4 | Characterization of stationary points and vibrational analisys

In this chapter we cover how to characterize stationary points that are found via geometry optimization. We will discuss harmonic vibrational analysis and the computation of vibrational frequencies.

4.1 Characterization of stationary points via the second derivative of the energy

One of the most important applications of quantum chemistry is to find and identify the stationary points of potential energy surfaces. Recall that a stationary point is a molecular geometry for which all first derivatives are null

$$\frac{\partial E(\mathbf{R})}{\partial \mathbf{R}_i}\bigg|_{\mathbf{R}=\mathbf{R}^*} = 0, i = 1, \dots, N_{\text{atoms}}$$
 (4.1)

Points on the PES that correspond minima are stable molecular geometries, while saddle points correspond to transition states.

How can we distinguish minima from transition states? As we discussed in the Chapter on the Born–Oppenheimer approximation, we need to know the curvature of the PES near a stationary point. It helps that near a stationary point the PES is a **quadratic function** of the nuclear coordinates. To appreciate this point consider the case of a diatomic molecule A-B with potential energy curve E(r) as a function of the bond distance r. At a stationary point r^* , we have that E'(r) = 0. Therefore, if we write the Taylor series for E(r) centered around r^* it can be simplified to

$$E(r) = E(r^*) + \underbrace{E'(r^*)(r - r^*)}_{=0} + \frac{1}{2}E''(r^*)(r - r^*)^2 + \frac{1}{6}E'''(r^*)(r - r^*)^3 + \dots$$

$$= E(r^*) + \frac{1}{2}E''(r^*)(r - r^*)^2 + \dots$$
(4.2)

In the second line we keep only the leading terms in the quantity $r - r^*$ [assuming that $E''(r^*) \neq 0$]. This shows that near the stationary point, the PEC E(r) looks like a parabola centered around r^* . The curvature of this parabola depends on $E''(r^*)$,

the **second derivative** of the PEC at the stationary point. If we compute $E''(r^*)$ then we can determine if the stationary point of a diatomic is a minimum or a transition state.

4.2 Characterization of stationary points in polyatomic molecules

For polyatomic molecules the energy is a function of all the atomic coordinates and the Taylor series near a stationary point is given by a more complex formula

$$E(\mathbf{R}) = E(\mathbf{R})^* + \frac{1}{2} \sum_{ij}^{3N_{\text{atoms}}} \left. \frac{\partial^2 E(\mathbf{R})}{\partial R_i \partial R_j} \right|_{\mathbf{R} = \mathbf{R}^*} (R_i - R_i^*) (R_j - R_j^*) + \dots$$
 (4.3)

In writing this equation, we use the symbols R_i and R_j to indicate the x,y,z coordinates of all atoms, which in total are $3N_{\rm atoms}$. This expression is still a quadratic function in the displacements $R_i - R_i^*$ and $R_j - R_j^*$, but the second derivative is replaced with the **Hessian matrix**, the matrix of mixed second derivatives

$$H_{ij} = \left. \frac{\partial^2 E(\mathbf{R})}{\partial R_i \partial R_j} \right|_{\mathbf{R} = \mathbf{R}^*} \tag{4.4}$$

To characterize the nature of a stationary point it is necessary to consider the eigenvalues of the Hessian matrix at the stationary point. If none of the eigenvalues is equal to zero, we can distinguish three cases:

- If all the eigenvalues are positive, then a stationary point is a local minimum.
- If some eigenvalues are positive and some are negative, then a stationary point is a saddle point. Transition states are characterized by only one negative eigenvalue.
- If the Hessian has only negative eigenvalues then the energy is maximum.

4.3 Vibrational frequencies

Once we compute the Hessian matrix, it is possible to compute approximate harmonic vibrational frequencies. Vibrations in a molecule are due to the motion of the nuclei. So to obtain information about vibrations we have to solve the nuclear Schrödinger equation assuming the Born-Oppenheimer approximation

$$[\hat{T}_{n} + E(\mathbf{R})]\Psi_{\nu}(\mathbf{R}) = E_{\nu}\Psi_{\nu}(\mathbf{R}) \tag{4.5}$$

This equation describes the quantum mechanical states of nuclei that move on the potential energy surface $E(\mathbf{R})$. The electrons never appear explicitly in this equation, but their effect is implicitly included in the PES (which we get from the electronic Schrödinger equation).

Solving the nuclear Schrödinger equation near a stationary point is particularly simple because the potential is quadratic

$$E(\mathbf{R}) = E(\mathbf{R})^* + \frac{1}{2} \sum_{ij} H_{ij} (R_i - R_i^*) (R_j - R_j^*)$$
 (4.6)

For a diatomic molecule, the Hamiltonian is equivalent to that of a **quantum** harmonic oscillator. This can be more easily seen if we change coordinate from r to $x = r - r^*$

$$\hat{T}_{n} + E(\mathbf{R}) = -\frac{1}{2\mu} \frac{d^{2}}{dr^{2}} + \underbrace{E(r^{*})}_{V_{0}} + \frac{1}{2} E''(r^{*}) (r - r^{*})^{2}$$

$$= \underbrace{\left(-\frac{1}{2\mu} \frac{d^{2}}{dx^{2}} + \frac{1}{2} kx^{2}\right)}_{\text{barrack extilator}} + V_{0}$$
(4.7)

where V_0 is the energy at the bottom of the well (the energy you get when you optimize the geometry of the diatomic molecule), $\mu = (M_A + M_B)/M_A M_B$ is the reduced mass of the diatomic molecule, and $k = E''(r^*)$ is the force constant.

The quantum harmonic oscillator is one of those few problems for which we know analytical solutions to the Schrödinger equation. The eigenvalues of the quantum harmonic oscillator are given by (in atomic units)

$$E_{\nu} = V_0 + \sqrt{\frac{k}{\mu}} \left(\nu + \frac{1}{2} \right) = V_0 + \omega \left(\nu + \frac{1}{2} \right), \quad \nu = 0, 1, 2, \dots$$
 (4.8)

where ω is the angular frequency of the oscillator and v is the vibrational quantum number. The ground state of the harmonic oscillator (the minimum energy) is obtained for v=0 and it is equal to

$$E_0 = V_0 + \frac{1}{2}\omega {4.9}$$

Note that this quantity is always **higher** than the energy at the bottom of the potential (V_0) by an amount equal to $\frac{1}{2}\omega!$ This amount of energy is called the **zero-point energy**, and it is due to the quantum nature of vibrations.

For polynuclear molecules, the solution to the nuclear Schrödinger equation can be found by diagonalization of the mass-weighted Hessian matrix

$$\tilde{H}_{ij} = \frac{H_{ij}}{\sqrt{M_i M_j}} \tag{4.10}$$

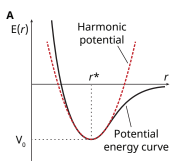
Diagonalizing the mass-weighted Hessian matrix means finding a set of eigenvectors $\mathbf{u}^{(a)}$ and corresponding eigenvalues λ_a that satisfy the equation

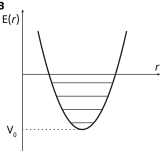
$$\sum_{i} \tilde{H}_{ij} u_{j}^{(\alpha)} = \lambda_{\alpha} u_{i}^{(\alpha)} \tag{4.11}$$

Each eigenvector/eigenvalue pair corresponds to a molecular vibrational mode α , which we call a **normal vibrational mode**. Each mode behaves as an independent harmonic oscillator with angular frequency

$$\omega_{\alpha} = \sqrt{\lambda_{\alpha}} \tag{4.12}$$

and has an associated quantum number $v_{\alpha} = 0, 1, 2, \dots$





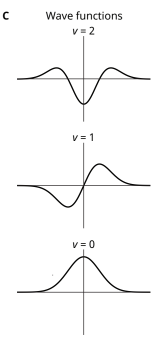
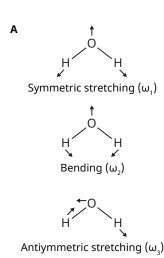


Fig. 4.1: The potential energy surface of a diatomic molecule near the equilibrium bond distance (r_{eq}) can be approximated by a quadratic function (parabola, see **A**). **B** and **C** show the energy levels and wave functions of the diatomic molecule vibrating with a quadratic potential.



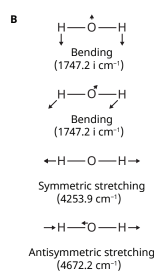


Fig. 4.2: Normal modes of water at the equilibrium geometry (**A**) and at the linear transition state (**B**) computed with Hartree–Fock theory and the def2-TZVP basis.

Once the mass-weighted Hessian matrix is diagonalized, the eigenvalues of the nuclear Schödinger equation (nuclear energy levels) are given by

$$E_{\nu} = V_0 + \sum_{V_{\alpha}} \omega_{\alpha} \left(\frac{1}{2} + V_{\alpha} \right) \tag{4.13}$$

where the quantities v_{α} are the quantum numbers for each vibrational mode. For a general molecule, the zero-point energy is given by

$$E_0 = V_0 + \frac{1}{2} \sum_{\nu_a} \omega_a \tag{4.14}$$

which again we see is higher than the energy at the bottom of the well.

The harmonic vibrational analysis introduced here can be performed with the majority of quantum chemistry codes. There are three main applications of a vibrational analysis

- The **sign** of the eigenvalues λ_a tell us information about the curvature of the PES at the stationary point. If all the λ_a are positive the stationary point is a minimum. **This corresponds to the case where all the frequencies are real**, since $\omega_a = \sqrt{\lambda_a}$. The presence of one or more **negative** eigenvalues indicates that there is a vibrational mode with a negative curvature. In this case a stationary point is a transition state. This case corresponds to finding an imaginary frequency since if an eigenvalue λ_a is negative, its square root will be an imaginary number, $\omega_a = i\sqrt{|\lambda_a|}$.
- The value of the individual vibrational frequencies ω_a approximate the vibrational frequencies measured in IR absorption spectroscopy. The eigenvectors of the mass-weighted Hessian tell us how the atoms are displaced in a vibrational mode, and can therefore tell us if a given mode is a bond stretching mode, a bending mode, etc.
- The sum of the vibrational frequencies tell us information about the zeropoint energy of a molecule. Accurate thermodynamic computations must include the zero-point energy to include nuclear vibrational effects and ultimately achieve good agreement with experiment. We will come back to this topic later.

The example below shows the vibrational analysis of the water molecule using Hartree–Fock and the def2-TZVP basis. The vibrational analysis shows that six modes are classified as "low-frequency" because they are close to zero (post-proj low-frequency mode). Some of these numbers are imaginary, and that's due to small numerical noise. These six low-frequency modes are trivial modes corresponding to three translations (along the x, y, or z direction) and three rotations (around the x, y, or z axis), hence the label "TR".

Modes 7, 8, and 9 corresponds to three molecular vibrational modes. Starting from the highest frequency mode, we have

• Mode 9 (ω_3 = 4213.5 cm⁻¹), an antisymmetric stretching of the two O–H bonds.

- Mode 8 (ω_1 = 4111.5 cm⁻¹), a symmetric stretching of the two O–H bonds.
- Mode 7 ($\omega_2 = 1734.7 \text{ cm}^{-1}$), a H-O-H bending mode.

Each normal mode has also an associated symmetry. The A_1 modes correspond to **totally symmetric modes**. This means that if we transform the modes according to any of the symmetries of water, the normal mode does not change. Mode 9 is of type B_2 , which means that if we rotate the molecule by 180° along the vertical axis that includes the oxygen atom, then the normal mode changes sign.

The calculation also tells us the IR activity (measured in km/mol) of each normal mode. This information allows us to predict or relate our results to an experimental spectrum. All three normal modes are active, with mode 7 and 9 being the most intense.

```
==> Harmonic Vibrational Analysis <==
post-proj low-frequency mode: 0.0001i [cm^-1] (TR)
post-proj low-frequency mode: 0.0001i [cm^-1] (TR)
post-proj low-frequency mode: 0.0000i [cm^-1] (TR)
post-proj low-frequency mode: 0.0001 [cm^-1] (TR)
Vibration
                                  7
                                                       8
                                                                             9
                          1734.7180
                                                                       4213.5468
Freq [cm^-1]
                                                 4111.5086
Irrep
                             A1
                                                    A1
                                                                            B2
Reduced mass [u] 1.0835

Force const [mDyne/A] 1.9210

Turning point v=0 [a0] 0.2531

RMS dev v=0 [a0 u^1/2] 0.1863
Reduced mass [u]
                              1.0835
                                                   1.0444
                                                                          1.0839
                                                  10.4022
                                                                         11.3375
                                                   0.1674
                                                                          0.1624
                                                    0.1210
                                                                          0.1195
                             101.9411
IR activ [km/mol]
                                                   17.0610
                                                                          76.6343
                              2495.8730
                                                                         6062.3558
Char temp [K]
                                                   5915.5454
               1 0
    2 H
    3 H
                         -0.00 -0.42 0.56 -0.00 -0.59 -0.39 0.00 0.57 0.42
```

The information listed at the bottom shows how each atom moves in each of the normal modes. For each atom, each triplet of numbers under a normal mode indicate how much is the atom displaced in the *x*, *y*, and *z* directions. This information can be used to analyze the nature of the normal modes in a molecule.

If we now repeat the same computation for linear water (after optimizing the geometry) we find seven low-frequency modes, two of which have a large imaginary frequency, $\omega = 1747.2$ i cm $^{-1}$. These results tell us that linear water corresponds to a transition state. The two imaginary modes, are the two orthogonal bending mode that transform linear H-O-H into the bent ground state of H $_2O$.

6 CHAPTER 4. CHARACTERIZATION OF STATIONARY POINTS AND VIBRATIONAL ANALISYS

==> Harmonic Vibrational Analysis <==

•	•	•	

	<pre>post-proj low-frequency</pre>	mode:	1747.1869i	[cm^-1] (V)	
	<pre>post-proj low-frequency</pre>	mode:	1747.1869i	[cm^-1] (V)	
	<pre>post-proj low-frequency</pre>	mode:	0.0000i	[cm^-1] (TR)	
	<pre>post-proj low-frequency</pre>	mode:	0.0000	[cm^-1] (TR)	
	post-proj low-frequency	mode:	0.0000	[cm^-1] (TR)	
	post-proj low-frequency	mode:	0.0000	[cm^-1] (TR)	
	post-proj low-frequency	mode:	0.0000	[cm^-1] (TR)	
	Vibration	1		2	
	Freq [cm^-1]	1747.1869i		1747.1869i	
	Irrep				
	Reduced mass [u]	1.1259		1.1259	
	Force const [mDyne/A]	-2.0250		-2.0250	
	Turning point v=0 [a0]	0.0000		0.0000	
	RMS dev v=0 [a0 $u^1/2$]	0.0000		0.0000	
<pre>IR activ [km/mol]</pre>		551.4304		551.4304	
Char temp [K]		0.0000		0.0000	
	Vibration	8		9	
Freq [cm^-1]		4253.9467		4672.2019	
	Irrep		Ag	B1u	
	Reduced mass [u]		1.0078	1.1259	
Force const [mDyne/A]		10.7453		14.4807	
Turning point v=0 [a0]		0.1676		0.1513	
	RMS dev v=0 [a0 $u^1/2$]		0.1190	0.1135	
	<pre>IR activ [km/mol]</pre>		0.0000	829.5184	
	Char temp [K]	6:	120.4822	6722.2582	