## Programming Project 1: Frequencies

# Computing frequencies and normal modes from the Hessian matrix

Write a script to read in a Hessian matrix, compute the corresponding frequencies and normal modes, and print them to a file in Jmol's .xyz format. You should import your Molecule class from Project 0 and make use of it in this script.

#### Extra Files

 $filename \qquad \qquad description$ 

molecule.xyz sample .xyz file for  $H_2O$ 

hessian.dat sample Hessian matrix for H<sub>2</sub>O

## **Equations**

Let N be the number of atoms and let  $m_A$  and  $(x_A, y_A, z_A)$  be the mass and Cartesian coordinates of the  $A^{\text{th}}$  atom.

$$(\mathbf{H})_{AB} = \frac{\partial^2 E}{\partial X_A \partial X_B} \qquad (X_{3A-2}, X_{3A-1}, X_{3A-0}) = (x_A, y_A, z_A) \qquad \text{for } A \in \{1, \dots, N\}$$
 (1)

$$(\tilde{\mathbf{H}})_{AB} = \frac{(\mathbf{H})_{AB}}{\sqrt{M_A M_B}} \qquad (M_{3A-2}, M_{3A-1}, M_{3A-0}) = (m_A, m_A, m_A) \qquad \text{for } A \in \{1, \dots, N\}$$
 (2)

$$\tilde{\mathbf{H}}\,\tilde{\mathbf{q}}_A = k_A\tilde{\mathbf{q}}_A \tag{3}$$

$$(\mathbf{q}_A)_B = \frac{(\tilde{\mathbf{q}}_A)_B}{\sqrt{M_B}} \qquad \text{for } A, B \in \{1, \dots, N\}$$

$$k_A \left[ \frac{E_h \operatorname{rad}^2}{a_0^2 u} \right] = k_A \left( \frac{E_h[J]}{a_0[m]^2 u[kg]} \right) \left[ \frac{\operatorname{rad}^2}{\operatorname{s}^2} \right] \qquad \nu_A \left[ \operatorname{Hz} \right] = \frac{1}{2\pi} \left[ \frac{\operatorname{cyc}}{\operatorname{rad}} \right] \cdot \sqrt{k_A \left[ \frac{\operatorname{rad}^2}{\operatorname{s}^2} \right]} \qquad \tilde{\nu}_A \left[ \operatorname{cm}^{-1} \right] = \frac{\nu_A \left[ \operatorname{Hz} \right]}{c \left[ \operatorname{cm/s} \right]}$$
(5)

## Procedure

- 1. read in the Hessian matrix (defined in equation 1) from hessian.dat
- 2. build a Molecule object from molecule.xyz
- 3. build the mass-weighted Hessian matrix (equation 2)
- 4. compute the eigenvalues  $(k_A)$  and eigenvectors  $(\tilde{q}_A)$  of the mass-weighted Hessian matrix (equation 3)
- 5. un-mass-weight the eigenvectors (equation 4) to get normal coordinates
- 6. determine the spatial frequencies  $\tilde{\nu}_A$  in cm<sup>-1</sup> from your force constants  $k_A$ , which are in atomic units (equation 5)
- 7. write the normal modes to a file in .xyz format, with frequencies neatly displayed in the comment lines

## Description of .xyz format for normal modes

The file format for visualizing a single normal mode in Jmol is

which amounts to a standard .xyz geometry file (distance units Å) with the displacement vector for each atom printed next to its Cartesian coordinates. For visualizing multiple motions, this format is simply repeated with an intervening empty line.

```
COMMENT LINE 1
            dx11 dy11 dz11
A1 x1 y1 z1
A2 x2 y2 z2 dx12 dy12 dz12
AN xN yN zN dx1N dy1N dz1N
COMMENT LINE 2
A1 x1 y1 z1
             dx21 dy21 dz21
A2 x2 y2 z2
             dx22 dy22 dz22
. . .
AN xN yN zN dx2N dy2N dz2N
. . .
COMMENT LINE N
A1 x1 y1 z1
             dxN1 dyN1 dzN1
A2 x2 y2 z2
             dxN2 dyN2 dzN2
AN xN yN zN
             dxNN dyNN dzNN
```