

Potential Energy Surface Transition State

Making INPUT 1

- First lines are input of method and basis set
 - #P HF/STO-3G pop=reg
- Empty line
- Title of the calculation: Anything is OK
- Empty line
- Charge and spin multiplicity: usually we consider neutral molecule so charge 0, multiplicity is number of unpaired electrons +1, usually we consider filled electron so 1

Making INPUT 2

- Then define the molecule either using **XYZ** or **Z-matrix** input

O1	0.000000	0.754227	-0.058812	O1
O2	0.000000	-0.754227	-0.058812	O2 1, RO1O2
H3	-0.742068	-1.085986	0.470499	H3, 2, RO2H3, 1, AO1O2H3
H4	0.742068	1.085986	0.470499	H4, 1, RO1H4, 2, AO2O1H4, 3, DH3O2O1H4

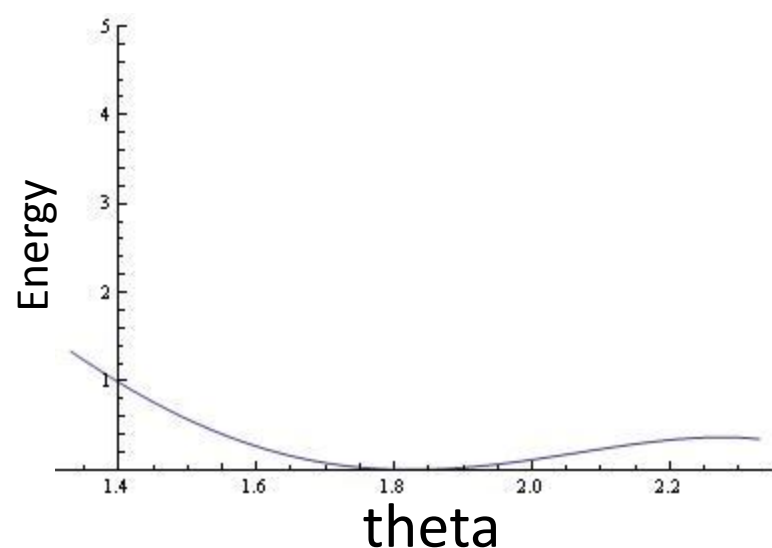
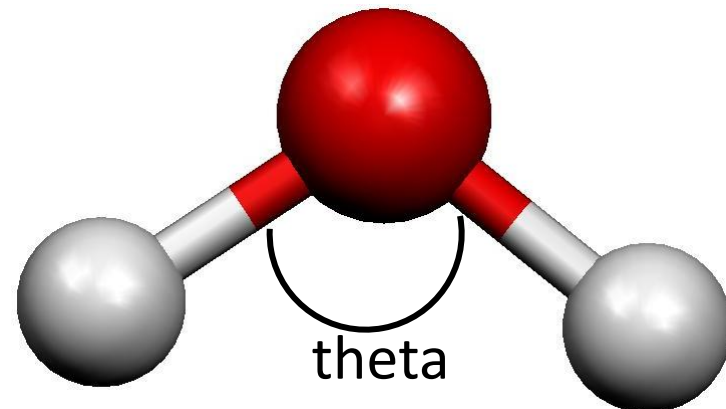
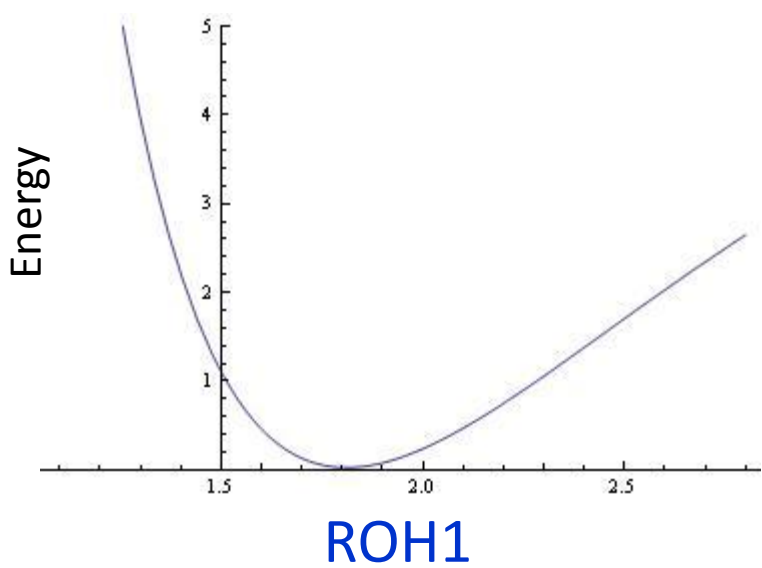
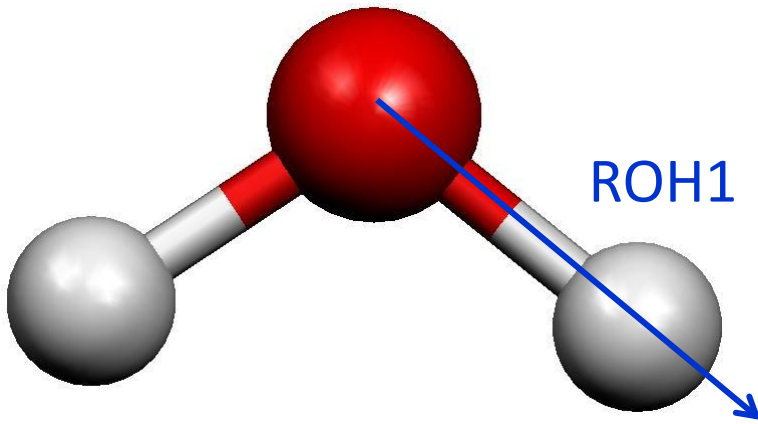
RO1O2=1.50845307
RO2H3=0.97
RO1H4=0.97
AO1O2H3=110.0
AO2O1H4=110.0
DH3O2O1H4=109.

- Then end with one blank line

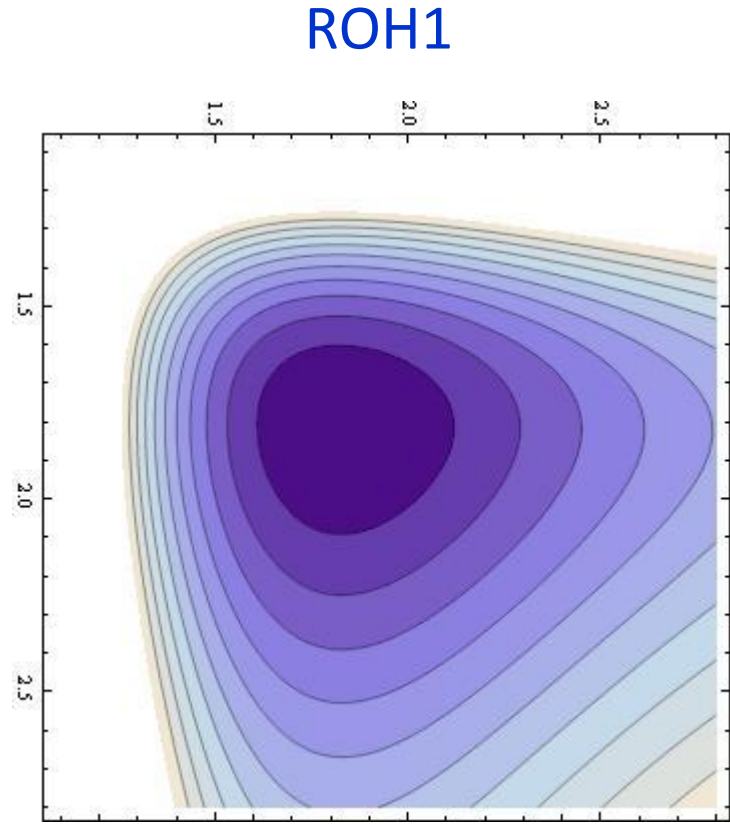
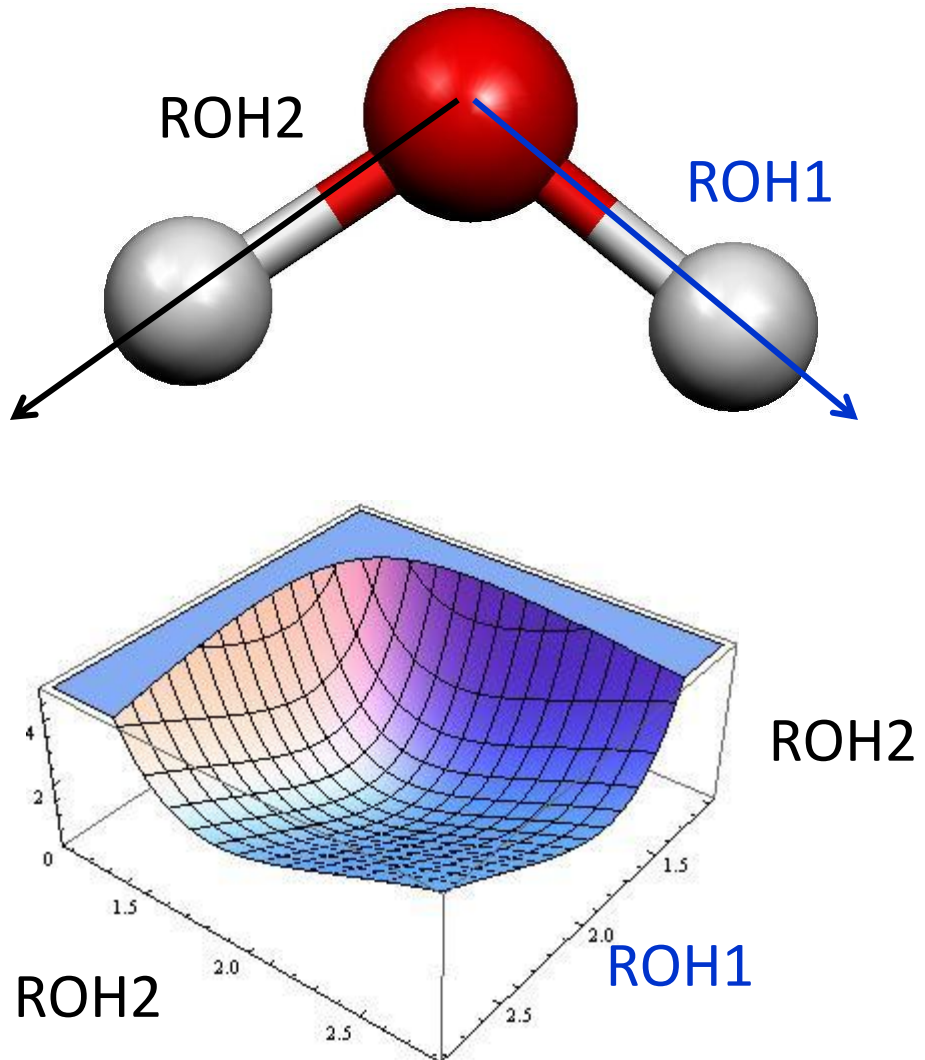
Review

- Last week we did diatomic molecules with only one bond so we only have to think about forming a bond or breaking that bond
- Today we will consider something more complex molecules
- Then we will make and break a bond at the same time REACTION

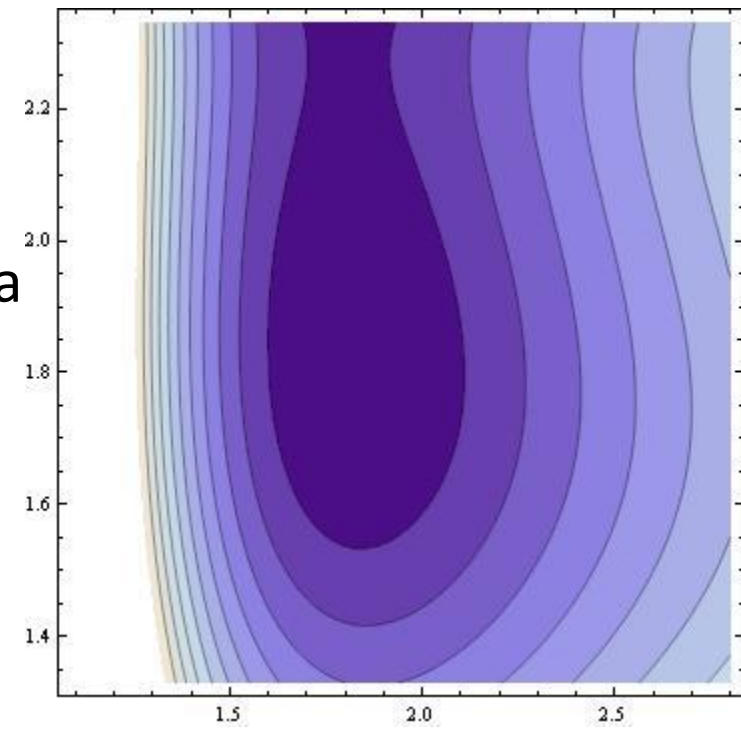
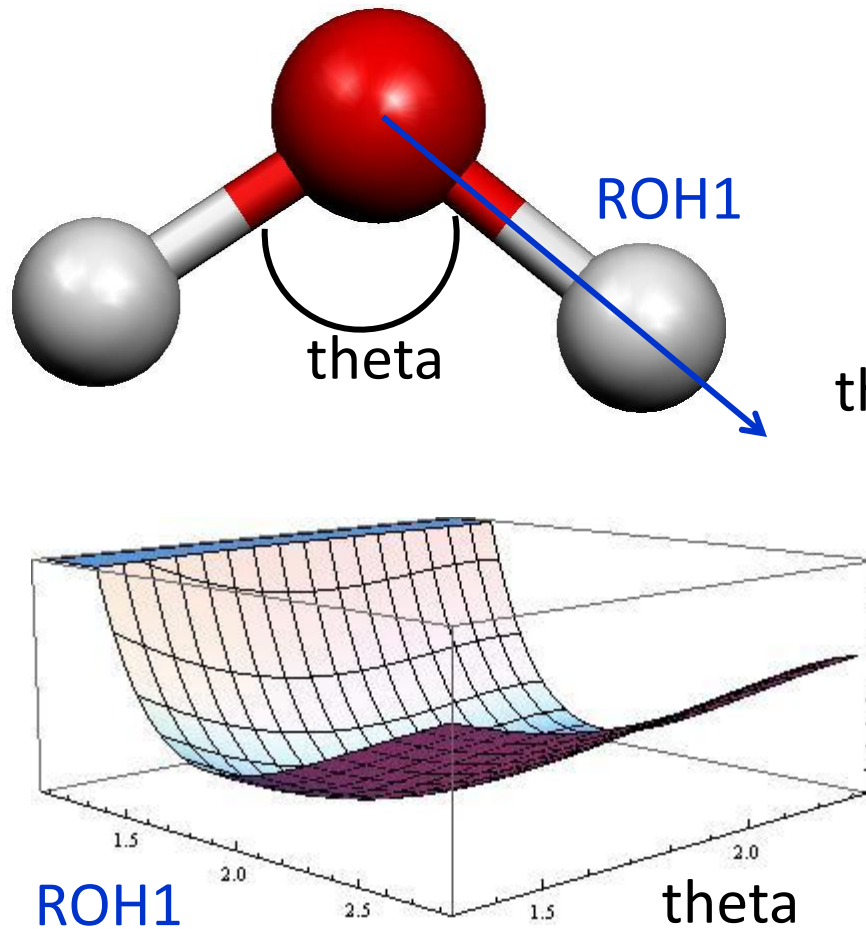
Water Molecule Potential Energy Curve



Water Molecule: Stretch Stretch Potential Energy Surface



Water Molecule: Stretch Bend Potential Energy Surface



Normal Mode Vibration

First we consider mass –weight Cartesian Coordinates

Next expand the potential energy to second term near equilibrium

$$V = V(0) + \sum_I^{3N} K_I q_I + \frac{1}{2} \sum_I^{3N} \sum_J^{3N} K_{IJ} q_I q_J = \frac{1}{2} \sum_I^{3N} \sum_J^{3N} K_{IJ} q_I q_J$$

Total energy is given as



Cross term

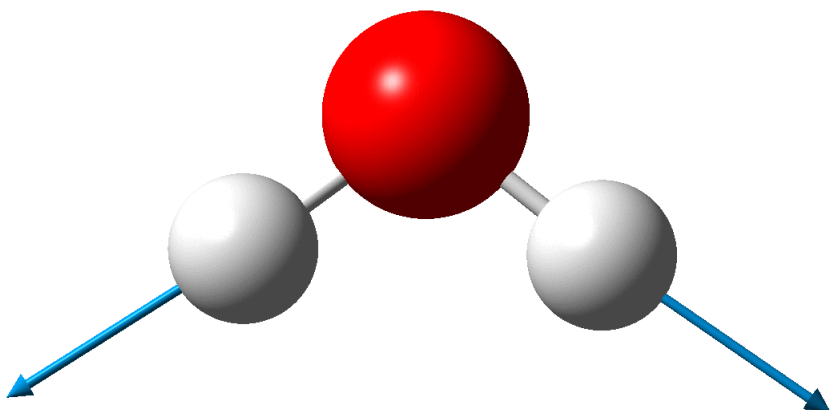
If we **diagonalize** K_{IJ}

Motion described by $3N-6$ (5 for linear molecule) **NORMAL MODES**

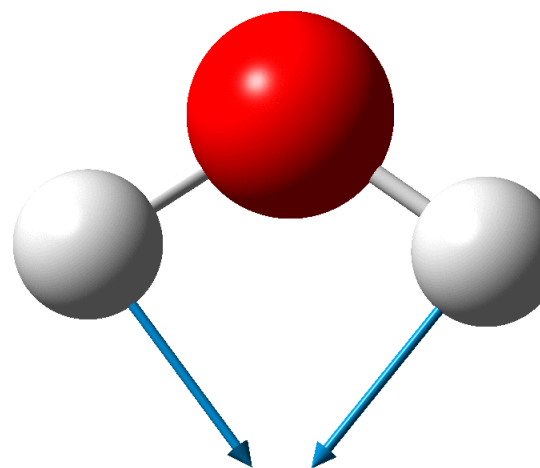
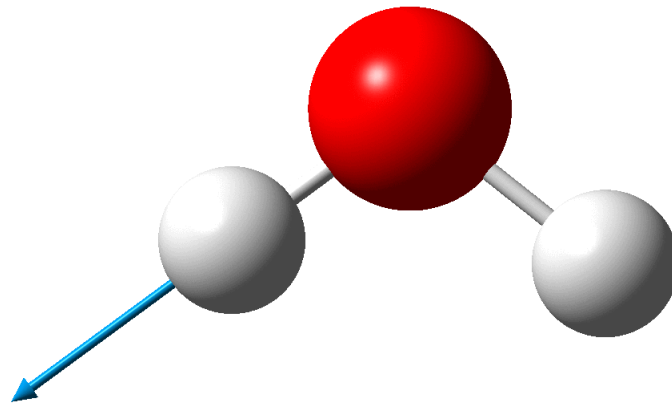
5/6 Modes that are ignored have zero λ_I TRANSLATION/ROTATION

Normal Modes of Water

Symmetric stretch



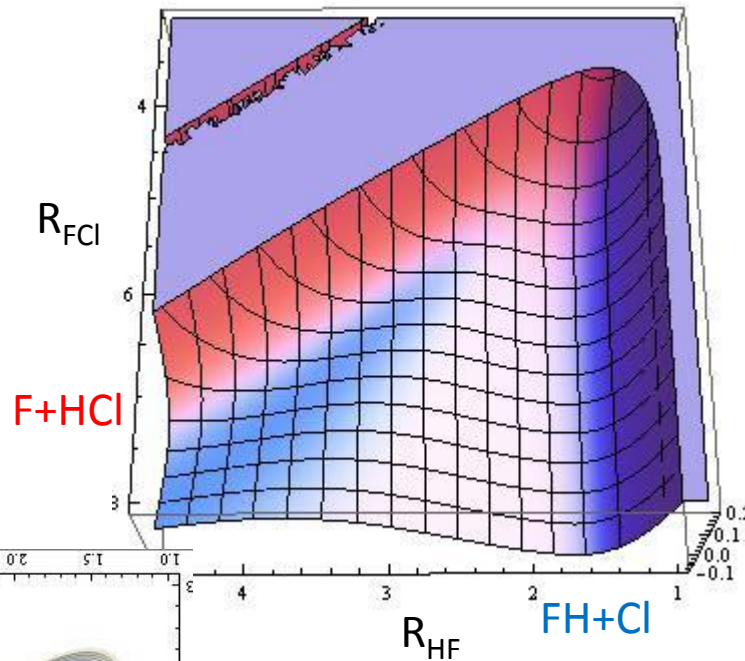
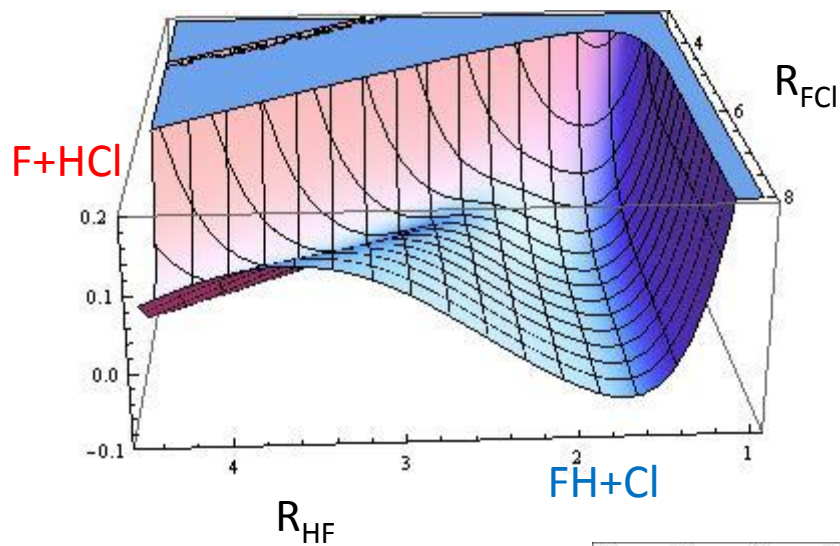
Antisymmetric stretch



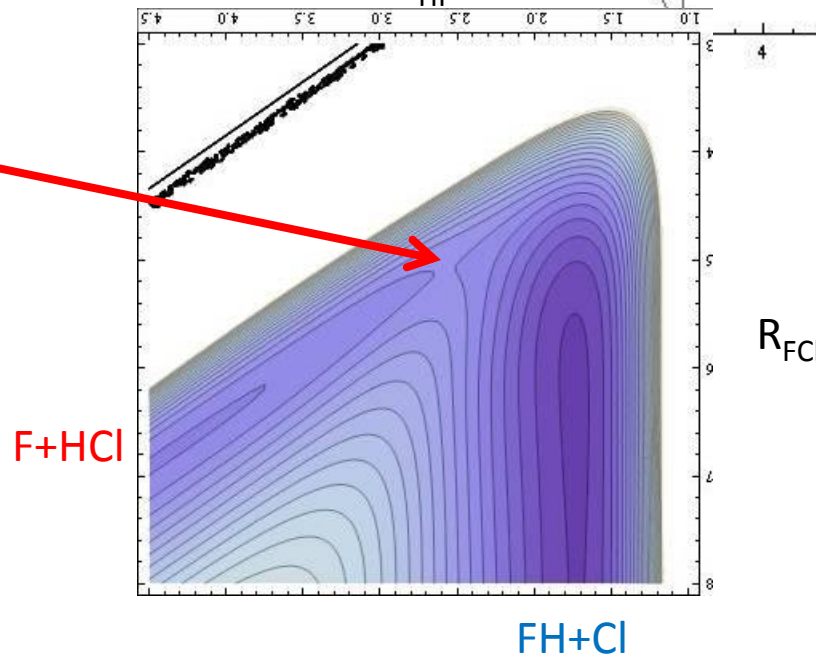
Bend

$\text{F} + \text{HCl} \rightarrow \text{FH} + \text{Cl}$ Potential Energy Surface

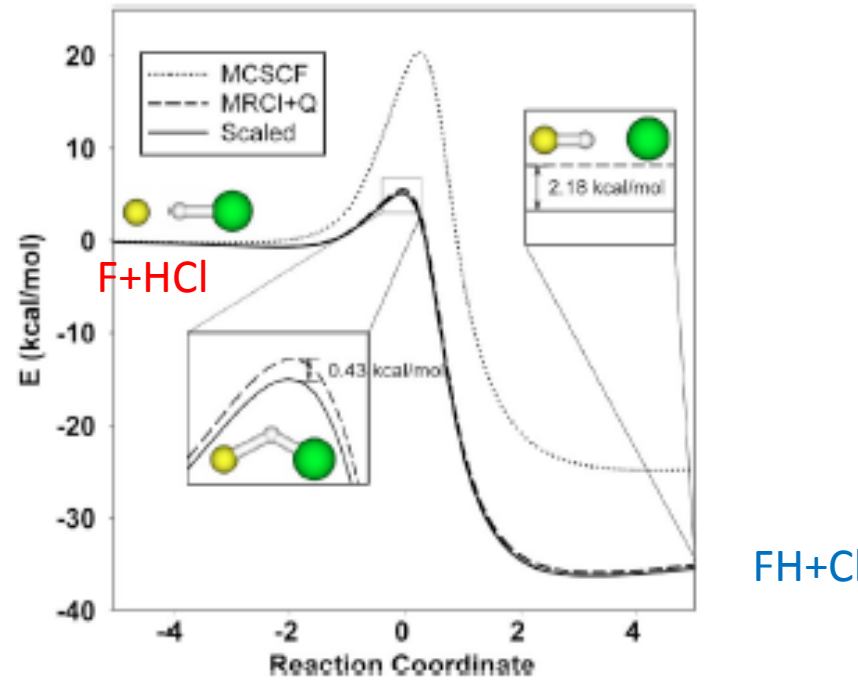
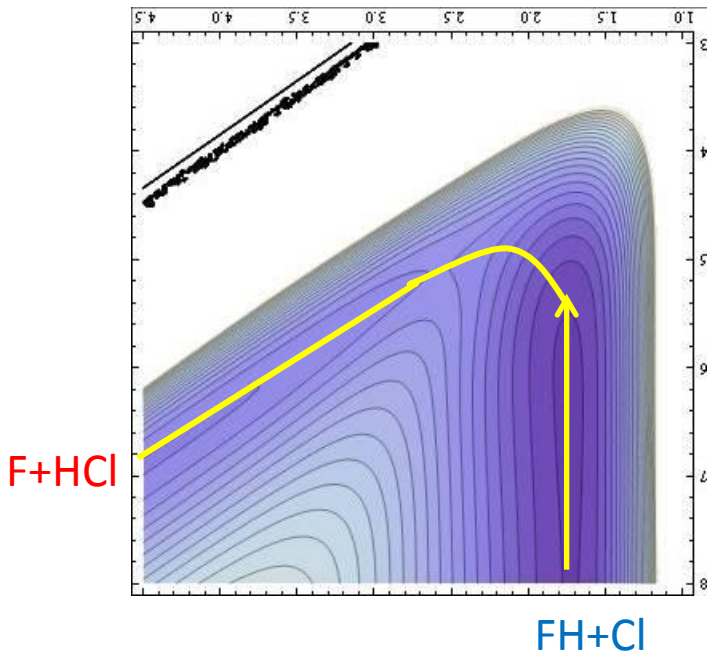
Hydrogen Abstraction Reaction, break a HCl bond form a HF bond



Potential Barrier



Potential Energy Curve along IRC

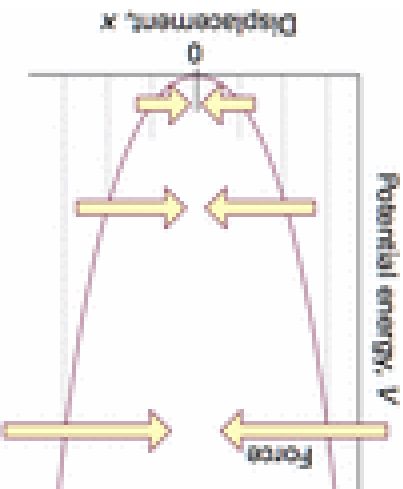


You need enough energy to go over the potential barrier to react!
The minimum energy path that connect the barrier with reactant and product is called intrinsic reaction path
For $F+HCl$ case you need about 5 kcal/mol of for reaction to occur

How to quantify Barrier?

- At the barrier the first derivative is zero and the second derivative is negative for one degree of freedom and the rest are always positive. In Gaussian you can use the freq keyword to check if there is imaginary frequency

Harmonic Oscillator approximation at barrier



$$F = m \frac{d^2x}{dt^2} = -kx \quad k: \text{force constant} \quad \omega = \sqrt{\frac{k}{m}}$$

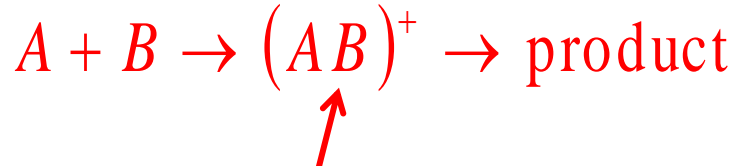
In the case of barrier the force constant k is negative so ω is imaginary

Find the potential Barrier

- Input a geometry that is close to the barrier and use the keyword Opt=(ts)
- Check if the optimized structure is a barrier by calculating the frequency by the freq keyword
- Use the IRC keyword to follow the reaction to the reactant and product keyword
IRC(forward), IRC(backward)

Transition State Theory

- Once you have the barrier you can estimate the thermal rate constant of the reaction from the calculation results! (Assuming that the reaction proceeds along the IRC)



Activated Complex
(Geomtry at Potential Barrier)

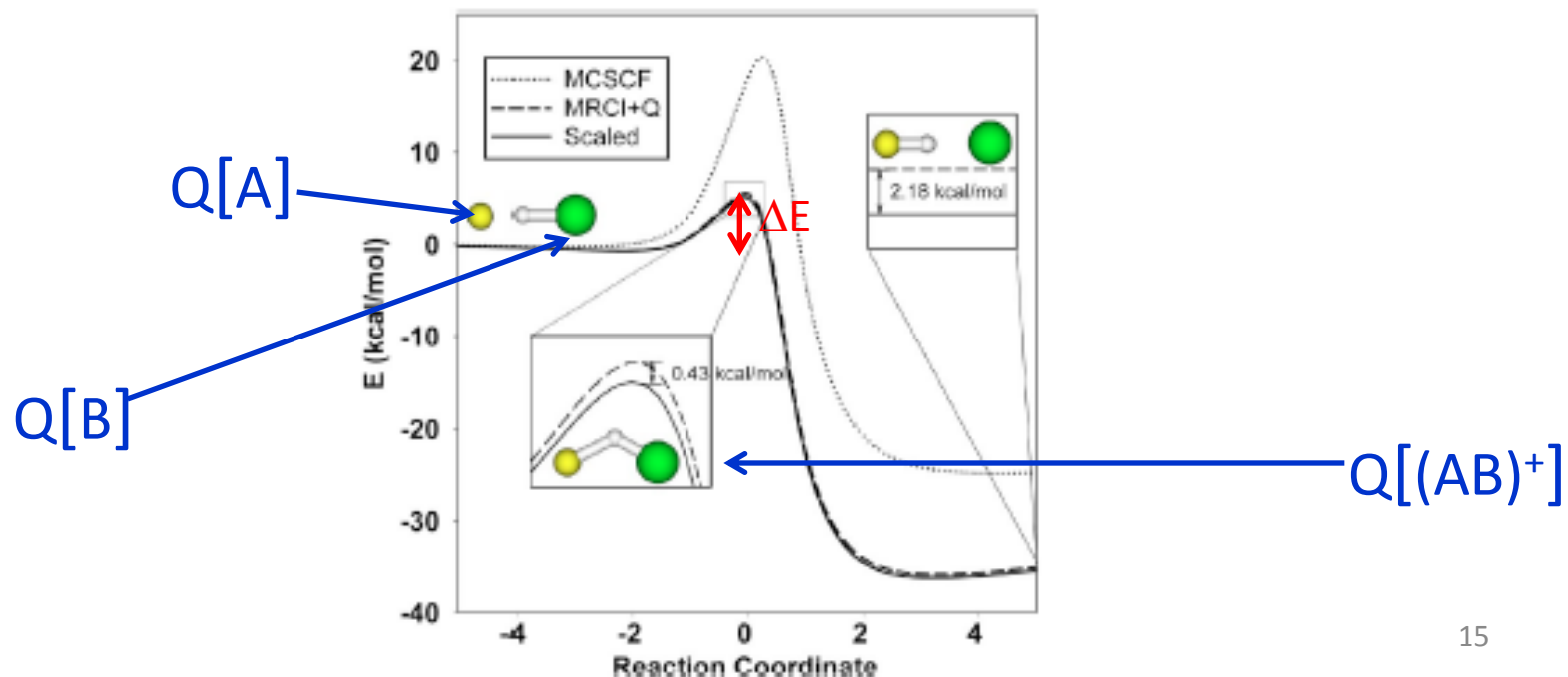
$$-\left(\frac{d[A]}{dt}\right) = \frac{k_B T}{h} K^+ [A][B]$$

K^+ is the thermal rate constant that depends on temperature

Things Needed To Calculate Rate Constant

$$K^+ = \frac{Q[(AB)^+]}{Q[A]Q[B]} \exp\left[-\frac{\Delta E}{k_B T}\right]$$

Q is the
 ΔE is the



Partition Function 1

$$Q_{trans} =$$

Usually you consider 1atm pressure

$$\begin{aligned} Q_{electronic} &= D_0 \exp[-\varepsilon_0/k_B T] + D_1 \exp[-\varepsilon_1/k_B T] \\ &\quad + D_2 \exp[-\varepsilon_2/k_B T] + \dots \\ &\rightarrow D_0 \end{aligned}$$

D

Partition Function 2

$$Q_{rot} =$$

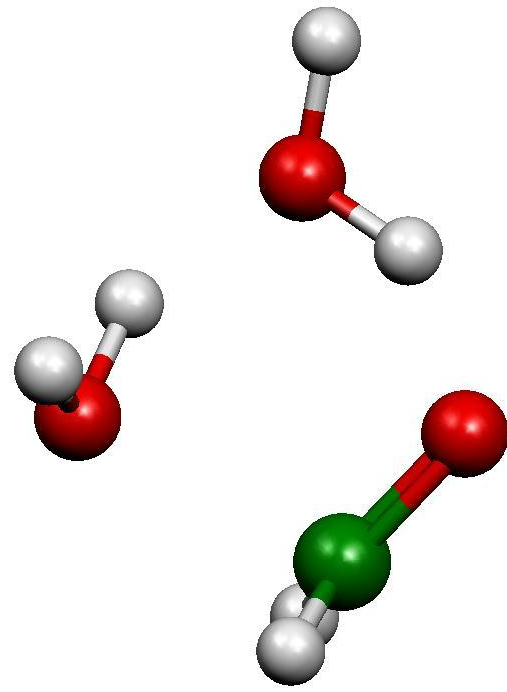
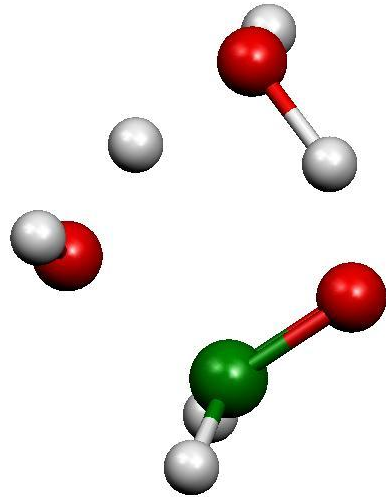
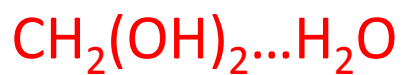
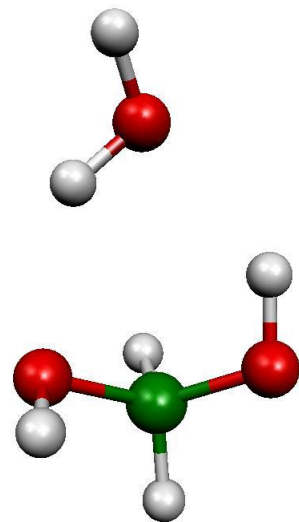
$$\theta_{A,B,C} = \frac{h^2}{8\pi^2 I_{A,B,C} k_B}$$

I is

$$Q_{vib} =$$

ν_i is

Methanediol Dehydration



Output of TS Optimization

140.109.112.238:22 - Tera Term VT

File	Edit	Setup	Control	Window	Resize	Help
DL5	0.04647	0.00000	0.00000	-0.00001	-0.00001	0.04646
DL6	2.12133	0.00000	0.00000	-0.00001	-0.00001	2.12132
	Item	Value	Threshold	Converged?		
Maximum Force		0.000000	0.000450	YES		
RMS Force		0.000000	0.000300	YES		
Maximum Displacement		0.000017	0.001800	YES		
RMS Displacement		0.000007	0.001200	YES		
Predicted change in Energy=-3.389145D-12						
Optimization completed.						
-- Stationary point found.						
<hr/>						
! Optimized Parameters !						
! (Angstroms and Degrees) !						
<hr/>						
! Name	Definition	Value	Derivative	Info.	!	!
! R1	R(1,2)	1.2942	-DE/DX =	0.0		!
! R2	R(1,3)	1.7415	-DE/DX =	0.0		!
! R3	R(1,4)	1.0985	-DE/DX =	0.0		!
! R4	R(1,5)	1.0964	-DE/DX =	0.0		!
! R5	R(2,9)	1.2826	-DE/DX =	0.0		!
! R6	R(3,6)	1.1819	-DE/DX =	0.0		!
! R7	R(3,7)	0.9694	-DE/DX =	0.0		!
! R8	R(6,8)	1.2546	-DE/DX =	0.0		!
! R9	R(8,9)	1.1596	-DE/DX =	0.0		!
! R10	R(8,10)	0.9652	-DE/DX =	0.0		!
! A1	A(2,1,3)	108.3108	-DE/DX =	0.0		!
! A2	A(2,1,4)	118.899	-DE/DX =	0.0		!
! A3	A(2,1,5)	117.0873	-DE/DX =	0.0		!

ts2.log lines 1288-1316/1913 68%

Output of TS Optimization

140.109.112.238:22 - Tera Term VT

File Edit Setup Control Window Resize Help

- Thermochemistry -

Temperature 298.150 Kelvin. Pressure 1.00000 Atm.

Atom 1 has atomic number 6 and mass 12.00000
Atom 2 has atomic number 8 and mass 15.99491
Atom 3 has atomic number 8 and mass 15.99491
Atom 4 has atomic number 1 and mass 1.00783
Atom 5 has atomic number 1 and mass 1.00783
Atom 6 has atomic number 1 and mass 1.00783
Atom 7 has atomic number 1 and mass 1.00783
Atom 8 has atomic number 8 and mass 15.99491
Atom 9 has atomic number 1 and mass 1.00783
Atom 10 has atomic number 1 and mass 1.00783

Molecular mass: 66.03169 amu.

Principal axes and moments of inertia in atomic units:

	1	2	3
Eigenvalues --	208.97954	286.92164	456.13868
X	0.69691	-0.60799	0.38035
Y	-0.56629	-0.14111	0.81204
Z	0.44004	0.78131	0.44264

This molecule is an asymmetric top.

Rotational symmetry number 1.

Rotational temperatures (Kelvin) 0.41446 0.30187 0.18988

Rotational constants (GHZ): 8.63597 6.29001 3.95656

1 imaginary frequencies ignored.

Zero-point vibrational energy 198551.1 (Joules/Mol)

Output of TS Optimization

140.109.112.238:22 - Tera Term VT

File Edit Setup Control Window Resize Help

Full mass-weighted force constant matrix:

Low frequencies ----1369.6748 -12.0457 -3.0710 0.0008 0.0014 0.0015

Low frequencies --- 15.2049 164.8661 367.0583

***** 1 imaginary frequencies (negative Signs) *****

Diagonal vibrational polarizability:

68.2855903 48.1711286 42.8890866

Harmonic frequencies (cm**-1), IR intensities (KM/Mole), Raman scattering activities (A**4/AMU), depolarization ratios for plane and unpolarized incident light, reduced masses (AMU), force constants (mDyne/Å), and normal coordinates:

	1		2		3
	A		A		A
Frequencies --	-1369.6747		164.8612		367.0574
Red. masses --	1.1492		2.2751		4.6712
Frc consts --	1.2702		0.0364		0.3708
IR Inten --	225.1406		15.4534		62.1548

Atom	AN	X	Y	Z	X	Y	Z	X	Y	Z
1	6	0.07	0.00	-0.05	-0.02	0.21	-0.01	0.07	0.04	-0.17
2	8	0.02	-0.02	0.03	-0.06	-0.15	-0.02	0.13	0.09	-0.15
3	8	-0.04	0.01	0.01	-0.02	-0.10	-0.03	-0.29	0.04	0.17
4	1	-0.02	0.01	0.00	-0.29	0.45	-0.26	-0.18	0.06	-0.13
5	1	-0.06	-0.02	0.05	0.25	0.44	0.26	0.02	0.02	-0.20
6	1	0.08	-0.41	0.56	0.00	-0.06	0.05	-0.16	-0.02	-0.13
7	1	-0.05	0.02	-0.02	0.13	-0.13	-0.23	-0.36	0.00	0.55
8	8	0.00	0.01	-0.02	0.09	0.04	0.06	0.13	-0.16	0.10
9	1	-0.56	0.40	-0.07	0.01	-0.11	-0.05	0.27	-0.15	-0.03
10	1	0.06	0.04	-0.08	0.02	0.33	0.08	0.08	0.25	0.09

4	5	6
A	A	A

Hartree Fock Projects 2

- Vinyl alcohol $\text{H}_2\text{C}=\text{CHOH}$, CH_3CHO isomer energy and barrier energy (IRC check)
- $\text{R}_1\text{R}_2\text{C}=\text{O}$ optimization effect of substituent on the $\text{C}=\text{O}$ vibrational frequency and intensity
- $\text{CO}_2 + \text{H}_2\text{O}$ carbonic acid formation barrier
- $\text{R}_1\text{R}_2\text{R}_3\text{C}-\text{CR}_4\text{R}_5\text{R}_6$ and $\text{R}_1\text{R}_2\text{C}=\text{CR}_3\text{R}_4$ torsion barrier and potential curve