# 1 Forces and Interactions

November 15, 2023

```
[7]: import numpy as np
     import matplotlib.pyplot as plt
     from numpy.linalg import norm
     from scipy.constants import c,epsilon_0,e,physical_constants
     %config InlineBackend.figure format = 'retina'
     # the lines below set a number of parameters for plotting, such as label font \Box
      ⇔size,
     # title font size, which you may find useful
     plt.rcParams.update({'font.size': 14,
                           'font.family': 'sans-serif',
                           'axes.titlesize': 16,
                           'axes.labelsize': 18,
                           'axes.labelpad': 14,
                           'lines.linewidth': 1,
                           'lines.markersize': 10,
                           'xtick.labelsize' : 18,
                           'ytick.labelsize' : 18,
                           'xtick.top' : True,
                           'xtick.direction' : 'in',
                           'ytick.right' : True,
                           'ytick.direction' : 'in',})
```

## 0.0.1 Interactions involving polar molecules

Many molecules exhibit a dipole or even higher moments due to the fact that the charges are not evenly distributed over the molecular structure. Some of the atoms exhibit a stronger tendency to accept charges than others. This is typically measured by electronegativity and provides an idea of whether atoms rather donate or accept a charge when binding to other atoms. While homo-atomic bonds therefore do not have dipole moments, hetero-atomic bonds do (see table).

Bond	Dipole moment [D	]
C-C	0	
C-N	0.22	
О-Н	1.51	
F-H	1.94	
N=O	2.0	

Molecule	Dipole moment [D]
hexane	0
water	1.85
ethanol	1.7
acetone	2.9

The dipole moment of a molecule is measured by the displacement of two charges  $\pm q$  from each other:

$$\vec{u} = q\vec{l}.\tag{1}$$

Its direction is from the negative to the positive side. It creates an electric field that is given by

$$\vec{E} = \frac{3(\vec{u} \cdot \hat{r})\hat{r} - \vec{u}}{4\pi\epsilon_0 \epsilon r^3},\tag{2}$$

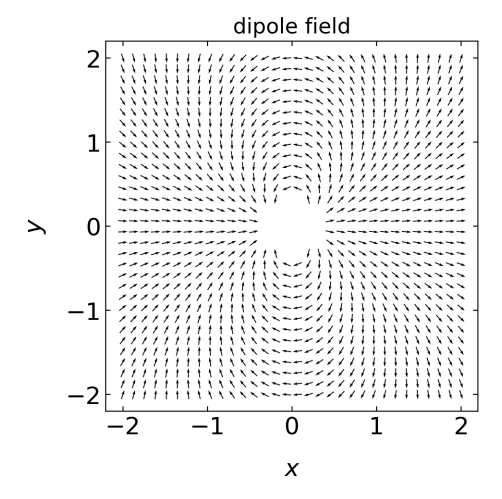
where  $\hat{r} = \vec{r}/|r|$ . The dipole self-energy, i.e., the energy to create the dipole in a solvent is given by

$$\mu^{0} = \frac{1}{4\pi\epsilon_{0}\epsilon} \left[ \frac{q^{2}}{2a} + \frac{q^{2}}{2a} - \frac{q^{2}}{l} \right]. \tag{3}$$

For l=2a this results in  $\mu^0=q^2/(8\pi\epsilon_0\epsilon a)=u^2/(4\pi\epsilon_0\epsilon l^3)$  and thus yields a similar dependence of the chemical potential on the dielectric function  $\epsilon$  as in the case of a single charge. The result is a similar dependence of the solubility on the dielectric function.

The plot below shows the electric field of a dipole:

```
Ex.flat[E.flat[:]>E_max]=np.nan
Ey.flat[E.flat[:]>E_max]=np.nan
# plot vecor field
plt.quiver(x, y, Ex/E, Ey/E, pivot='middle', headwidth=3, headlength=5,scale=40)
plt.xlabel('$x$')
plt.ylabel('$y$')
plt.title('dipole field')
plt.show()
```



**Ion–dipole interaction** The interaction energy of a dipole with a charge can be calculated by

$$w(r) = -\frac{qQ}{4\pi\epsilon_0\epsilon} \left[ \frac{1}{r - \frac{1}{2}l\cos(\theta)} - \frac{1}{r + \frac{1}{2}l\cos(\theta)} \right] = -\frac{qQ}{4\pi\epsilon_0\epsilon r^2}\cos(\theta) = -uE\cos(\theta), \tag{4}$$

where the last two equations are assuming that the distance between both objects r is much larger than the extent of the dipole l itself. From the last equation, we see that the interaction can be either

attractive or repulsive. An angle  $\theta=0^{\circ}$  results in an attractive interaction, while  $\theta=180^{\circ}$  yields repulsive interaction. Using a single charge (e.g., an Na<sup>+</sup> ion) and a dipole of  $u=1.85\,\mathrm{D}$  (water molecule) results in an interaction energy of about  $39\,k_{\mathrm{B}}T$ . Ions align and bind polar molecules like water, for example. The alignment is caused by the torque  $\tau=\vec{u}\times\vec{E}$ . For arbitrary polar molecules this is called **solvation**, while for water the term **hydration** is used. The strength of the hydration can effect the mobility of ions in solution as it makes them effective charges and is of interest, for example, in the study of ion transport through ion channels, as this requires the stripping of the hydration shell.

## **Example: Sodium Ion Hydration**

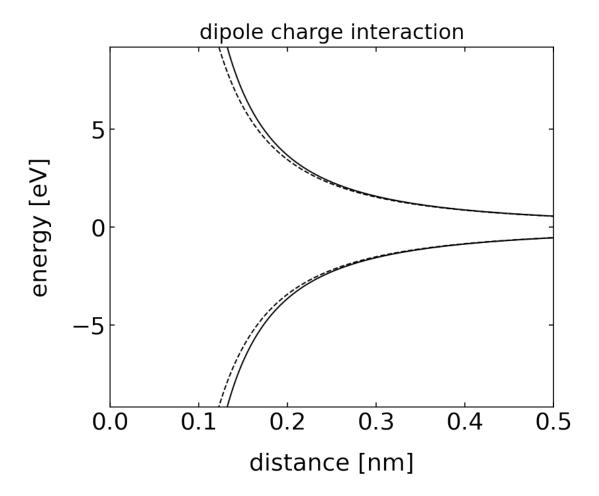
The plot below shows the ion dipole interaction for a sodium ion and a water molecule as a function of distance.

```
[]: def ion_dipole(u,r,l,theta,epsilon):
    q=u/l
    f=e*q/(4*np.pi*epsilon*epsilon_0)
    return(-f*(1/(r-0.5*l*np.cos(theta))-1/(r+0.5*l*np.cos(theta))))
[]: D=physical_constants["atomic unit of electric dipole mom."][0]

J2eV=physical_constants["electron volt-joule relationship"][0]
```

```
[]: r=np.linspace(0.1e-9,0.5e-9,200)
l=0.02e-9
```

```
[]: plt.figure(figsize=(6,5))
    plt.plot(r*1e9,ion_dipole(1.8*D,r,1,0,1)/J2eV,'k--')
    plt.plot(r*1e9,ion_dipole(1.8*D,r,1,np.pi,1)/J2eV,'k--')
    plt.plot(r*1e9,ion_dipole(1.8*D,r,1*5,0,1)/J2eV,'k')
    plt.plot(r*1e9,ion_dipole(1.8*D,r,1*5,np.pi,1)/J2eV,'k')
    plt.ylim(-9.2,9.2)
    plt.xlim(0,0.5)
    plt.xlabel("distance [nm]")
    plt.ylabel("energy [eV]")
    plt.title("dipole charge interaction")
    plt.show()
```

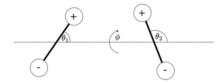


The table below shows some selected hydration properties of ions. The hydrated radius determines the diffusion of the ion in water. The hydration number is the number of orientationally bound water molecules. Typically, smaller ions have a larger hydration number.

Ion	Bare Ion radius (nm)	Hydrated radius (nm)	Hydration number
Na+	0.095	0.36	4
Mg2+	0.065	0.43	6
Cl-	0.181	0.33	1
OH-	0.176	0.3	3

## 0.0.2 Dipole-Dipole Interactions

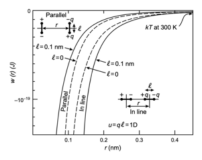
The interaction of two dipole will depend on three angles, the two angles of the dipoles with the connecting axis  $\theta_1$ ,  $\theta_2$  and the angle  $\phi$  between the two planes which contain the individual dipoles and the axis. The figure below shows the corresponding geometry.



The energy of the two dipoles at a distance r follows then without a detailed calculation with

$$w(r,\theta_2,\theta_2,\phi) = -\frac{u_1u_2}{4\pi\epsilon_0\epsilon r^3}\left[2\cos(\theta_1)\cos(\theta_2) - \sin(\theta_1)\sin(\theta_2)\cos(\phi)\right] \tag{5}$$

The image below (taken from the book by Israelachvili) shows the dipole dipole interaction energy for the **parallel** and the **in-line** configuration as indicated. As shown, the in-line configuration with facing opposite charges is more favorable.



The energies that are typically found are also larger than the thermal energy at room temperature and thefore lead to ordering effects.

## 0.0.3 Rotating Dipoles, angle averaged Potential

The equations we derived so far deliver the energy for a specific fixed orientation of dipoles for example. However, molecules with dipoles may rotate and undergo rotational Brownian motion driven by thermal energy. This rotational diffusion can be very fast and the rotational sampling creates an average interaction that may scale differently with the distance. To get an effective distance dependence in the interaction, we have to integrate the Boltzmann factor over the orientational degrees of freedoms, i.e.

$$\exp\left(\frac{w(r)}{k_BT}\right) = \frac{\int \exp\left(-\frac{w(r,\Omega)}{k_BT}d\Omega\right)}{\int d\Omega} = \left\langle \exp\left(-\frac{w(r,\Omega)}{k_BT}d\Omega\right)\right\rangle \tag{6}$$

with

$$d\Omega = \sin(\theta)d\theta d\phi$$

and

$$\int d\Omega = 4\pi$$