions ($A+B\rightarrow P$), and bimolecular reactions (both unimolecular reactions (both unimolecular dissociation and unimolecular isomerization) are treated using Hinshelwood-Lindemann theory; bimolecular association reactions are treated with a chemical activation mechanism.

2.1 Hinshelwood—Lindemann mechanism for unimolecular reactions

Before we proceed to consider energy-resolved rate constants, we first examine the classical Lindemann mechanism for unimolecular reaction $A \rightarrow P$:

Step 1: A + M
$$\xleftarrow{k_1(T)}{k_c(T)}$$
 A* + M
Step 2: A* $\xrightarrow{k_2(T)}$ P

In the original Lindemann model, the rate constants are assumed to be temperature-dependent and energy-independent. The thermal activation rate constant k_1 is modeled as:

$$k_1 = Z \exp(-E_0 / k_{\rm B}T) \tag{1}$$

where E_0 is called the threshold energy, k_B is Boltzmann's constant, and Z is the bimolecular collision rate constant (in units of cm³ molecule⁻¹ s⁻¹) between the reactant A and the bath gas M:

$$Z = \sigma \langle v_{\rm rel} \rangle \tag{2}$$

where σ is a velocity-independent collision cross section between A and M, and $\langle v_{rel} \rangle$ is the average relative velocity. The rate constant k_c of the de-activation step is simply equal to Z, which assumes that every collision of the energized molecule A* with M leads to de-energization (this is called the strong-collision assumption). The dissociation rate constant k_2 is simply the high-pressure-limit Arrhenius pre-exponential factor A_{uni}^{∞} . Using the steady-state approximation for A*, the unimolecular dissociation rate constant

 k_{uni} (which is defined as $-\frac{1}{[A]}\frac{d[A]}{dt}$) is:

$$k_{\rm uni} = \frac{k_1[M]}{1 + k_c[M]/k_2}$$
(3)

In the high-pressure-limit ([M] >> 1),

$$k_{\text{uni}}^{\infty} = k_1 k_2 / k_c = A_{\text{uni}}^{\infty} \exp(-E_0 / k_{\text{B}}T)$$
(4)

and the unimolecular dissociation is a (pseudo-) first-order reaction; in the low-pressure-limit ([M] \approx 0), $k_{uni}^0 = k_1[M]$, and the dissociation reaction is of second order.

As discussed next, our approach goes beyond the original theory mentioned above.

In our approach, $k_{uni}^{\infty}(T)$ is computed using VTST with multidimensional tunneling; from now on, we will use canonical VTST (CVT) and small curvature tunneling (SCT) as an example of high-pressure-limit rate constants for our description in this manual (although in practice one may, for example, use other tunneling methods if desired). The "threshold energy" E_0 of the unimolecular dissociation reaction (step 2) is set equal to the temperature-dependent high-pressure-limit Arrhenius activation energy

 $E_{a}^{\text{CVT/SCT}}(T)$, which is computed as:

$$E_{a} = \begin{cases} \frac{E(T^{4} + 2T_{0}T^{3} - T_{0}^{2}T^{2})}{(T^{2} + T_{0}^{2})^{2}} + nRT & \text{endothermic reaction} \\ \frac{E(T^{4} + 2T_{0}T^{3} - T_{0}^{2}T^{2})}{(T^{2} + T_{0}^{2})^{2}} + \frac{nRT^{2}}{T + T_{0}} & \text{exothermic reaction} \end{cases}$$
(5)

where E (kcal/mol) and T_0 (K) are fitting parameters in the following fitting formula for fitting the computed high-pressure-limit rate constants:

$$k = \begin{cases} A(\frac{T}{300})^{n} \exp\left[-\frac{E(T+T_{0})}{R(T^{2}+T_{0}^{2})}\right] & \text{endothermic reaction} \\ A(\frac{T+T_{0}}{300})^{n} \exp\left[-\frac{E(T+T_{0})}{R(T^{2}+T_{0}^{2})}\right] & \text{exothermic reaction} \end{cases}$$
(6)

and $A_{\text{uni}}^{\infty}(T)$ is computed as

$$A_{\rm uni}^{\infty}(T) = k_{\rm uni}^{\infty}(T) \exp[E_{\rm a}^{\rm CVT/SCT}(T)/RT]$$
⁽⁷⁾

The value of $A_{uni}^{\infty}(T)$ is now temperature-dependent and we shall call it the frequency factor when it is used in SS-QRRK theory (see below).

The energization and dissociation rate constants are treated as energy-dependent using SS-QRRK theory, and the collisional de-energization rate constant is computed based on modified strong collision model, which is discussed next. In particular, we consider the following thermal activation mechanism:

Step 1:
$$A(T) + M \xleftarrow{k_1(E; T)}{k_c(T)} A^*(E) + M$$

Step 2: $A^*(E) \xrightarrow{k_2(E)} P$

Step 1 is the thermal activation step, in which the thermally equilibrated fluoroform molecules (denoted as A(T) in the mechanism to denote that species A is at temperature T) collide with bath gas M to produce the rovibrationally excited reactant $A^*(E)$ with total rovibrational energy E randomized among all modes. The rate constant of thermal activation k_1 is a function of energy E and is parametrically dependent on T. The de-activation collisional rate constant k_c is assumed to be temperature-dependent as in Lindemann–Hinshelwood theory; it is computed as

$$k_c(T) = Z\beta_c \tag{8}$$

where Z is the Lennard-Jones collision rate constant, and β_c is the collision efficiency.

To model $k_1(E; T)$ and $k_2(E)$, we use single-frequency quantum Rice-Ramsperger-Kassel theory (QRRK), in which rotation is not treated explicitly, and the molecule consists of *s* identical oscillators with vibrational frequency \overline{v} , where we express \overline{v} in wave numbers (cm⁻¹). In practice, we set *s* equal to number of vibrational degree of freedom (3*N*-6 for non-linear molecule), and \overline{v} is chosen to be the reactant geometric mean frequency. The geometric mean frequency is computed as:

$$\overline{v} = \left(\prod_{i=1}^{s} v_i\right)^{1/s} \tag{9}$$

where v_i is the vibrational frequency of normal mode *i*, and *s* is the number of vibrational degrees of freedom.

We denote the energy of the reactant molecule as E and set E equal to $nhc \overline{v}$, where h is Planck's constant, c is the speed of light, and n is $E/hc \overline{v}$. In the QRRK model of the microcanonical rate constant at energy E, a unimolecular reaction happens when a specific vibrational mode associated with the reaction coordinate possesses a critical energy (or threshold energy) E_0 equal to $mhc \overline{v}$. Note that n and m are not usually integers.

First consider $k_1(E,T)$. The equilibrium constant for step 1 is the fraction of the molecules with *n* quanta of excitation, which is

$$K^{\text{QRRK}}(n,T) = \exp(\frac{-nhc\bar{\nu}}{k_{\text{B}}T})[1 - \exp(\frac{-hc\bar{\nu}}{k_{\text{B}}T})]^{s} \frac{(n+s-1)!}{n!(s-1)!}$$
(10)

The equilibrium constant equals the ratio of forward to reverse rates:

$$K^{\text{QRRK}}(n,T) = k_1^{\text{QRRK}}(E = nhc\overline{\nu};T)/k_c(T)$$
(11)

Substituting Eq. (10) into Eq. (11) and solving for the forward rate yields $k_1^{\text{QRRK}}(E;T)$.

Next consider the QRRK rate constant $k_2(E)$; this is equal to the frequency factor times the fraction of molecules with at least *m* quanta in one chosen mode, and for a quantum mechanical oscillator with *s* identical frequencies \overline{v} and temperature *T*, this is given by

$$k_2^{\text{QRRK}}(E = nhc\overline{\nu}) = A^{\text{QRRK}} \frac{n!(n-m+s-1)!}{(n-m)!(n+s-1)!}$$
(12)

In SS-QRRK, we set the frequency factor A^{QRRK} equal to $A^{\infty}_{\text{uni}}(T)$ given by Eq. (7), and we set the threshold energy equal to the high-pressure local Arrhenius activation energy given by Eq. (5), which yields

$$m = E_{a}^{\text{CVT/SCT}}(T) / hc\overline{\nu}$$
(13)

The analog of Eq. (3) is

$$k_{\text{uni}}(T,p) = \sum_{n=m}^{+\infty} \frac{k_1^{\text{QRRK}}(E = nhc\overline{\nu};T)[M]}{1 + \frac{k_c(T)[M]}{k_2^{\text{QRRK}}(E = nhc\overline{\nu})}}$$
(14)

where $k_c(T)$ is obtained by Eq. (8), $k_1^{\text{QRRK}}(E;T)$ is obtained by Eqs. (10) and (11), and $k_2^{\text{QRRK}}(E)$ is from Eq (12).

Now, it can be shown why we choose the frequency factor and threshold energy by Eqs. (7) and (5). In the high-pressure-limit, Eq. (14) reduces to:

$$k_{\text{uni}}(T,p) = \sum_{n=m}^{+\infty} \frac{k_1(E = nhc\overline{\nu};T)k_2(E = nhc\overline{\nu})}{k_c(T)}$$
(15)

and, by carrying out the summations, Eq. (4) is obtained. The high-pressure-limit canonical rate constants computed by SS-QRRK theory are therefore same as the ones determined by CVT/SCT theory. Therefore our SS-QRRK microcanonical rate constant formalism can build in variational effects, multidimensional tunneling based on the directly calculated potential energy surface along the tunneling path, and vibrational and torsional anharmonicity with approximately the same computational effort that CVT/SCT allows one to include them efficiently in the high-pressure limit.

2.2 Chemical activation mechanism for bimolecular association

The chemical activation mechanism for a bimolecular reaction with an associated intermediate is:

$$\begin{array}{ccc} A+B \xleftarrow{k_{1}(T)} & AB^{*} & \xrightarrow{k_{2}(E)} P \\ & & \downarrow k_{c}[M] \\ & & AB \end{array} \tag{Type 1}$$

and the chemical activation mechanism for a for bimolecular association reaction is:

$$A+B \xleftarrow{k_1(T)}{\longleftarrow} AB* \xrightarrow{k_c[M]} AB$$
(Type 2)

where A and B are reactants, AB* is the ro-vibrationally excited unimolecular state, AB is the stabilized unimolecular state (i.e., AB is the intermediate in a type-1 reaction, and it is the product of a type-2 reaction), and P is the product in a type-1 reaction.

Applying the steady-state approximation to the chemical activation mechanism chemical activation mechanism yields

$$k_{\text{stab}} = \frac{d[AB]}{[A][B]dt} = k_1^{\text{MS-CVT/SCT}} \sum_{E_0}^{+\infty} \frac{k_c[M] f(E)}{k_c[M] + k_{-1}^{\text{QRRK}}(E) + k_2^{\text{QRRK}}(E)}$$
(16)

$$k_{\rm P} = \frac{d[{\rm P}]}{[{\rm A}][{\rm B}]dt} = k_1^{\rm MS-CVT/SCT} \sum_{E_0}^{+\infty} \frac{k_2^{\rm QRRK}(E)f(E)}{k_{\rm c}[{\rm M}] + k_{-1}^{\rm QRRK}(E) + k_2^{\rm QRRK}(E)}$$
(17)

for reaction type 1 and

$$k_{\text{stab}} = \frac{d[AB]}{[A][B]dt} = k_1^{\text{MS-CVT/SCT}} \sum_{E_0}^{+\infty} \frac{k_c[M] f(E)}{k_c[M] + k_{-1}^{\text{QRRK}}(E)}$$
(18)

for reactions type 2, where f(E) is the fraction of energized species (AB*) at energy *E*, which is given by

$$f(E) = \frac{k_{-1}^{\text{QRRK}}(E)K(E)}{\sum_{E_0}^{+\infty} k_{-1}^{\text{QRRK}}(E)K(E)}$$
(19)

where

$$K(E) = \exp(\frac{-hv}{k_{\rm B}T})[1 - \exp(\frac{-hv}{k_{\rm B}T})]^s \frac{(n+s-1)!}{n!(s-1)!}$$
(20)

In eqs (16), (17), and (18), [M] is the concentration of the bath gas, which is calculated using the ideal gas law, [M] = p / RT, E_0 is the threshold energy, which is computed using Eq. (5), and $k^{QRRK}(E)$ denotes a quantum RRK theory energy-resolved rate constant,

which is computed using Eq. (12) with the corresponding *A* factor and threshold energy. The summations in eqs (16), (18), and (19) are evaluated with a step size of one quantum $(h\overline{v})$. The quantity *n*! is computed using a gamma function or Stirling's approximation.

2.3 Energy transfer parameters

The collision efficiency is computed by solving the following equation:

$$\frac{\beta_{\rm c}}{1 - \beta_{\rm c}^{1/2}} = \frac{\left| \langle \Delta E \rangle \right|}{F_E k_{\rm B} T} \tag{21}$$

where $\langle \Delta E \rangle$ is the average vibrational energy transferred during both energization and de-energization processes (note that $|\langle \Delta E \rangle|$ is smaller than the often encountered $\langle \Delta E \rangle_{down}$ parameter that is the average energy transferred in collisions in which the vibrational energy goes down); and F_E is the thermal fraction of unimolecular states above the threshold energy and is defined in Troe's work (Troe, J. J. Chem. Phys. 1977, 66, 4745). For moderate-sized molecules, F_E is nearly unity (Troe, J. J. Phys. Chem. 1979, 83, 114).

The temperature dependence of F_E can be treated with Troe's method when it is needed: F_E can be computed using Troe's definition:

$$F_E = \frac{\int_{E_0}^{+\infty} \rho(E) e^{-E/k_{\rm B}T} dE}{k_B T \rho(E_0) e^{-E_0/k_{\rm B}T}}$$
(22)

or it can be computed using the Whiten-Rabinovitch approximation (J. Troe, *J. Phys. Chem.*, 1979, 83, 114).

The collision efficiency can also be estimated as:

$$\beta_{\rm c} = \left(\frac{\left<\Delta E\right>_{\rm down}}{\left<\Delta E\right>_{\rm down} + F_E k_{\rm B} T}\right)^2 \tag{23}$$

where $\langle \Delta E \rangle_{\text{down}}$ is the averaged energy transferred for *only* the de-activation process, by which we mean for only collisions in which the vibrational energy goes down.

The Lennard-Jones collision rate constant k_{LJ} is the product of the hard-sphere collision rate constant k_{HS} and the dimensionless reduced collision integral $\Omega_{2,2}^*$, which is defined by Hirschfelder et al (Hirschfelder, J. O.; Curtiss, C. F.; Bird, R. B. In Molecular Theory of Gases and Liquids; Wiley and Sons: London, 1967.). Therefore

$$k_{\rm c} = \beta_{\rm c} \Omega_{2,2}^* k_{\rm HS} \tag{24}$$

The values of the reduced collision integral can be evaluated by numerical integration and have been fitted by other workers with various simple algebraic expressions. Usually, we choose Troe's fitting expression (Troe, J. J. Chem. Phys. 1977, 66, 4745; Troe, J. J. Chem. Phys. 1977, 66, 4758) for computing $\Omega_{2,2}^*$:

$$\Omega_{2,2}^{*} = \begin{cases} [0.697 + 0.5185 \log_{10}(\frac{k_{\rm B}T}{\varepsilon_{\rm A-M}})]^{-1} & \frac{k_{\rm B}T}{\varepsilon_{\rm A-M}} \in [3,300] \\ [0.636 + 0.567 \log_{10}(\frac{k_{\rm B}T}{\varepsilon_{\rm A-M}})]^{-1} & \frac{k_{\rm B}T}{\varepsilon_{\rm A-M}} \notin [3,300] \end{cases}$$
(25)

where ε_{A-M} is the Lennard-Jones interaction parameter between molecule A and M and is computed as the geometric average of ε_{A-A} and ε_{M-M} .

The hard–sphere collision rate constant $k_{\rm HS}$ is computed as:

$$k_{\rm HS} = \pi (\frac{d_{\rm A} + d_{\rm M}}{2})^2 \sqrt{\frac{8k_{\rm B}T}{\pi\mu}}$$
 (26)

where the vdW diameters d_A and d_M are computed from the Lennard-Jones parameters σ_{A-A} and σ_{M-M} using the relation $d = 2^{1/6}\sigma$, and μ is the reduced mass of A and M.

3. Fitting high-pressure-limit rate constants

Before carrying out the SS-QRRK calculations, the user should use *Polyrate* to compute the high-pressure-limit rate constants and possibly to perform multi-structural torsional anharmonicity corrections using MSTor code. For transition states without conformational complexity one can use the original VTST, and for transition states with conformational complexity one can use multi-path or multi-structural VTST; in either case one can include multidimensional quantum mechanical tunneling. After computing the high-pressure-limit rate constants, the user should use the following equation to fit the rate constants (different formula should be applied for endothermic and exothermic reactions):

$$k = \begin{cases} A(\frac{T}{300})^{n} \exp\left[-\frac{E(T+T_{0})}{R(T^{2}+T_{0}^{2})}\right] & \text{endothermic reaction} \\ A(\frac{T+T_{0}}{300})^{n} \exp\left[-\frac{E(T+T_{0})}{R(T^{2}+T_{0}^{2})}\right] & \text{exothermic reaction} \end{cases}$$
(27)

where *A* (for unimolecular reaction, its unit is s⁻¹; and for bimolecular reaction, its unit is cm³ molecule⁻¹ s⁻¹), *n* (unitless), *E* (unit: kcal/mol) and T_0 (unit: K) are fitting parameters. The ideal gas constant *R* is in the unit of kcal mo⁻¹ K⁻¹) with value 0.00198588. Then, the temperature-dependent Arrhenius activation energies can be computed from the local slopes of Arrhenius plots, which yields:

$$E_{\rm a} = \begin{cases} \frac{E(T^4 + 2T_0T^3 - T_0^2T^2)}{(T^2 + T_0^2)^2} + nRT & \text{endothermic reaction} \\ \frac{E(T^4 + 2T_0T^3 - T_0^2T^2)}{(T^2 + T_0^2)^2} + \frac{nRT^2}{T + T_0} & \text{exothermic reaction} \end{cases}$$
(28)

Finally, the local Arrhenius frequency factor *A*, which is also temperature-dependent, can be computed as:

$$A^{\infty}(T) = k^{\infty}(T) \exp[E_{a}(T)/RT]$$
(29)

4. Description of input

The input of the SS-QRRK code is template-based; users should use the template that corresponds to their type of calculation and then change the values accordingly. Next we shall discuss the input templates for SS-QRRK calculations and the meanings of the keywords. Some common keywords are only explained once (in section 4.1).

4.1 Unimolecular reaction

The input template for unimolecular reaction (A \rightarrow P) is:

8.789665E+11 26.1 1.013173E+12 26.2 1.167539E+12 26.5 1.361410E+12 26.7 2.658167E+12 28.5 sAstar 30. GeoMeFreq 580.880669720707 MaxE 8000. #END

Meanings of keywords:

numbTrxn	number of temperatures
Trxnlist	list of temperatures. This should be consistent with number of temperatures.
numbP	number of pressures
unitP	unit of the input pressures. Available options are: torr, bar, atm, Pa, kPa, lg10atm, lg10bar, lg10torr
Plist	List of pressures; this should be consistent with number of pressures that user input
ColliIntg	The form of the collision integral one uses. Available options are: Troe, Reid
avgEtrans	Average energy transferred parameter $\triangle E$. Available units are: cm**-1, kcal*mol-1, cal*mol-1. The temperature dependence can be incorporated by using: $\triangle E(T) = \triangle E(T/300)^n$. There are two types of $\triangle E$ parameter: $< \triangle E >_{all}$ (i.e. average energy transferred during both activation and deactivation collisions), $< \triangle E >_{down}$ (i.e., average energy transferred only for deactivation collisions). For example, "avgEtrans 740. cal*mol-1 0. all" means we are using $ < \triangle E >_{all} =740$ cal/mol, and the temperature-dependence coefficient $n=0$ (i.e., we do not consider <i>T</i> -dependence of $< \triangle E >_{all}$).
FE	The F_E parameter for the activated unimolecular states. If user specify a positive number, then it means that the value specified by the user

	specified is directly used as $F_{\rm E}$; if a value of -1 is specified, it means
	that F_E will be computed based on Whitten-Rabinovitch
	approximation (with an additional input file: vibinfo.inp). Please refer
	to section 4.4 for more information about the additional input files.
paramA	Lennard-Johns parameters for reactant A, in the order of σ (unit: Å), a
	(unit: Kelvin, i.e., ε/k_B), molar mass (unit: gram per mole).
paramBath	Lennard-Johns parameters for bath gas.
Method	For unimolecular dissociation or isomerization reaction, the keyword
	"HLQRRK" must be specified.
rxn1 AEa	To specify the A factor (temperature-dependent local frequency factor
	which is determined by eq. (29)) and activation energy (determined b
	eq. (28), in the unit of kcal/mol) for the high-pressure-limit
	unimolecular reaction $A \rightarrow P$. Under this keyword, the user should
	specify A in left column and Ea in right column; this information
	should be consistent with the order of the temperatures that one has
	specified under Trxnlist
sAstar	number of vibrational degree freedom of A
GeoMeFreq	The geometric mean frequency (in the unit of cm ⁻¹) of unimolecular
	reactant A. The geometric mean frequency is computed as eq. (9).
MaxE	The summation upper limit for quantum RRK theory (unit: kcal/mol);
	this value should be set to be high enough so that the summation is
	converged. A value of 8000 is recommended.

4.2 Bimolecular association

The input template for bimolecular association reaction $(A + B \rightarrow P)$ is as follows:

```
#BasicInfo
numbTrxn 3
Trxnlist
   298.15
   300.
   400.
numbP 4
unitP lg10atm
Plist
3.
-0.5
-0.75
-3.
#Collision
ColliIntg
             Troe
avgEtrans 92 cm**-1 0. all
       1.15
FE
             6.00 410.0 93.0
paramPstar
             2.93 38.0 2.0
paramBath
#FalloffMethod
Method ORRK
       NO
Rxn2
rxn1
        rate
2.7165E-14
2.8101E-14
1.3146E-13
#ORRKData
sPstar
             42.
GeoMeFreq
             1085.669208
MaxE
            8000.
rxnneg1AEa
                        29.78
29.8
1.5444E+13
1.5961E+13
2.9953E+13
                        30.25
#END
```

In section 4.1, we have already covered most common keywords. Here, we only explain

new keywords:

Method	For bimolecular reactions, the keyword "QRRK" must be specified in
	order to do SS-QRRK calculation. (Do not be confused with the
	"HLQRRK" keyword, which is specifically used for unimolecular
	reaction with Hinshelwood-Lindemann theory.)
Rxn2 N0	For bimolecular association reaction, which does not involve a stable
	intermediate, the option for "Rxn2" must be "NO".

rxn1	rate	User specified high-pressure-limit rate constant for the forward
		bimolecular association reaction $(A + B \rightarrow P)$.

rxnneg1AEa To specify the *A* factor (temperature-dependent local frequency factor, which is determined by eq. (29)) and activation energy (determined by eq. (28), in the unit of kcal/mol) for the high-pressure-limit unimolecular reaction $P \rightarrow A + B$ (i.e., the reverse reaction of bimolecular association). Under this keyword, the user should specify *A* in left column and *E*a in right column; this information should be consistent with the order of the temperatures that one has specified under Trxnlist

paramPstar Lennard-Johns parameters for the unimolecular product P, in the order of σ (unit: Å), ϵ (unit: Kelvin, i.e., ϵ/k_B), molar mass (unit: gram per mole).

sPstar The number of vibrational degree of freedom for P.

GeoMeFreq The geometric mean frequency (in the unit of cm⁻¹) of unimolecular product P.

4.3 Bimolecular reaction with an intermediate

P) is: #BasicInfo numbTrxn 3 Trxnlist 298.15 300. 400. numbP 3 unitP lg10atm Plist 0.25 0. -0.25 #Collision ColliIntg Troe avgEtranš 92 cm**-1 0. all FE 1.15 6.00 410.0 93.0 paramAB 2.93 38.0 2.0 paramBath #FalloffMethod Method QRRK Rxn2 YES rxn1 rate 2.1632E-16 2.2860E-16 2.4896E-15 #QRRKData sABstar 42. 1092.49876 GeoMeFreq MaxE 8000. rxnneg1AEa 1.7242E+12 26.55 1.9273E+12 26.61 4.3192E+13 28.62 rxn2AEa 2.0228d+12 23.12 2.2603d+12 23.18 3.4447d+13 24.95 #END

The input template for bimolecular reaction with a stable intermediate $(A + B \rightarrow AB \rightarrow$

In section 4.1 and 4.2, we have already covered most common keywords. Here, we only explain new keywords:

Method For bimolecular reactions with a stable intermediate, the keyword "QRRK" must be specified in order to do SS-QRRK calculation. (Do not be confused with the "HLQRRK" keyword, which is specifically used for unimolecular reaction.)

Rxn2 YES	For bimolecular reactions with a stable intermediate, the "Rxn2" must
	be set to be "YES" to indicate the existence of a second-step reaction.
rxn2AEa	To specify the A factor (temperature-dependent local frequency factor,
	which is determined by eq. (29)) and activation energy (determined by
	eq. (28), in the unit of kcal/mol) for the high-pressure-limit
	unimolecular reaction AB \rightarrow P (i.e., the second-step reaction). Under
	this keyword, the user should specify A in left column and Ea in right
	column; this information should be consistent with the order of the
	temperatures that one has specified under Trxnlist
rxnneg1AEa	To specify A factor and activation energy for high-pressure-limit
	reaction AB \rightarrow A + B
rxn1 rate	User specified high-pressure-limit rate constant for the forward
	bimolecular association reaction (A + B \rightarrow AB).
GeoMeFreq	The geometric mean frequency (in the unit of cm ⁻¹) of unimolecular
	intermediate AB.
paramAB	Lennard-Johns parameters for the unimolecular intermediate AB, in
	the order of σ (unit: Å), ϵ (unit: Kelvin, i.e., ϵ/k_B), molar mass (unit:
	gram per mole).
sABstar	The number of vibrational degree of freedom for AB.

4.4 Other input files

When the "FE" option is specified as "-1", the F_E factor is computed using the Whitten-Rabinovitch approximation. This is the most convenient option for computing F_E . When the F_E factor is computed by the Whitten-Rabinovitch approximation, vibinfo.inp is needed as additional input file.

In vibinfo.inp, the vibrational zero-point energy (in kcal/mol, for the unimolecular species) is specified on the first line, followed by the vibrational frequencies (in cm⁻¹).

39.76669 / ZPE in kcal/mol	
67.6166	
74.9266	
81.1074	
95.0748	
114.2145	
125.9215	
321.9349	
399.5684	
410.5671	
417.2896 438.9300	
456.7933	
400.7955 543.3220	
615.2602	
665.2141	
674.6209	
860.1783	
886.6351	
926.5675	
933.1321	
934.0968	
937.4362	
1909.0212	
2053.4916	
2114.4137	
2134.0820	
2148.7552	
2153.5027	
2153.8935	
2159.6340	

5. Test runs

We have included five test runs in

Polyrate2016-2A/util/SS-QRRK/testrun/ and the corresponding output files are in Polyrate2016-2A/util/SS-QRRK/testo/

Test run1: Unimolecular reaction, with user specified $F_{\rm E}$ =1.15

Test run2: Unimolecular reaction, with F_E being computed by Whiten-Rabinovitch

Test run3: Bimolecular reaction with an intermediate

Test run4: Unimolecular reaction, with $F_{\rm E}$ being computed from density of states

Test run5: Bimolecular association reaction