## 2.3 Debye-Hückel theory - activity coefficient of solutions

The long range and strength of the Coulomb interaction between ions typically dominates the non ideal behavior of ionic solutions. It is as well the reason that even for extremely diluted solutions these non idealities have to be taken into account. Opposite charges attract each other. Thus it is more likely that anions are found closely to cations, and vice versa. Although the over all solution is electrically neutral counter ions are on average more likely to be found close to any ion. This spherical haze of counter ions (ionic atmosphere) around a central ion has on (time) average the same charge but with opposite sign than the central ion, leading to a shielding of the central ion, as schematically shown in Fig. 2.2a).



Figure 2.2: a) Schematic drawing of the short range order of ions in a solution; b) Debye-Hückel limiting case (red straight line) and the result of extended Debye Hückel theory

The corresponding lowering of the energy is described by the difference to the ideal molar Gibbs energy, and thus is identical to  $RT \ln \gamma_{\pm}$ . Before discussing the details we will just summarize the final results: At very low concentration the activity coefficient can be calculated from the Debye-Hückel limiting law

$$\log(\gamma_{\pm}) = -|z_{\pm}z_{-}| A\sqrt{I} \quad , \tag{2.17}$$

where  $z_+$  and  $z_-$  are the number of charged cations and anions, A is a constant (depending on temperature and solvent properties; e.g. A = 0.509 for aqueous solution at 25°C), and I is the dimensionless ionic strength, defined as

$$I = \frac{1}{2} \sum_{i} z_i^2 \left(\frac{m_i}{m^0}\right) \quad . \tag{2.18}$$

As discussed in the following subsection and illustrated in Fig. 2.2 b) Eq. (2.17) holds for strong electrolytes, low concentration (up to  $10^{-2}$  mol/l), and low charge numbers.

For higher concentration or ions with larger charge numbers as shown in Fig. 2.3 the extended Debye Hückel (or Davies) equation can be used:

$$\log(\gamma_{\pm}) = -\frac{|z_{\pm}z_{-}| A\sqrt{I}}{1 + B\sqrt{I}} + CI \quad .$$
(2.19)



Figure 2.3: a) Table for some activity coefficients  $\gamma_{\pm}$  in water at 298 K; b) measured activity coefficients of CaCl<sub>2</sub>

Before going into any details some general aspects of the Debye Hückel approach are summarized

- the classical Poisson equation (i.e. a differential equation) is solved which relates local charge densities  $\rho(r)$  to second derivatives of the electrical potential (energy)
- the situation becomes "difficult" because the local energy distribution E(r) in turn defines the charge distribution via the Boltzmann factor
- instead of "solving" the complicated differential equation a good guess for the charge density distribution  $(\propto \exp(-r/r_D))$  is chosen for which a self consistent solution for the Debye length  $r_D$  is calculated
- the final results, i.e. the difference to the random distribution of charges without shielding is translated into an activity.

It is worth to go into all the tedious details of the solution because similar systems like the Debye layer at metal surfaces or the Helmholtz layers at solid electrolyte interfaces are solved in much the same way.