The low frequency modes of solvated ions and ion pairs in aqueous electrolyte solutions: iron(II) and iron(III) chloride

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We have investigated the hydration dynamics of solvated iron(II) and iron(III) chloride. For this, THz/FIR absorption spectra of acidified aqueous FeCl₂ and FeCl₃ solutions have been measured in a frequency range of 30–350 cm⁻¹ (±1–10 THz). We observe a nonlinear concentration dependence of the absorption, which is attributed to the progressive formation of chloro-complexes of Fe(II) and Fe(III), respectively. By principal component analysis of the concentration dependent absorption spectra, we deduced the molar extinction spectra of the solvated species Fe²⁺ + 2Cl⁻ and FeCl⁺ + Cl⁻, as well as FeCl₂⁺ + 2Cl⁻ and FeCl₃⁺ + Cl⁻. In addition, we obtain ion association constants log K_FeCl₂ = −0.88(5) and log K_FeCl₃ = −0.32(16) for the association of Fe²⁺ and Cl⁻ to FeCl⁺ and the association of FeCl₂⁺ and Cl⁻ to FeCl₃⁺, respectively. We performed a simultaneous fit of all the effective extinction spectra and their differences, including our previous results of solvated manganese(II) and nickel(II) chlorides and bromides. Thereby we were able to assign absorption peaks to vibrational modes of ion–water complexes. Furthermore, we were able to estimate a minimum number of affected water molecules, ranging from ca. 7 in the case of FeCl⁺ + Cl⁻ to ca. 21 in the case of FeCl₂⁺ + Cl⁻.

1 Introduction

Ion hydration phenomena have attracted the attention of many researchers.¹⁻⁴ Whereas macroscopically ion hydration is well understood, at the molecular level it is still part of an ongoing and sometimes controversial discussion.³⁻⁸ Recently, the solvation dynamics of alkali and alkaline earth metals have been studied in detail.²⁻³,⁶⁻⁹ Most of these studies indicate prevalently self confined cations and anions in aqueous phase at low and moderate concentrations.²⁻⁶⁻⁸,¹⁰ In certain cases, however, cooperative effects between the ions have been postulated.⁹

The hydration properties of transition metal ions have only recently been studied.¹¹⁻¹³ The behaviour of ferrous (Fe²⁺) and ferric (Fe³⁺) iron in aqueous solution with respect to their tendency towards formation of distinct chloro-complexes are of special interest to the fields of iron metabolism and biological function,¹₄,¹₅ geochemical and hydrothermal studies,¹⁶,¹⁷ isotopic fractionation¹⁸ and chemistry of natural waters.¹⁹ In spite of a broad range of technical applications, simultaneous quantitative analysis of aqueous iron(II) and iron(III) at high concentrations still remains a challenge. Calorimetry,²⁰ isotachophoresis,²¹ ion chromatography,¹⁹⁻²² and spectrophotometry,¹⁷,²₃⁻²⁷ are restricted to the millimolar concentration range, since the formation of ion associates complicates data evaluation at higher concentrations.

While the masses of Fe²⁺ and Fe³⁺ are practically the same, the ionic radii of the ions differ considerably (0.92 Å for Fe²⁺ and 0.79 Å for Fe³⁺) due to the different charge states. Fe³⁺ has a much higher charge density, which causes substantially different hydration behavior compared to Fe²⁺. Besides the octahedral hexaqua complex [Fe(H₂O)₆]³⁺, several chloro-complex species have been identified in acidic solutions of ferrous chloride, which are: octahedral monochloro-complex [Fe(H₂O)₅Cl]⁺, dichloro-complex [Fe(H₂O)₄Cl₂], and tetrahedral tetrachloro-complex [FeCl₄]⁻.²⁻²⁸ The latter species, however, is formed exclusively at very high chloride excess and/or high temperatures.²⁸,³⁰ Similarly, the complexes formed by ferric chloride are: octahedral hexaqua complex [Fe(H₂O)₆]³⁺, monochloro-complex [Fe(H₂O)₅Cl]⁺, dichloro-complex trans-[Fe(H₂O)₄Cl₂], trichloro-complex [Fe(H₂O)₃Cl]₃ and tetrahedral tetrachloro-complex [FeCl₄]⁻.¹⁷,²¹⁻⁴¹ Similarly to the ferrous salt, the highest order chloro-species is only found at high chloride excess and/or high temperatures.

So far, the equilibrium formation constant of the ferrous monochloro-complex at room temperature has been determined by spectrophotometry²⁸,²⁹ and potentiometry⁴¹ to be 0.69 kg mol⁻¹, 0.43 kg mol⁻¹ and 0.75 kg mol⁻¹, respectively. The formation constant of the dichloro-complex has been determined spectrophotometrically to be 0.018 kg² mol⁻².²⁸
In contrast to ferrous chloride, there have been many attempts to determine equilibrium constants for the formation of ferric chloro-complexes in aqueous phase using spectrophotometry\(^{17,26,32,33,44-49}\) and potentiometry\(^{20,51}\). Most of them, however, did not include the concept of ion activity, and thus there is a clear divergence in the values of formation constants for solutions of different ionic strength. Only a handful of publications present an estimate of equilibrium constants at zero ionic strength, either by extrapolation\(^{32,33,44,51}\) or by introducing activity coefficients for the ions.\(^{17,26,49}\) While the values for the formation constant of the monochloro-complex at room temperature are in good agreement, ranging from 19.1 to 33.1 kg mol\(^{-1}\),\(^{17,14,49,51}\) there are large inconsistencies present for the formation constant of the dichloro-complexes ranging from 2 to 14 kg mol\(^{-1}\).\(^{17,26,33,44,49,51}\) The trichloro-complex formation constant is considerably smaller, ranging from 0.04 to 0.08 kg mol\(^{-1}\).\(^{17,26,33,44}\) Note that cumulative formation constants have been converted to stepwise formation constants.

Despite these previous efforts, several questions remain unresolved. For instance, are the aforementioned methods sensitive only to the formation of first shell complexes, or could other associates also play a role? Do the water molecules incorporated in ferrous and ferric complexes exhibit a different dynamic behaviour compared to bulk water molecules? Is water beyond the first hydration shell affected by the ions and ion associates as well?

In the present study, we investigate the hydration behavior of iron(II) and iron(III) chloride in water with respect to the formation of ion pairs, as well as the ion pair absorption spectra.

Here we investigate the concentration dependent THz/FIR absorption spectra of iron(II) and iron(III) chloride in water with respect to the formation of ion associates. Using principal component analysis (PCA) as an unbiased mathematical procedure in conjunction with a chemical equilibrium model, we are able to dissect the experimental spectra into the absorption features of the various ions and complex species. This enables us to extract information about the vibrational properties of the solvated ions and ion associates, as well as their dynamical hydration shells. This study provides a proof of principle for the applicability of THz/FIR spectroscopy as an analytical tool for the simultaneous determination of Fe(II) and Fe(III) chloride at concentrations up to the solubility limit.

2 Materials and methods

2.1 Materials

The sample solutions were prepared by dissolving the weighed out amounts of FeCl\(_2\)-4H\(_2\)O and FeCl\(_3\)-6H\(_2\)O (Sigma Aldrich, 99% purity) in HPLC grade water. In order to prevent oxidation and hydroxide formation, all solutions were acidified with HCl to an H\(^+\) concentration of 1 M, corresponding to pH 0. For determination of water concentration in the samples, densities were measured at 20 °C using an Anton-Paar DMA58 density meter.

2.2 FTIR measurements

THz/FIR absorption measurements were carried out using a Bruker Vertex 80v FTIR spectrometer equipped with a liquid helium cooled silicon bolometer from Infrared Laboratories as detector (for more specific details about the experimental setup, see Schmidt et al.\(^7\) and supporting information therein). During the complete series of measurements, the sample compartment was constantly purged with technical grade dry nitrogen to avoid humidity effects of the air. The temperature of the sample cell was kept constant at 20 ± 0.2 °C with a commercial high precision thermostat. For each single spectrum, 256 scans with a resolution of 2 cm\(^{-1}\) were averaged. The sample solutions were placed in a standard Bruker liquid cell between two parallel TPX windows of 4 mm thickness with a Kapton spacer of ca. 25 µm thickness. The exact sample layer thickness was determined prior to each measurement using the Fabry–Pérot etalon effect of the empty cell.

Using Lambert–Beer’s law, the frequency dependent absorption coefficient \(\alpha(\nu)\) of an aqueous solution is expressed as

\[
\alpha_{\text{solute}}(\nu) = \frac{1}{d} \log \left( \frac{I_{\text{solution}}(\nu)}{I_{\text{water}}(\nu)} \right) + \alpha_{\text{water}}(\nu)
\]

where \(d\) is the sample thickness, \(I_{\text{water}}(\nu)\) and \(I_{\text{solution}}(\nu)\) are the transmitted intensities of the water reference and the sample, respectively, and \(\alpha_{\text{water}}(\nu)\) is a fit of the absorption spectrum of water.\(^{52,55}\) In this way, artefacts due to reflections at the cell windows are minimized.

The contribution of HCl to the absorption was determined in an independent measurement and subtracted from all subsequent measurements, assuming a weak interaction with the coexisting ions.

For further data evaluation, we subtract the expected absorption of water in the sample to get the effective ionic absorption \(\alpha_{\text{ion}}^{\text{eff}}\):

\[
\alpha_{\text{ion}}^{\text{eff}}(\nu) = \alpha_{\text{solute}}(\nu) - \frac{C_w}{C_w^0} \alpha_{\text{water}}(\nu)
\]

where \(C_w\) and \(C_w^0\) are the actual water concentrations in the solution and in pure water, respectively, as determined by density measurements. Deviations from linear concentration dependence can be examined by comparing the effective molar
extinction $e_{\text{eff}}$ of each sample, which in turn is determined by dividing $e_{\text{ion}}$ by the actual salt concentration $c_s$:

$$e_{\text{ion}}(\tilde{\nu}) = \frac{e_{\text{eff}}(\tilde{\nu})}{c_s}$$

(3)

It is important to mention here that any changes in solvation water absorption induced by the ions are inherent in the effective ionic extinction $e_{\text{eff}}$. We assume that the interaction between each ion and surrounding water molecules can be understood by taking into account two contributions. Since the absorption properties of water are modified in the vicinity of ions, subtraction of the bulk water absorption from the sample absorption results in a negative contribution with the line shape of the water spectrum. The hydrated ion, however, has additional low frequency modes which can be described in terms of rattling modes and/or vibrational modes of the ion–water complexes.7,8

3 Experimental results

Sample solutions with concentrations ranging from 0.5 M to 3.5 M of FeCl2 and FeCl3, respectively, were measured using FTIR spectroscopy. The absorption spectra and the effective molar extinction spectra of both salts, respectively, are presented in Fig. 1 and 2. The positive value of the extinction over the entire frequency range can be attributed to the solvated ions. By examination of Fig. 1B and 2B it is obvious that the two iron chlorides have distinct nonlinear absorption features (marked with black arrows).

4 Spectral dissection

In order to deconvolute the experimental spectra into distinct components, we performed a principal component analysis (PCA) on the $e_{\text{eff}}$ spectra. Using this method (which has been described in detail before52), we were able to reduce both data sets of FeCl2 and FeCl3 to linear combinations of two components each, which include more than 96% of the information in both cases. The loading vectors and concentration dependent scores of the principal components of both salts are shown in Fig. 3.

Although for both salts the results of the PCA are rather similar, the underlying mechanism seems to be different. Comparing different sets of association constants from literature (cf. Section 2) it becomes evident that in the case of FeCl2, the solvated ions and the first ion associate FeCl+ dominate over the whole concentration range, while in the case of FeCl3, the first and second associates, FeCl2+ and FeCl3+, dominate the solution’s composition.

For the further evaluation of our data we make the assumption that next to Fe2+, FeCl+, FeCl2+, FeCl3+ and Cl− the contribution of other species to the observed absorption is negligible.
Any changes in the concentration dependent absorption spectrum are attributed to a shift in the equilibrium:

\[ \text{Fe}^{2+} + \text{Cl}^- \rightleftharpoons \text{FeCl}^{+} \tag{4} \]

\[ \text{FeCl}^{2+} + \text{Cl}^- \rightleftharpoons \text{FeCl}_2^{+} \tag{5} \]

The equilibrium constants \( K_{\text{FeCl}_2} \) and \( K_{\text{FeCl}_3} \) are defined by the ratio of product and educt concentrations \( c \) and their respective activity coefficients \( \gamma \) according to the following equations:

\[ K_{\text{FeCl}_2} = \frac{c_{\text{FeCl}^+}}{c_{\text{Fe}^{2+}} \cdot c_{\text{Cl}^-}} \cdot \frac{\gamma_{\text{FeCl}^+}}{\gamma_{\text{Fe}^{2+}} \cdot \gamma_{\text{Cl}^-}} \tag{6} \]

\[ K_{\text{FeCl}_3} = \frac{c_{\text{FeCl}_2^+}}{c_{\text{FeCl}^+} \cdot c_{\text{Cl}^-}} \cdot \frac{\gamma_{\text{FeCl}_2^+}}{\gamma_{\text{FeCl}^+} \cdot \gamma_{\text{Cl}^-}} \tag{7} \]

The activity coefficients of the mixed compounds FeCl\(^+\), FeCl\(^{2+}\) and FeCl\(^{3+}\) are not known. However, in first order approximation the activity coefficients of the iron containing species in both the numerator and denominator are similar (\( \gamma_{\text{FeCl}^+} \approx \gamma_{\text{Fe}^{2+}} \) and \( \gamma_{\text{FeCl}_2^+} \approx \gamma_{\text{FeCl}^+} \)).\(^{32}\) This approximation is based on the assumption that for solutions of high ionic strength the activity coefficient is not determined by the charge of a species, but rather by the chemical structure. Experimental activity coefficients\(^{56,57}\) were used to determine the Pitzer parameters for FeCl\(_2\) and FeCl\(_3\) according to the model of Harvie and Weare,\(^{58}\) which are shown in Table 1. These parameters allow us to predict the activity coefficient \( \gamma_{\text{Cl}^-} \) in solutions of any ionic strength.

The respective underlying models for the effective ionic absorption are the following:

\[ \chi_{\text{ion,FeCl}_2} = \chi_{\text{single}} + \chi_{\text{pair}} + \chi_{\text{triple}} \tag{8} \]

\[ \chi_{\text{ion,FeCl}_3} = \chi_{\text{pair}} + \chi_{\text{triple}} + \chi_{\text{triple}} \tag{9} \]

Although we assume that the concentration of free Fe\(^{3+}\) is close to zero, the association constant for the formation of FeCl\(^{2+}\) cannot be neglected. Therefore we used the mean of four literature values in our calculation (19.0,\(^{49}\) 26.3,\(^{17}\) 30,\(^{54}\) and 33.1).\(^{55}\)

We performed a fit of the scores of the first two principal components with \( K_{\text{FeCl}_2} \) and \( K_{\text{FeCl}_3} \), respectively, as parameters, as has been described before.\(^{52}\) Clearly the fitted lines match the scores quite well (see insets in Fig. 3, solid lines). The resulting values for the equilibrium constants are presented in Table 2.

In addition to the equilibrium constants, we were also able to deduce the molar extinction spectra of the different solvated species. The result is displayed in Fig. 4. Note that each of these spectra contains the anion contribution corresponding to the stoichiometry of the salt \( (\chi_{\text{single,FeCl}_2} = \chi_{\text{FeCl}^+} + 2\chi_{\text{Cl}^-}; \chi_{\text{pair,FeCl}_2} = \chi_{\text{FeCl}^+} + \chi_{\text{Cl}^-}; \chi_{\text{pair,FeCl}_3} = \chi_{\text{FeCl}_2^+} + 2\chi_{\text{Cl}^-}; \chi_{\text{triple,FeCl}_3} = \chi_{\text{FeCl}_2^+} + \chi_{\text{Cl}^-}) \).

The extinction spectrum of Fe\(^{2+}\) + 2Cl\(^-\) in Fig. 4A is dominated by one broad band at 150 cm\(^{-1}\) and the tail of a feature that peaks at > 350 cm\(^{-1}\). The spectrum of FeCl\(^+\) + Cl\(^-\) shows two resonances centered at 145 cm\(^{-1}\) and 230 cm\(^{-1}\), as well as a high frequency wing.

The extinction spectrum of FeCl\(^{2+}\) + 2Cl\(^-\) in Fig. 4B displays two strong peaks at 190 cm\(^{-1}\) and 315 cm\(^{-1}\). The spectrum of FeCl\(^3+\) + Cl\(^-\) , however, shows a clear resonance around 170 cm\(^{-1}\) and at least two overlapping spectral features between 250 cm\(^{-1}\) and 350 cm\(^{-1}\).

## 5 Global fit

In our previous studies, we have performed a global fit of the aqueous ionic spectra of MnCl\(_2\), NiCl\(_2\), MnBr\(_2\) and NiBr\(_2\) and their differences, in order to pin down precise peak positions of

### Table 1

<table>
<thead>
<tr>
<th>Compound</th>
<th>( \beta(0) )</th>
<th>( \beta(1) )</th>
<th>( C^0 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>FeCl(_2)</td>
<td>0.339 ± 0.005</td>
<td>1.48 ± 0.06</td>
<td>-0.019 ± 0.001</td>
</tr>
<tr>
<td>FeCl(_3)</td>
<td>0.42 ± 0.01</td>
<td>-0.028 ± 0.002</td>
<td>7.0 ± 0.4</td>
</tr>
</tbody>
</table>

### Table 2

<table>
<thead>
<tr>
<th></th>
<th>( K (M^{-1}) )</th>
<th>\log K</th>
</tr>
</thead>
<tbody>
<tr>
<td>FeCl(_2)</td>
<td>0.13(1)</td>
<td>-0.89(4)</td>
</tr>
<tr>
<td>FeCl(_3)</td>
<td>0.49(8)</td>
<td>-0.32(8)</td>
</tr>
</tbody>
</table>
spectral features as well as to determine the effect of the ions on the water spectrum.\textsuperscript{52}

In the global fit, the anion and cation bands are modeled by a modified damped harmonic oscillator function:

\[ \tilde{\epsilon}_{DH}(\tilde{\nu}) = \frac{a_0 w_0^2 \tilde{\nu}^2}{4\pi \left[ \tilde{\nu}_0^{-2} \tilde{\nu}^2 + \left( \frac{w_0^2}{4\pi} - \frac{w_0^2}{\tilde{\nu}^2} \right)^2 \right]} \]  

where \( a_0 \) is the amplitude, \( w_0 \) the width and \( \tilde{\nu}_0 \) the center frequency of the mode. The corresponding center frequency of an unperturbed Brownian harmonic oscillator is given as:

\[ \tilde{\nu}_0 = \left( \frac{w_0^2}{4\pi} + \frac{w_0^2}{\tilde{\nu}^2} \right)^{1/2}. \]

The low and high frequency part of the spectra are modeled using the corresponding modes of the water spectrum\textsuperscript{52} scaled by factors \( n_{HF} \) and \( n_{HF} \). In addition to this, a negative contribution is present in each spectrum (cf. Section 3.2), which can be modeled by the water extinction spectrum scaled by a factor \( n_{hydration} \).

Here we extend the model to include the four spectra of FeCl\textsubscript{2} and FeCl\textsubscript{3}. The inclusion of these in the global fit results for MnCl\textsubscript{2} and NiCl\textsubscript{2} are reported for comparison. The determined fit parameters are listed in Table 3.

We observe that several species have identical parameters, \textit{e.g.} the linewidth of ionic bands, reducing the complexity of the fit. As an example for the various contributions to a fitting curve of one experimental spectrum, the spectrum of Fe\textsuperscript{2+} + 2Cl\textsuperscript{-} is displayed in Fig. 5.

While trying to fit the spectra of the ion associates, it is possible that overlapping bands of anionic and cationic species cause an ambiguity in the assignment. Sharma \textit{et al.} previously have overcome this challenge by including the difference spectra of each metal chloride and bromide into the fit, thereby uncoupling the anion from cation contributions.\textsuperscript{53} Since in the present work we lack the spectra of ferrous and ferric bromides for a direct comparison, we proceeded in an iterative approach: As a first step,
we included only the spectrum of Fe$^{2+} + 2\text{Cl}^-$ into the fit of the manganese(II) and nickel(II) chloride and bromide spectra, thereby determining the $\text{Cl}^-$ mode. In a second step, this mode was then fixed in the spectra of FeCl$^+ + \text{Cl}^-$, FeCl$_2^+ + 2\text{Cl}^-$ and FeCl$_3^+ + \text{Cl}^-$ prior to inclusion of these spectra into the fit. After the global fit, the parameters of the chloride mode were fixed to the slightly changed new values. This was repeated iteratively until the change in the chloride fit parameters was smaller than the uncertainty. In Table 3, the fixed parameters are marked with an asterisk (*).

### 6 Discussion

#### 6.1 Association constants

Prior to describing and comparing different values of ion-association constants, it is essential to emphasize that the results are method dependent. One reason for this is the different degrees of experimental sensitivity towards other ion associates besides contact ion pairs. In fact when it comes to ion association, the real number of different species is much larger than the simple one-step reaction scheme suggests. As described by Marcus and Hefter, the ions undergo a series of reaction steps from (a) solvent separated ion pairs, in which both ions still have their full hydration sphere, over (b) solvent shared ion pairs, in which the hydration spheres are overlapping, to (c) contact ion pairs, in which one ion penetrates the hydration sphere of the other. The system becomes even more complex if a third ion is included. For example, an equilibrium constant found by the anion exchange method may be smaller than the one determined using potentiometry, since the first is only sensitive towards contact ion pairs, while the latter is also sensitive to the presence of other ion pairs.

Here we performed a principal component analysis of our concentration dependent THz/FIR absorption spectra of FeCl$_2$ and FeCl$_3$ solutions, and fitted the scores with association constants $K_{\text{FeCl}_2}$ and $K_{\text{FeCl}_3}$, respectively, as fitting parameters. The logarithm of the determined parameters $K_{\text{FeCl}_2}$ and $K_{\text{FeCl}_3}$ is $-0.89(4)$ and $-0.32(8)$, respectively. These numbers are well below the values determined by other methods ranging from $-0.37$ to $-0.12$ in the case of FeCl$_2$ and 0.32 to 1.16 in the case of FeCl$_3$. This indicates that the THz/FIR absorption is especially sensitive to the formation of contact ion pairs, while the other methods might be more susceptible to solvent shared and solvent separated ion pairs.

#### 6.2 Band assignment

##### 6.2.1 The low frequency cation bands

The simultaneous fit of all spectra provides precise information about the spectral bands of the ions and ion-associates in aqueous phase. Each spectrum can be partitioned into a varying number of absorption bands, which we attribute to vibrations of the iron–water complexes. Out of these, one low frequency mode (denoted CM 1 in Table 3) is assigned to a cationic band centered between 130 and 170 cm$^{-1}$. A similar band in the spectra of magnesium chloride and bromide at around 180 cm$^{-1}$ has been attributed to a vibration of the hexaqua complex of Mg$^{2+}$, called $\Theta$ mode, in which the metal ion, together with two axial H$_2$O, moves against four equatorial H$_2$O molecules. We propose that analogous vibrations of the octahedral water complexes of the metal ions Mn$^{2+}$, Ni$^{2+}$ and Fe$^{2+}$, as well as the chloro-substituted complexes of Fe$^{2+}$ and Fe$^{3+}$, are responsible for the bands between 130 and 170 cm$^{-1}$. For the described vibrational geometry, the respective reduced masses of the hexaqua complexes increase in the order Mn$^{2+} < $ Fe$^{2+} < Ni$^{2+}$ (40.21 g mol$^{-1}$, 40.39 g mol$^{-1}$, 40.93 g mol$^{-1}$), which in absence of other effects would result in a red-shift of the band. The metal–oxygen distance, however, decreases in the order Mn$^{2+} > $ Fe$^{2+} > Ni$^{2+}$ (218 pm, 213 pm, 207 pm) due to increased ligand field stabilization. In absence of other effects, this strengthening of the M$^{2+}$–O bond would lead to a blue-shift. The blue-shift in the order Mn$^{2+} < $ Fe$^{3+} < Ni$^{2+}$, that we in fact observe, is therefore the result of the counteracting effects due to mass reduction and bond strengthening. Using the relationship $k = (2\pi\nu)^2\mu$ between effective force constant $k$, vibrational frequency $\nu$ and reduced mass $\mu$, we can calculate effective force constants of 44 N m$^{-1}$, 52 N m$^{-1}$ and 65 N m$^{-1}$ for the vibrational mode of the hexaqua complexes of Mn$^{2+}$, Fe$^{2+}$ and Ni$^{2+}$, respectively.

By exchanging a water ligand by chloride, the reduced mass for the $\Theta$ vibration of the octahedral Fe$^{2+}$ complex is increased to 43.27 g mol$^{-1}$ (Cl$^-$ axial) or to 45.08 g mol$^{-1}$ (Cl$^-$ equatorial).
In absence of other effects, this increase would cause a red-shift of the $\Theta$ mode by ca. 4–6% or 6–9 cm$^{-1}$. In good agreement with this, we observe a red-shift of 6 cm$^{-1}$ to 142 cm$^{-1}$.

Moving from Fe$^{2+}$ to Fe$^{3+}$, the reduced mass of the mono-chloro complex remains the same, while the bond strength increases due to the higher charge of the metal center. Accordingly, the $\Theta$ mode is blue-shifted by 12 cm$^{-1}$ to 154 cm$^{-1}$. Exchanging another water molecule (in trans-position) by Cl$^-$, the reduced mass for the $\Theta$ vibration is increased to 45.68 g mol$^{-1}$ (Cl$^-$ axial) or to 48.98 g mol$^{-1}$ (Cl$^-$ equatorial). In absence of any other effect, this increase is expected to lead to a red-shift of 3–4% or 5–6 cm$^{-1}$. In fact, we observe a much larger red-shift of 12 cm$^{-1}$. We attribute this to a weakening of the bond strength by the partial charge compensation of the metal center by the negatively charged ligands.

### 6.2.2 Other ionic bands

In all spectra we observe a second absorption feature around 170–270 cm$^{-1}$ which is attributed to the cation. Possible assignments include: (a) The IR-active $\Psi$ mode described by Funkner et al., in which the central metal ion in an octahedral complex moves together with the four equatorial ligands against the two axial ligands; (b) a mode of the chloride ligand interacting with its hydration shell similar to the free anion; (c) a mode of hydration water (found at 157 cm$^{-1}$ for bulk water). Two observations are noteworthy: first, the second mode of Fe$^{2+}$ at 210 cm$^{-1}$ is almost identical with regard to center frequency, amplitude and width to the mode of Mn$^{2+}$ at 214 cm$^{-1}$. And second, comparing FeCl$^+$ to FeCl$^{2+}$, the second cation mode is red-shifted by 17 cm$^{-1}$ or 10%, which is very similar to the 8% red-shift of the first cation mode.

For Fe$^{2+}$ we fitted a small amplitude mode at fixed center frequency of 0 cm$^{-1}$, which we attribute to a relaxational process.

We observe only one band which can be attributed to the chloride anion in the spectra of Mn$^{2+}$ + 2Cl$^-$, Fe$^{2+}$ + 2Cl$^-$ and Ni$^{2+}$ + 2Cl$^-$ at 184 cm$^{-1}$, which is close to the chloride band around 190 cm$^{-1}$ found for other salts. Due to the increased complexity of the spectra of FeCl$^+$ + Cl$^-$, FeCl$^{2+}$ + 2Cl$^-$ and FeCl$_2$ + Cl$^-$, we decided to fix the values of the anion band to the values deduced for Fe$^{2+}$ + 2Cl$^-$ prior to the fit.

### 6.2.3 The linewidths

Since the spectra of Mn$^{2+}$ + 2Cl$^-$, Fe$^{2+}$ + 2Cl$^-$ and Ni$^{2+}$ + 2Cl$^-$ should lack any contributions of ion associates, we will call the bands in these spectra free ion bands. The linewidth for all free ion bands has been fitted as one parameter, which agrees well with the experimental spectra. The linewidth reflects the perturbation or damping of the vibrational modes of the ion–water complexes. Hence as a result we state that all free ions are subject to the same disturbance, which means that they are all connected to the same thermal bath.

The linewidth of the modes attributed to the chloro-complexes of Fe$^{2+}$ and Fe$^{3+}$ varies. The first and third band of FeCl$^+$ have the same width as the free ion bands (231 cm$^{-1}$), while the width of the second band of FeCl$^+$ is slightly larger (248 cm$^{-1}$). For all the other bands we observe a smaller width, ranging from 98 cm$^{-1}$ to 205 cm$^{-1}$. Distinct linewidths indicate that these modes are either connected to a different set of thermal bath states, or connected to the same bath, but with a different coupling parameter.

At this point, we can only speculate about the underlying molecular mechanism. We propose that the width of modes of solvated ions depends on the librational motions of the surrounding water molecules. These librations act as a random force causing line broadening. According to calculations by Vila Verde et al., water molecules between two ions (in case of a solvent-shared ion pair) or close to their point of contact (in case of a contact ion pair), are slowed down cooperatively. We therefore interpret the reduced linewidth as a consequence of inhibited librational motion of water close to the ion associates.

### 6.3 Hydration water

The low frequency mode at 117 cm$^{-1}$ (called WM in Table 3) is attributed to a hydration water mode. It is most likely either due to a shifted hindered translational mode of the first shell hydration water, or due to an increased line strength of a second shell mode, which is not IR active in a bulk water environment. In addition to this, there are high and low frequency components attributed to the relaxational and librational modes of hydration water, scaled by factors $n_{HF}$ and $n_{LF}$. Finally, there is a negative contribution to the effective ionic extinction in the shape of the bulk water spectrum scaled by a factor $n_{hyd}$. Since the relaxational and librational parts are compensated by $n_{LF}$ and $n_{HF}$, the negative contribution is dominated by the shifted translational band of hydration water. The scaling factor $n_{hyd}$ provides an estimate of the minimum number of water molecules affected by the ions.

For the extinction spectra of Mn$^{2+}$, Fe$^{2+}$ and Ni$^{2+}$ we can use the same fitting parameters for $n_{hydration}$ and $n_{LF}$, while the effect on the librational mode of water ($n_{HF}$) is slightly higher for Fe$^{2+}$ compared to Mn$^{2+}$ and Ni$^{2+}$, $n_{hydration}$ is in the range of 14–15. In a previous study we found that HCl affects ca. 5 water molecules and attributed this effect mainly to the anion. If we transfer this result to the present study, a minimum number of 4–5 water molecules are affected by the metal ions Mn$^{2+}$, Fe$^{2+}$ and Ni$^{2+}$, which is in good agreement with a value of six water molecules of an octahedral geometry.

For the ion-associate FeCl$^+$ + Cl$^-$, the value obtained for the minimum number of affected water molecules is $n_{hydration} = 7.3$, which is distinctly lower than for all other species. Assuming again a number of 5 water molecules affected by Cl$^-$, the chloro-complex affects only a minimum number of 2–3 water molecules. This can only be explained if both the ferrous and chloride ions lose part of their influence on hydration water upon ion pairing.

For FeCl$^{2+}$ + 2Cl$^-$ we obtain a lower limit of $n_{hydration} = 21.4$ affected water molecules, with similar values for the high and low frequency components ($n_{LF} = 23.2$ and $n_{HF} = 25.0$). Subtracting the proposed effect of the anions, we deduce that 11–12 water molecules are affected by FeCl$^{2+}$. This is surprisingly large, especially in comparison to FeCl$^+$, in which the Cl$^-$ ligand loses most of its effect on water upon ion pairing. We speculate that this indicates an effect beyond the first hydration shell of Fe$^{2+}$.

The minimum number of affected water molecules ($n_{hydration} = 16.9$) as well as the low and high frequency...
components \( n_{\text{lep}} = 20.3 \) and \( n_{\text{HF}} = 18.7 \) are smaller in the case of \( \text{FeCl}_2^+ + \text{Cl}^- \) compared to \( \text{FeCl}_2^{2+} + 2\text{Cl}^- \). This can be explained by a compensation, or an effective shielding, of the positive charge of the central metal ion by the two axial chloride ligands.

### 7 Quantitative analysis of FeCl\(_2\)/FeCl\(_3\) mixtures

As described in the previous sections, the absorption characteristics of FeCl\(_2\) and FeCl\(_3\) are distinctly different in the THz/FIR region. Their spectral signatures can therefore be used to distinguish ferrous and ferric iron species in aqueous solution.

Using Pitzer’s equations\(^{58}\) and the complex formation constants \( K_{\text{FeCl}_2} \) and \( K_{\text{FeCl}_3} \), the distribution of different ions and ion complex species can be predicted for FeCl\(_2\) and FeCl\(_3\) solutions of any concentration and even for any mixture of both salts (cf. Section 5). For a given water concentration, the extinction spectrum of water and the extinction spectra of the different complex species displayed in Fig. 4 can be used to predict the total absorption spectrum of any solution of FeCl\(_2\) and/or FeCl\(_3\) according to the following equation:

\[
x_{\text{tot}} = x_{\text{water}} + c_{\text{FeCl}_2} \left( x_{\text{FeCl}_2^+} + 2x_{\text{Cl}^-} \right) + c_{\text{FeCl}_3} \left( x_{\text{FeCl}_2^+} + x_{\text{Cl}^-} \right) + c_{\text{FeCl}_2^+} \left( 2x_{\text{FeCl}_2^+} + x_{\text{Cl}^-} \right) \tag{11}
\]

The concentrations \( c_x \) can be calculated as functions of density \( \rho \) and initial salt concentrations \( c_{\text{FeCl}_2}^{\text{init}} \) and \( c_{\text{FeCl}_3}^{\text{init}} \) of a given solution. It is therefore possible to determine the composition of an unknown mixed solution of FeCl\(_2\) and FeCl\(_3\) by fitting eqn (11) to the experimental absorption spectrum, using \( c_{\text{FeCl}_2}^{\text{init}} \) and \( c_{\text{FeCl}_3}^{\text{init}} \) as fitting parameters.

We have tested this method for a quantitative analysis of five sample solutions containing FeCl\(_2\) and FeCl\(_3\) in concentration ratios of 1 M/0 M, 0.75 M/0.25 M, 0.5 M/0.5 M, 0.25 M/0.75 M and 0 M/1 M. Each solution was acidified with 1 M HCl to prevent hydroxide formation and oxidation. The absorption spectra of these solutions after subtraction of the HCl contribution are presented in Fig. 6. The real concentrations (known from sample preparation) and measured concentrations (obtained from fitting) of FeCl\(_2\) and FeCl\(_3\) are plotted in Fig. 7.

### 8 Summary and conclusion

We have measured the THz/FIR absorption spectra of acidified aqueous FeCl\(_2\) and FeCl\(_3\) solutions at concentrations of 0.5 to 3.5 M in a frequency range of 30–350 cm\(^{-1}\). We observed a non-linear concentration dependence of the effective ionic extinction for both salts. This can be rationalized in terms of the formation of chloro-complexes of Fe(II) and Fe(III), respectively. We were able to dissect the spectra of FeCl\(_2\) into the contributions of single ions and ion pair and the spectra of FeCl\(_3\) into the contributions of ion pair and ion triplet using principal component analysis. In addition, we were able to determine the ion association constants for both salts.

The logarithm of the formation constant of the ion pair FeCl\(^+\) determined this way is \(-0.89(4)\); the logarithm of the formation constant of the ion triplet FeCl\(_2^+\) is \(-0.32(8)\). Both values are considerably lower than the values found in literature.\(^{17,26,28,33,42–44,49,51}\) However, we have to keep in mind that our measurements are susceptible mainly to the formation of contact ion pairs, while other methods might also be sensitive towards the formation of solvent separated or solvent shared ion pairs.

From the PCA of the FeCl\(_3\) spectra we extracted the molar extinction spectra of Fe\(^{2+}\) + 2Cl\(^-\) and FeCl\(^+\) + Cl\(^-\); from the PCA of the FeCl\(_2\) spectra we extracted the molar extinction spectra of FeCl\(_2^{2+}\) + 2Cl\(^-\) and FeCl\(_2^+\) + Cl\(^-\). We performed a global fit of all the extinction spectra and their differences, including the previous results for manganese(II) and nickel(II) chlorides and bromides.\(^{52}\)

For all cations we observe a peak centered around 150 cm\(^{-1}\), which we assign to the \( \Theta \) vibration of the octahedral aqua- and chloro-complexes. This peak shifts for different cations, which can be explained by changes in reduced mass and bond strength of the complexes.

\( \text{Mn}^{2+} + 2\text{Cl}^- \), \( \text{Ni}^{2+} + 2\text{Cl}^- \) and \( \text{Fe}^{2+} + 2\text{Cl}^- \) affect approximately the same amount of water, with a minimum number of affected water molecules of 14–15. The ion pair FeCl\(^+\) + Cl\(^-\) affects only half as much water, which we attribute to the
charge compensation of the paired ions. The ion pair FeCl$_2^+ + 2$Cl$^-$, on the other hand, affects a much larger number of water molecules (ca. 21), which is an indication for a hydration effect extending beyond the first hydration shell of Fe$^{3+}$. Again, the minimum number of affected water molecules is reduced upon association of another chloride ion to ca. 17 in the case of FeCl$_2^+ + $Cl$^-$. Furthermore, our analysis could be used successfully for a quantitative determination of concentrations of FeCl$_2$ and FeCl$_3$ in mixed solutions of total salt concentration of 1 M with errors of less than 0.08 M. Thus we have provided a proof of principle that this technique can be used as an analytical tool for highly concentrated salt solutions.

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