Tutorial 10: Reaction Ensemble and Constant pH ensemble

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Contents

1	Introduction	1
2	Theoretical Background 2.1 Constant pH method	3
3	First part: homogeneous aqueous solution of acidic species 3.1 System Setup	4 4
4	Second part: linear weak polyelectrolyes 4.1 Titration curves	7
5	Distribution of charges onto the chain	11

1 Introduction

This tutorial introduces the basic features for simulating tritatable systems via two different methods, the Constant pH method and the Reaction Ensemble method, pointing out differences and similarities between these two approaches and explaining the reasons to prefer the latter in some particular cases. The first part of this tutorial will consider a homogeneous aqueous solution of a titratable acidic species HA that can dissociate as

follows:

$$HA \Longrightarrow A^- + H^+,$$
 (1)

while second part will consider a polyelectrolyte chain formed by a series of titratable monomers bonded together.

Both constant pH and Reaction Ensemble methods are implemented via a Monte Carlo algorithm with the classical Metropolis-Hastings acceptation rule. Here we compare both methods based on the paper published by Landsgesell et. al[1].

2 Theoretical Background

In this tutorial we deal with chemical reactions of the following kind:

$$HA \Longrightarrow A^- + H^+.$$
 (2)

If $N_0 = N_{\rm HA} + N_{\rm A^-}$ is the number of titratable groups in solution, a degree of dissociation α can be defined:

$$\alpha = \frac{N_{A^-}}{N_0}. (3)$$

Another observable is the degree of association which is given by:

$$\overline{n} = \frac{N_{\rm HA}}{N_0} = 1 - \alpha.$$

The equilibrium concentration of each species is defined by the equilibrium reaction constant

$$K = \exp(\beta \sum_{i} \nu_i (\mu_i - \mu_i^0)) = \frac{a(A^-)a(H^+)}{a(HA)};$$
 (4)

for each species i $a(i) = e^{\beta(\mu_i - \mu_i^0)} = (c_i/c_i^0)\gamma_i$ denotes the relative activity, μ_i is the chemical potential, μ_i^0 is the chemical potential under some standard reference conditions, ν_i is the stoichiometric coefficient, γ_i is the activity coefficient of species i, c_i and c_i^0 are the concentration of the species i and its standard concentration, respectively. In the case of non-interacting particles (ideal gas) or very dilute solutions ($\gamma_i \approx 1$) activities are equal to c_i/c_i^0 . In chemical equilibrium, which is defined by $\Delta G = \sum_i \nu_i \mu_i = 0$, we obtain that $K = \exp(\beta \Delta (G - G^0)) = \exp(-\beta \sum_i \nu_i \mu_i^0)$. Therefore we have (in the case of no interactions):

$$\exp(\beta[(\mu_{\rm H^+} - \mu_{\rm H^+}^0) + (\mu_{\rm A^-} - \mu_{\rm A^-}^0) - (\mu_{\rm HA} - \mu_{\rm HA}^0)]) = \prod_i \left(\frac{c_i}{c_i^0} \gamma_i\right)^{\nu_i} = \text{const.}$$
 (5)

Equivalently, for reaction (2) and in case of a dilute solutions ($\gamma_i \approx 1$), equation (5) can be approximated as follows:

 $K_{\rm c} = \frac{c_{\rm H}c_{\rm A}}{c_{\rm HA}} = {\rm const},$ (6)

where K_c carries the dimension 1/volume.

2.1 Constant pH method

In the constant pH method, the acceptance probability for a reaction is

$$P_{\rm acc} = \min \left\{ 1, e^{\beta \Delta E_{\rm pot} \pm \ln 10(\text{pH - pK}_{\rm a})} \right\}, \tag{7}$$

and the acceptance probability of a reaction is $P_{\text{acc}} = \frac{N_{\text{HA}}}{N0}$ for a dissociation and $P_{\text{acc}} = \frac{N_{\text{A}}}{N0}$ for an association reaction [1]. Here ΔE_{pot} is the potential energy change due to the reaction, and pH - p K_{a} is an input parameter. pH and K_{a} represent, respectively, the concentration of H⁺ ions in the solution and the thermodynamic dissociation constant for HA species when the system is approximated as a dilute solution ($\gamma_i \approx 1$):

$$pH = -\log_{10} \frac{c_{H^+}}{c_{H^+}^0}; \tag{8}$$

$$K_{\rm a} = \frac{(c_{\rm H}^+/c_{\rm H}^0)(c_{\rm A}^+ - / c_{\rm A}^0)}{(c_{\rm HA}/c_{\rm HA}^0)}.$$
 (9)

The chemical prefactor ± 1 defines the direction of the reaction (+1 dissociation, -1 association). When a dissociation move is attempted, the titratable molecule HA is charged and a counterion H⁺ is randomly placed into the simulation box, while when an association move is attempted, a A⁻ is neutralized and a random counterion H⁺ is removed from the cell.

2.2 Reaction Ensemble method

A chemical reaction involving n species of type s_i and with stoichiometric coefficients ν_i can be written as

$$\sum_{i=1}^{n} \nu_i s_i = 0. {10}$$

For such a general reaction, the acceptance probability in the Reaction Ensemble method is defined as

$$P_{\text{acc}} = \min \left\{ 1, K_c^{\xi} V^{\bar{\nu}\xi} \prod_{i=1}^{n} \left[\frac{N_i^{0!}!}{(N_i^{0} + \xi \nu_i)} \right] e^{-\beta \Delta E_{\text{pot}}} \right\}.$$
 (11)

Here K_c is the ideal reacting gas quantity introduced above, V the volume, $\bar{\nu}$ the total change in the number of particles, N_0^i the number of particles prior to the reaction and ξ is the extent of the reaction, which could assume value +1 (forward reaction) or -1 (backward reaction). Each reaction is proposed with uniform probability. For reaction (2) eq. (11) can be simplified:

$$P_{\rm acc} = \min \left\{ 1, \left(K_c \frac{N_{\rm HA}}{N_{\rm A} - N_{\rm H+}/V} \right) e^{-\beta \Delta E_{\rm pot}} \right\}.$$
 (12)

Notice that in this case you can also define $K_a = K_c/c_0$.

The main difference in the two methods consists in the fact that in the Reaction Ensemble the system pH is determined via the actual proton concentration in the simulation box, while in Constant pH method it represents an input parameter which remains constant during all the simulation.

3 First part: homogeneous aqueous solution of acidic species

3.1 System Setup

We start defining some important input parameters and setting the particles randomly inside the box:

For each negative charged A^- particle (type = 1) we put in the simulation box the respective counterion H^+ (type = 2) in order to maintain the electroneutrality of the

system. Notice that the implementation in Espresso requires that the dimension of the equilibrium constant is consistent with its internal unit of volume; rules for a correct conversion of K_a (experimental) to K_c (in internal units) are explained in the user guide: http://espressomd.org/html/doc/advanced_methods.html. The next step is to define the reaction:

```
RE=None
mode=="reaction_ensemble"
if(mode=="reaction_ensemble"):
    RE = reaction_ensemble.ReactionEnsemble(temperature=1, exclusion_radius=1)
elif(mode=="constant_pH_ensemble"):
    RE = reaction_ensemble.ConstantpHEnsemble(temperature=1, exclusion_radius=1)
    RE.constant_pH=7
    RE.add_reaction(gamma=K_diss, reactant_types=[0], reactant_coefficients=
        [1], product_types=[1, 2], product_coefficients=[1, 1], default_charges=
        {0: 0, 1: -1, 2: +1})
print(RE.get_status())
system.setup_type_map([0, 1, 2])
```

You can switch from one method to the other simply by changing the "mode" parameter. Now the system is ready to be simulated:

```
Nreac = 10**4 #number of association/dissociation attempt to be performed
for i in range(Nreac):
    RE.reaction()
```

Notice that, as this level, we're not taking into account electrostatic interactions between charged species. If you compare the input equilibrium constant K_c with the effective one you can calculate at the end of a simulation run: $K_c^{(\text{eff})} = \frac{\langle c_{\text{A}} - \rangle \langle c_{\text{H}} + \rangle}{\langle c_{\text{HA}} \rangle}$ one observes that both are equal in the case of no interactions present. This is not true when e.g. electrostatic interaction are enabled due to the fact the latter introduces an excess chemical potential.

3.2 Dissociation degree versus concentration

Performing several simulations with constant N_0 but varying the dimension of the simulation box it is possible to obtain the dissociation degree α (eq. 3) as a function of the concentration of titratable units. Results are shown in figure 1.

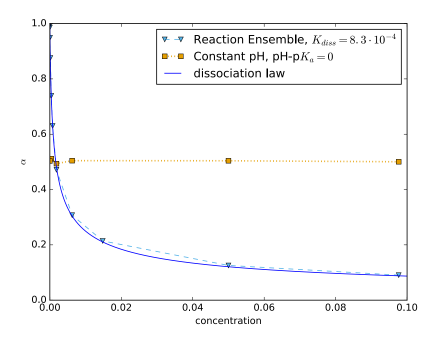


Figure 1: Dissociation degree α as function of the concentration of tritatable groups. Green: Constant pH method; violet: Reaction Ensemble method; black: ideal behavior.

As you can see, only the curve obtained with the Reaction Ensemble method fits the ideal behavior described by the dilution law

$$K_a = \frac{\alpha^2}{1 - \alpha} c_{\text{titr}},\tag{13}$$

where $c_{\rm titr}$ is the concentration of titratable units in solution, and this is due to the fact that the acceptance rule in Constant pH method does not depend on the volume of the system. As α differs, so will the number of counterions in the cell. This could have a strong impact on screening effects when electrostatic interactions are taken into account. Moreover, in the Constant pH method the real chemical nature of counterions is unknown. In fact, when a HA molecule dissociates at high pH value, the generated H⁺ would react instantaneously with an OH⁻ ions, so the positively charged ions that remains in solution must represent a different species (e.g. a Na⁺ cation).

4 Second part: linear weak polyelectrolyes

Weak polyelectrolytes are a family of responsive materials whose application spans the range from flocculation-induced water purification to tissue targeted drug delivery of expensive or cytotoxic medicines. Their properties, however, depend on their specific chemical environment, with details such as pH, salt concentration and valency, and their topology markedly modifying the chemical behavior of their ionizing groups. Here we're going to study the titration curves of a linear weak polyelectrolytes composed by 50 titratable units with both Reaction Ensemble and Constant pH methods. In this case, we'll enable electrostatic interactions between charged species in order to correctly describe their properties.

```
# setting up electrostatics
from espressomd import electrostatics
p3m = electrostatics.P3M(prefactor = l_bjerrum*temperature, accuracy=1e-3)
system.actors.add(p3m)
First of all, we define a few important parameters:
# system parameters defined in terms of internal unit sigma = 3.55 Angstrom
box_1 = 56.134
1 \text{ bjerrum} = 2.0
temperature = 1.
system = espressomd.System(box_1=[box_1]*3)
# setup partilcles
# when you have to handle with many differet type of species in your system,
# make sure to define variables and dictonaries that can be helpful to reduce
# mistakes when you write your scripts.
N_P = 1
              #number of chains
MPC = 50
              #monomers per chain
NO = N_P*MPC #total number of monomers
nNaOH = 0
              #number of initial Na+OH-
nHC1 = 0
              #number of initial H+Cl- (additional H+'s)
type_HA = 0
              # type 0 = HA
              # type 1 = A-
type_A = 1
type_H = 2
              # type 2 = H+
```

```
type_OH = 3  # type 3 = OH-
type_Na = 4  # type 4 = Na+
type_Cl = 5  # type 5 = Cl-

charges={}
charges[type_HA] = 0
charges[type_A] = -1
charges[type_H] = 1
charges[type_OH] = -1
charges[type_Na] = 1
charges[type_Cl] = -1
```

Here, we've defined the type and the charge of the polymer titratable units (HA \rightleftharpoons A⁻), counterions H⁺, and Na⁺OH⁻ and H⁺Cl⁻ ionic species, which can be inserted inside the simulation box. Let's set up the polymer, its counterions and bonding interactions between monomers.

To titrate a polyelectrolytic chain, which is composed by N_0 titratable units with a certain pK_a , in the Reaction Ensemble we need to modulate the pH of the environment. To do this, we need to add additional H^+ (increasing the pH) Obviously, for OH^- ions (to decrease it). We need also to introduce the respective counterions to preserve the electroneutrality of the system. We need also to take into account also the autoprotolysis of water.

```
K_diss = 0.002694 #eq constant HA <--> A-+ H+
```

```
#HA <--> A- + H+
RE.add_reaction(gamma=K_diss, reactant_types=[type_HA],
    reactant_coefficients=[1],
                                   product_types=[type_A, type_H],
    product_coefficients=[1,1], default_charges={type_HA: charges[type_HA],
    type_A: charges[type_A], type_H: charges[type_H]})
#H20 autoprotolysis
RE.add_reaction(gamma=(1/K_w), reactant_types=[type_H, type_OH],
                                                   product_coefficients=[],
reactant_coefficients=[1,1], product_types=[],
default_charges={type_H: charges[type_H], type_OH: charges[type_OH]})
 Due to the fact that, at a certain pH, the ionization degree of a weak polyelectrolytes
depends on its spatial conformation, in order to obtain correctly averaged values we have
also to couple the reaction algorithm with MD simulations.
# Integration parameters
system.time_step = 0.02
system.cell_system.skin = 1. #only for tutorial purposes
system.cell system.max num cells = 2744
system.thermostat.set_langevin(kT=temperature, gamma=1.0)
Finally, we are ready to run the simulation!
for i in range(12000):
RE.reaction()
system.integrator.run(300) # this is for tutorial only, too less integrations
if(i \% 50 == 0):
print(i,") HA", system.number_of_particles(type=type_HA), "A-",
 system.number_of_particles(type=type_A), "H+",
```

 $K_w = 10.**(-14)*0.02694**2$ #eq constant for autoprotolysis

4.1 Titration curves

For a solution of weak acidic molecules with a certain $pK_a = -\log_{10}(K_c/c^0)$, the trend of α as function of pH is described by the Henderson-Hasselbalch equation (black curve

system.number_of_particles(type=type_H), 'OH-',
system.number_of_particles(type=type_OH), 'Cl-',
system.number_of_particles(type=type_Cl), 'NA+',

system.number_of_particles(type=type_Na))

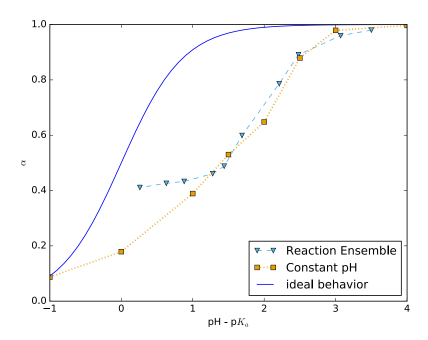


Figure 2: Titration curves for a weak polyelectrolyte chain composed by $N_0 = 50$ titratable monomers. Green: Constant pH method; violet: Reaction Ensemble method; black: ideal behavior.

in fig. 2):
$$pH = -\log_{10}(c_{H^+})/c^0 \end{red} \end{red} \end{red} \end{red} (14)$$

However, when titratable units are bonded together in a polyelectrolyte chain their effective $pK_a^{(eff)}$ differs from the ideal one pK_a ; this cam be described by the fact that when the charge carried by a dissociated monomer tends to partially inhibit the dissociation of its neighbors (at electrostatic interaction strengths present in water), and this results into a lower total degree of dissociation with respect to the non-bonded acidic units case. Anyway, this effect can be partially compensated by the presence of counterions, which are able to screen repulsive interactions between dissociated monomers. As you can observe in figure 2, Constant pH and Reaction Ensemble method results are very similar at high pH values, but they show very pronounced differences at low pH values. More in details, $pK_a^{(eff)}$ tends to the ideal one when the concentration of H^+ is

high. This depends on the fact that with Reaction Ensemble method we have to inject a strong acid (H⁺Cl⁻) in order to titrate the polymer, and this results in a more salty solution with a strong screening power. This behavior would be reversed in case of a weak poly-base, with superimposable curves at low pH values and the Reaction Ensemble one approaching the ideal one when the amount of OH⁻ ions becomes relevant.

5 Distribution of charges onto the chain

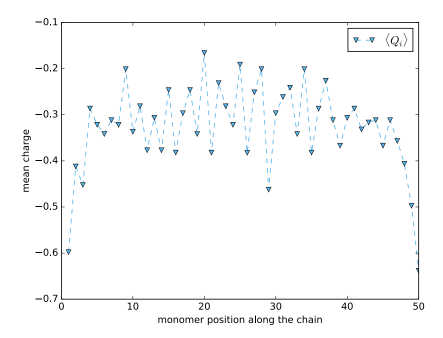


Figure 3: Mean charges of monomers as function of their position along the chain (calculated via Reaction Ensemble method) at pH - p $K_a = 2$ Notice that the curve is noisy because of the short duration of the simulation. Make sure to perform longer simulations to achieve a better convergence.

Figure 3 shows the mean charge assumed by each monomer during the simulation in function of its position along the chain. As you can observe, monomers lying at the extremities tend to be more charged than those that lie in the innermost regions. This could be easily explained thinking that the ends of the chain can better arrange in space

in order to minimize repulsive interactions between charges. This results do not depend on the method, i.e. the shape of the curve at a certain dissociation degree would be te same also with the Constant pH; however, as previously discussed, at a certain value pH - pK_a , the total dissociation degree depends on the method, so the Constant pH curve would results shifted to lower mean charge values for a weak poly-acid at high pH values.

References

[1] Jonas Landsgesell, Christian Holm, and Jens Smiatek. Simulation of weak polyelectrolytes: a comparison between the constant ph and the reaction ensemble method. *The European Physical Journal Special Topics*, 226(4):725–736, 2017.