Perform pure salt simulation (NH4CI)

- 1. Parameterisation methods
- 2. Measurements

Testing parameters

System settings: 0.15M 4-4-4 nm boxes (6 NH4, 6 Cl, 2022 H2O)

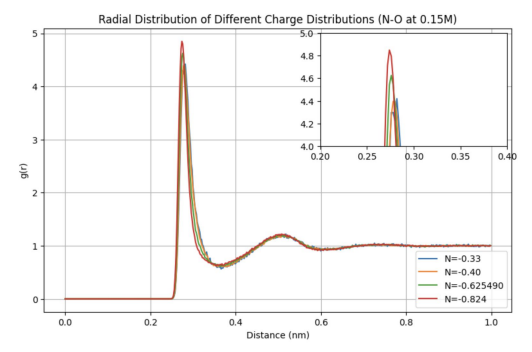
| | | Source |
|---------|------------------------------|--|
| charge0 | N: -0.33 H: +0.33 | Original CHARMM parameters |
| charge1 | N: -0.40 H: +0.35 | NH4+ partial charges using OPLS charges, as OPLS and CHARMM partial charges of NH3+ in Lysine are nearly identical [b] |
| charge2 | N: -0.625490 H: +0.406372 | Partial charges were derived using electrostatic potential fitting. Energy constants were taken from the CHARMM, version 22, forcefield for primary amines.[h] |
| charge3 | N: -0.824 H: +0.456 | NH4 charges by restrained ESP-fit method using ChelpG approach [a] |

RDF results (N-O)

| 200ns, 4-4-4 | r_max | CN | 500ns, 4-4-4 | r_max | CN | 200ns, 7-7-7 | r_max | CN |
|-----------------|-------|-------|-----------------|-------|-------|-----------------|-------|-------|
| 0.15M, q=-0.33 | 0.282 | 6.98 | 0.15M, q=-0.33 | 0.28 | 7.0 | 0.15M, q=-0.33 | 0.28 | 6.98 |
| 0.15M, q=-0.40 | 0.28 | 7.353 | 0.15M, q=-0.40 | 0.28 | 6.759 | 0.15M, q=-0.40 | 0.28 | 6.975 |
| 0.15M, q=-0.62 | 0.278 | 6.599 | 0.15M, q=-0.62 | 0.276 | 6.19 | 0.15M, q=-0.62 | 0.278 | 6.528 |
| 0.15M, q=-0.824 | 0.276 | 6.25 | 0.15M, q=-0.824 | 0.276 | 6.163 | 0.15M, q=-0.824 | 0.276 | 5.968 |
| 1M, q=-0.33 | 0.282 | 6.377 | | | | 1M, q=-0.33 | 0.282 | 6.488 |

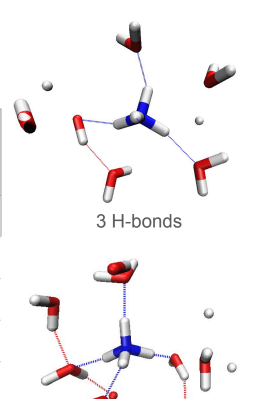
RDF results (N-O)

| 500ns, 7-7-7 | r_max | CN |
|-----------------|-------|-------|
| 0.15M, q=-0.33 | 0.282 | 6.98 |
| 0.15M, q=-0.40 | 0.28 | 6.975 |
| 0.15M, q=-0.62 | 0.278 | 6.528 |
| 0.15M, q=-0.824 | 0.276 | 5.968 |



Number of hydrogen bond

| 500ns, 4-4-4 | r_min | CN | Avg. no. of H-bond (4-4-4) | Avg. no. of H-bond (7-7-7) |
|-----------------|-------|-------|----------------------------------|----------------------------------|
| 0.15M, q=-0.33 | 0.368 | 7.0 | 2.15 | 2.25 |
| 0.15M, q=-0.40 | 0.364 | 6.759 | 2.25 | 2.34 |
| 0.15M, q=-0.62 | 0.354 | 6.19 | 2.52 | 2.61 |
| 0.15M, q=-0.824 | 0.36 | 6.163 | 2.13 | 2.82 |



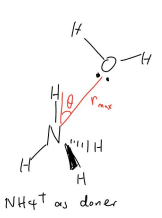
4 H-bonds

Hbond vmd plugin

Input



A hydrogen bond is formed between an atom with a hydrogen bonded to it (the donor, D) and another atom (the acceptor, A) provided that the distance D-A is less than the cut-off distance and the angle D-H-A is less than the cut-off angle

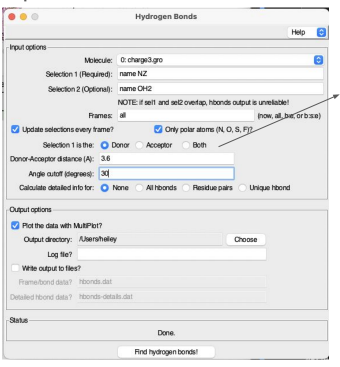


Parameters used in the calculation of hydrogen bonds:

- Atomselection 1: (name NZ) and (name "N.*" "O.*" "S.*" FA F1 F2 F3)
- Atomselection 2: (name OH2) and (name "N.*" "O.*" "S.*" FA F1 F2 F3)
- Update selections every frame: yes
- Initial frame: 0
- Frame step: 1
- Final frame: 2001
- Donor-Acceptor distance: 3.54
- Angle cutoff: 30
- Type: unique
- Write a file with H bond/frame data: yes
- Filename: /Users/heiley/Desktop/hbonds3.dat
- Details output file: /Users/heiley/Desktop/hbonds3-details.dat

Hbond vmd plugin

Input



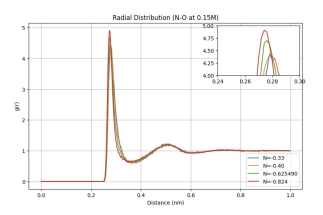
Donor vs acceptor

```
if {$DA == "D" || $DA == "both"} {
    set hbondsingleframe1 [measure hbonds $dist $ang $sel1 $sel2]
    set count1 [llength [lindex $hbondsingleframe1 0]]
}
if {$DA == "A" || $DA == "both"} {
    set hbondsingleframe2 [measure hbonds $dist $ang $sel2 $sel1]
    set count2 [llength [lindex $hbondsingleframe2 0]]
}
lappend framecount $f

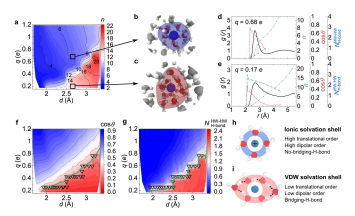
lappend numHbonds [expr $count1 + $count2]
```

Explanation of lower CN as |q_N| increases

| 500ns, 4-4-4 | r_max | CN |
|-----------------|-------|-------|
| 0.15M, q=-0.33 | 0.282 | 6.98 |
| 0.15M, q=-0.40 | 0.28 | 6.975 |
| 0.15M, q=-0.62 | 0.278 | 6.528 |
| 0.15M, q=-0.824 | 0.276 | 5.968 |



- The coordination number n drops sharply from $n \ge 12$ to n = 4 8 at qc(d) as q increases (Fig. <u>1a</u>). This result indicates that an ion with low charge density, behaving like an electrically neutral particle, forms a thick solvation shell, whereas an ion with high charge density develops a thin, well-developed solvation shell (Fig. <u>1a-c</u>). We call the former the "VDW solvation shell" and the latter the "ionic solvation shell" according to their dominant interactions to form the shell.
- In the VDW solvation shell, a water molecule in the original second shell location penetrates the open space between the first and the second shell while stabilised through its H-bonding with two molecules in the first shell (Fig. 1d, e).
- As q increases, the ion-induced water dipole reorientation destroys these bridging H-bonds, kicking the penetrating water out from the first shell. We have confirmed this mechanism in Supplementary Fig. 3 by directly monitoring the water dipole ordering and H-bonding in a non-equilibrium simulation.



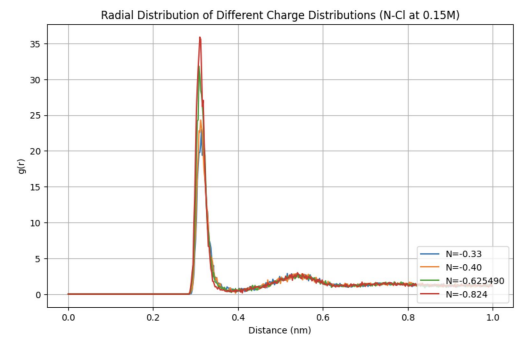
Shi, R., Cooper, A.J. & Tanaka, H. Impact of hierarchical water dipole orderings on the dynamics of aqueous salt solutions. *Nat Commun* 14, 4616 (2023). https://doi.org/10.1038/s41467-023-40278-x

RDF results (N-CI)

| 200ns, 4-4-4 | r_max | CN | 500ns, 4-4-4 | r_max | CN | 200ns, 7-7-7 | r_max | CN |
|-----------------|-------|-------|-----------------|-------|-------|-----------------|-------|-------|
| 0.15M, q=-0.33 | 0.314 | 0.078 | 0.15M, q=-0.33 | 0.316 | 0.08 | 0.15M, q=-0.33 | 0.314 | 0.082 |
| 0.15M, q=-0.40 | 0.314 | 0.08 | 0.15M, q=-0.40 | 0.314 | 0.084 | 0.15M, q=-0.40 | 0.314 | 0.084 |
| 0.15M, q=-0.62 | 0.312 | 0.091 | 0.15M, q=-0.62 | 0.312 | 0.097 | 0.15M, q=-0.62 | 0.314 | 0.097 |
| 0.15M, q=-0.824 | 0.31 | 0.102 | 0.15M, q=-0.824 | 0.31 | 0.107 | 0.15M, q=-0.824 | 0.31 | 0.107 |
| 1M, q=-0.33 | 0.314 | 0.429 | | | | 1M, q=-0.33 | 0.314 | 0.429 |

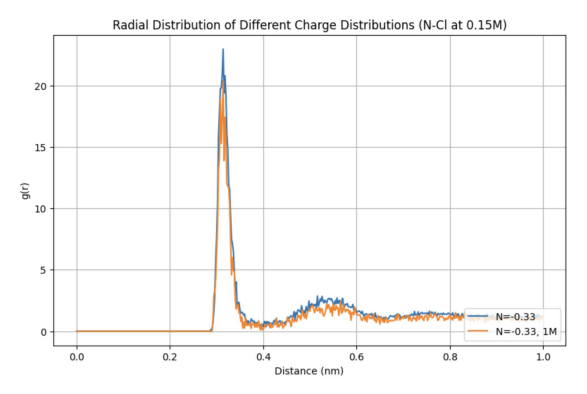
RDF results (N-CI)

| 500ns | r_max | CN |
|-----------------|-------|-------|
| 0.15M, q=-0.33 | 0.314 | 0.082 |
| 0.15M, q=-0.40 | 0.314 | 0.084 |
| 0.15M, q=-0.62 | 0.314 | 0.097 |
| 0.15M, q=-0.824 | 0.31 | 0.107 |



RDF results (N-CI)

| 200ns | r_max | CN |
|----------------|-------|-------|
| 0.15M, q=-0.33 | 0.314 | 0.082 |
| 1M, q=-0.33 | 0.314 | 0.429 |



Research on NH4 conduction in ion channel

- [a] Detailed Mechanism for AmtB Conducting NH4 /NH3: Molecular Dynamics Simulations
 - Optimised charges by restrained ESP-fit method using ChelpG approach
 - N: -0.824; H: +0.456

[b] Ammonium Recruitment and Ammonia Transport by E. coli Ammonia Channel AmtB

- NH4+ partial charges using OPLS charges
- NH4+ bond and bond angle using CHARMM27 parameters, same as NH3 moiety of Lysine

[c] Molecular Dynamics Simulations on the Escherichia coli Ammonia Channel Protein AmtB: Mechanism of Ammonia/ Ammonium Transport

- Apply Mulliken charges at the HF/6-31G(d)
- LJ parameters same as amide groups

[d] Ammonium Transporters Achieve Charge Transfer by Fragmenting Their Substrate

- NH4+ and surrounding water molecules and protein side chains are described by a polarizable force field based on the classical Drude oscillator, (62-65) and parametrized to reproduce both the free energy of hydration and the ion–protein interactions
- Pair-specific LJ parameters between the ions and atoms of the ligands are adjusted to reproduce the ab initio interaction energies
- Full geometry optimizations of the complexes between these ligands and the three cations Na+, K+, and NH4+ are performed at the MP2(FC)/6-311++G(d,p) level using Gaussian 09 program

[g] Ammonium Transport Proteins with Changes in One of the Conserved Pore Histidines Have Different Performance in Ammonia and Methylamine Conduction

- The force field parameters of ammonia and methylamine were obtained from server SwissParam. [52]
- In order to validate the force field parameters, the salvation free energy of ammonia and methylamine in water was calculated using free energy perturbation (detailed protocol shown in Text S1).
- The calculated solvation free energies for ammonia and methylamine are −5.00±0.24 and −4.49±0.18 kcal/mol (Figure S1) respectively, very similar to the experiment values (−4.31 kcal/mol for ammonia and −4.57 kcal/mol for methylamine [53]).
- Therefore the parameters used here should be suitable for simulations.

Research on NH4 conduction in ion channel

| | Method | Force field, water | Charges |
|---------|---|--|------------------------|
| Default | Parameters same as methylammonium MAMM N from primary NH3+, phosphatidylethanolamine | CHARMM36, TIP3P | N: -0.33 H: +0.33 |
| [a] | NH4 charges by restrained ESP-fit method using ChelpG approach | GROMOS87, SPC | N: -0.824 H: +0.456 |
| [b] | NH4+ partial charges using OPLS charges, as OPLS and CHARMM partial charges of NH3+ in Lysine are nearly identical | CHARMM27, H atoms were added to crystal waters using PSFGEN in NAMD | N: -0.40 H: +0.35 |
| [c] | N, H charges using Mulliken charges at the HF/6-31G(d) | CHARMM, TIP3P | Not specified |
| [d] | NH4+ and surrounding water molecules and protein side chains are described by a polarizable force field based on the classical Drude oscillator, and parametrised to reproduce both the free energy of hydration and the ion–protein interactions. Pair-specific LJ parameters between the ions and atoms of the ligands are adjusted to reproduce the ab initio interaction energies. Full geometry optimizations of the complexes between these ligands and the three cations Na+, K+, and NH4+ are performed at the MP2(FC)/6-311++G(d,p) level using Gaussian 09 program. | CHARMM27, TIP3P | 1 |
| [e] | | CHARMM36, TIP3P | Adapted from [a] |

Research on NH4 conduction in ion channel (2)

| | Method | Force field, water | Charges |
|---------|--|--------------------|------------------------------|
| Default | Parameters same as methylammonium MAMM N from primary NH3+, phosphatidylethanolamine | CHARMM36, TIP3P | N: -0.33 H: +0.33 |
| [9] | Parameters of ammonia from server SwissParam Validated with salvation free energy of ammonia in water using free energy perturbation | CHARMM27, TIP3P | |
| [h] | Parameters were obtained using the Extensible Computational Chemistry Environment (ECCE)49 after performing quantum mechanics calculations using NWChem Geometrically optimised at RHF level with 6-31111G(3df,3pd)basis set with Pople(3df,3pd) as a polarization function,and with Pople-style as a diffusion function Partial charges were derived using electrostatic potential fitting. Energy constants were taken from the CHARMM, version 22, forcefield for primary amines. | CHARMM, TIP3P | N: -0.625490 H: +0.406372 |

| | Elements | Energy constant (K _b) ^a | Bond length |
|---------------|----------|--|-------------|
| | | [kcal/(mol Ų)] | (Å) |
| Stretching | N—H | 403.0 | 1.003 |
| | | Energy constant $(K_{\theta})^a$ | Bond angle |
| | | [kcal/(mol rad²)] | (°) |
| Bending | H-N-H | 44.0 | 109.471 |
| | | Partial charge | |
| Electrostatic | N | -0.625490 | |
| | Н | +0.406372 | |
| | | | |

Other NH4 parameterisations

| | Method | Force field, water | Charges |
|---------------|--|----------------------------|--------------------------|
| Default | N from primary NH3+, phosphatidylethanolamine H from polar H+ | CHARMM36, TIP3P | N: -0.33 H: +0.33 |
| Madrid [5] | N, H charges using OPLS-based model, scaled by 0.85 | Madrid-2019, TIP4P/2005 | N: -0.3400 H: +0.2975 |

Measurements

RDF data (Na-O)

| | No. of OH2 in first hydration shell | First peak | First minimum |
|-----------------|-------------------------------------|------------|---------------|
| Simulation [1] | 5.702 | 2.36 | 3.18 |
| Literature [2] | 4.0-8.0 | 2.36 | |
| MC in TIP4P [3] | 6.2 | 2.50 | |
| Madrid [6] | 5.4 | 2.33 | |

RDF data (N-O)

| | No. of OH2 in first hydration shell | First peak | First minimum |
|--------------------|-------------------------------------|------------|---------------|
| Simulation [1] | 6.581 | 2.82 | 3.66 |
| Literature [2] | 8.1-10.0 | 2.6-3.05 | |
| MC in TIP4P [3] | 7.3 | 2.90 | |
| MC in TIP3P [4] | 7.0-7.1 | 2.63-2.70 | |
| Madrid [5] | 5.5 | 2.66 | |
| NH3 MD [16] | | 3.25 | |
| NH3 Ab Initio [16] | | 2.75 | |

RDF plots of N_{NH4} -O

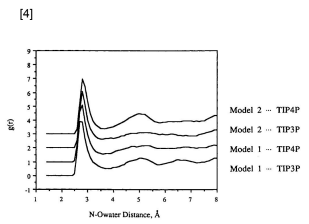
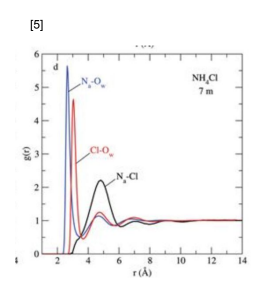
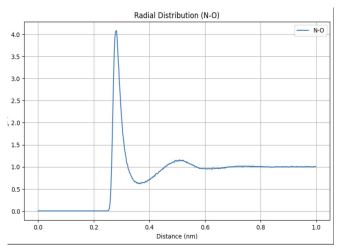


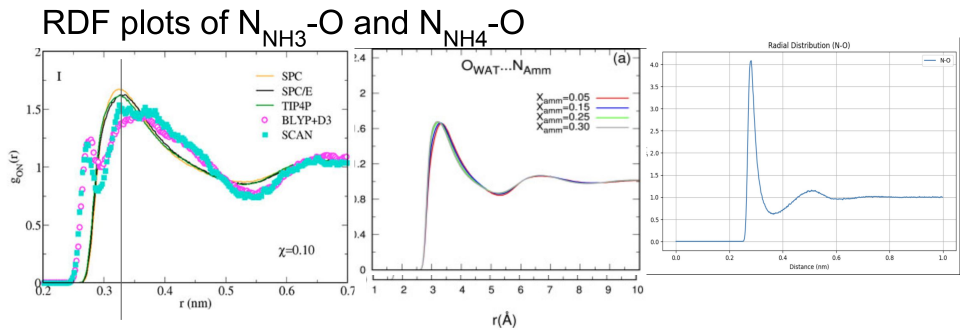
Figure 2. N— O_{water} radial distribution functions for model 1 and 2 in TIP3P and TIP4P.





NH4: First maximum found at r = 0.282nm Coordination number = 6.581 χ =0.02

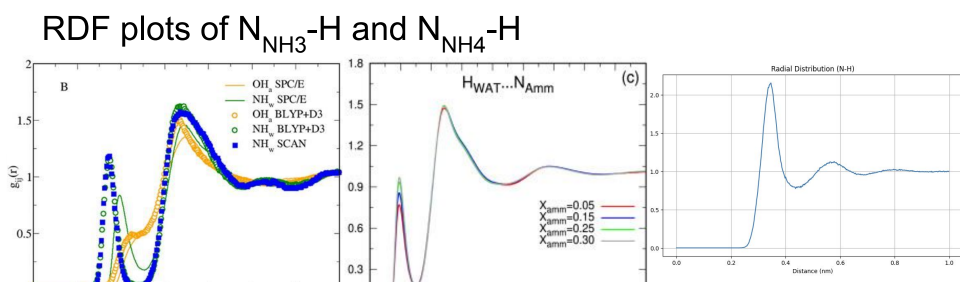
Measurements



NH3: MD (lines) and AIMD (symbols) at χ = 0.1 (NH3:H2O). The three different force fields used for water are represented by orange (SPC), black (SPC/E), and green (TIP4P) lines, while ab initio BLYP+D3 and SCAN correlations are indicated as magenta circles and cyan squares, respectively. [16]

NH3: MD simulation at $\chi = 0.05-0.30$ [17]

NH4: First maximum found at r = 0.282nm Coordination number = 6.581 $\chi = 0.02$



NH3: Comparison between SPC/E (lines), BLYP+D3 (circles), and SCAN (squares) radial distributions functions involving hydrogen atoms for T = 295 K and χ = 0.10. Correlations between atoms of the same molecule type are reported in (a), whereas cross-correlations are shown in (b). [16]

r (nm)

0.3

0.4

0.5

0.6

0.

0.2

r(Å) NH3: MD simulation at χ = 0.05-0.30 [17]

NH4 from original simulation: First maximum found at r = 0.344nm Coordination number = 20.891 $\chi = 0.02$

10

Measurements

RDF data (CI-O)

| | No. of OH2 in first hydration shell | First peak | First minimum |
|-----------------------|-------------------------------------|------------|---------------|
| Simulation, NaCl [1] | 7.911 | 3.16 | 3.84 |
| Simulation, NH4Cl [1] | 7.227 | 3.16 | 3.80 |
| Literature [2] | 5.9-8.5 | 3.1-3.3 | |
| MC in TIP4P [3] | 7.6 | 3.25 | |
| Madrid [6] | 5.9 | 3.05 | |

Free energy calculations

| | Absolute hydration energy of NH4 | Absolute hydration energy of Na | Relative hydration energies between NH4+ and Na+ |
|------------------------------|----------------------------------|---------------------------------|--|
| Experiment [7] | -78.6 | | -21.7 kcal/mol |
| Free energy perturbation [4] | -79.55 (+ Born) -98.85 | | -28.87 kcal/mol |
| Empirical model [8] | -68.1 | -87.2 | -19.1 kcal/mol |
| MC simulation [3] | -73.1 | -92.3 | -19 kcal/mol |

Osmotic coefficient at 1M/1m

| | NH4CI | NaCl |
|---------------------------------------|--------|-----------|
| Isopiestic vapour pressure method [9] | 0.897 | |
| MC simulations [10] | | 0.91-0.92 |
| Experiment [11] | | 0.94 |
| MD simulations [12] | | 0.93 |
| Experiment [13] | 0.927 | |
| Hückel equation calculation [14] | 0.902 | |
| Experiment [15] | 0.9671 | |

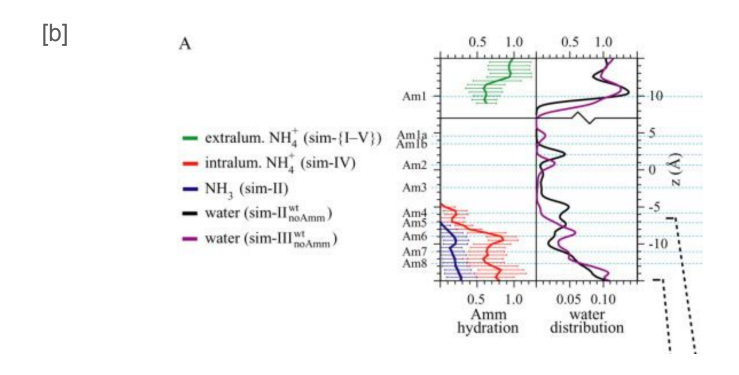
References

- [1] https://github.com/Heiley-W/project/blob/main/RDF-analysis.ipynb
- [2] Ionic radii in aqueous solutions https://doi.org/10.1021/cr00090a003
- [3] Halide, Ammonium, and Alkali Metal Ion Parameters for Modeling Aqueous Solutions https://doi.org/10.1021/ct600252r
- [4] Free Energy Calculations Involving NH4+in Water https://doi.org/10.1002/jcc.540120106
- [5] Further extension of the Madrid-2019 force field: Parametrization of nitrate and ammonium ion https://doi.org/10.1063/5.0177363
- [6] A force field of Li+, Na+, K+, Mg2+, Ca2+,Cl-, and in aqueous solution based on the TIP4P/2005 water model and scaled charges for the ions https://doi.org/10.1063/1.5121392
- [7] Ref 33 and 22 of [4] respectively
- [8] A simple empirical model describing the thermodynamics of hydration of ions of widely varying charges, sizes, and shapes https://doi.org/10.1016/0301-4622(94)00051-4
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- [10] Osmotic and activity coefficients from effective potentials for hydrated ions https://doi.org/10.1103/PhysRevE.55.5689
- [11] Ref 20 of [10]
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- [14] Mean Activity Coefficients and Osmotic Coefficients in Aqueous Solutions of Salts of Ammonium Ions with Univalent Anions at 25 °C https://doi.org/10.1021/je300474k
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- [16] The structure of water-ammonia mixtures from classical and ab initio molecular dynamics https://doi.org/10.1063/5.0220328
- [17] Effects of concentration and pressure on the aqueous solvation structure of ammonia and composition dependent ion solvation scenario in water-ammonia mixtures https://doi.org/10.1016/j.fluid.2020.112507

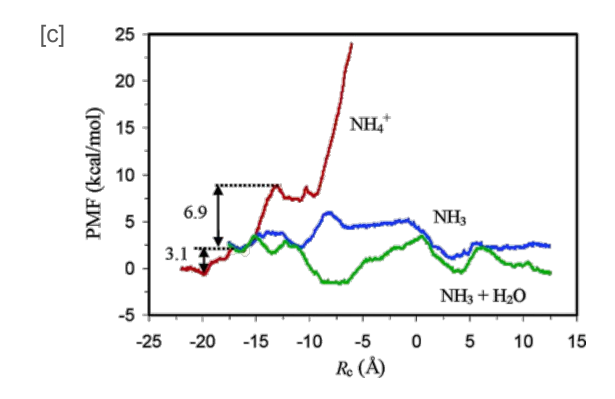
References

- [a] Detailed Mechanism for AmtB Conducting NH4+/NH3: Molecular Dynamics Simulations https://doi.org/10.1529/biophysi.106.090191
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- [e] A two-lane mechanism for selective biological ammonium transport https://doi.org/10.1101/849562
- [f] The mechanism of ammonia transport based on the crystal structure of AmtB of Escherichia coli https://doi.org/10.1073/pnas.0406475101
- [g] Ammonium Transport Proteins with Changes in One of the Conserved Pore Histidines Have Different Performance in Ammonia and Methylamine Conduction https://doi.org/10.1371/journal.pone.0062745
- [h] Free-energy profiles for ions in the influenza M2-TMD channel https://doi.org/10.1002/prot.22376

Research on NH4 conduction in ion channel



Research on NH4 conduction in ion channel



Research on NH4 conduction in ion channel

[f] Experiment

