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■ Pressure-induced ion pairing in MgSO₄ solutions: Implications for the oceans of icy worlds

C. Schmidt¹*, C.E. Manning²

Abstract

At ambient temperature, liquid water transforms from a low-density to a high-density dynamic structure at $\sim\!0.2$ GPa. The transition persists in electrolyte solutions; however, its effects on solute properties are unknown. We obtained Raman spectra of 0.5–2.0 molal MgSO4 solutions at 21 °C and 10 -4 to ~1.6 GPa. Above about 0.4 GPa, we observed an increase in the MgSO4 contact ion pair abundance with pressure, regardless of concentration. This phenomenon contravenes the general rule that dissolved salts dissociate upon compression, and is likely caused by the structural collapse in the solvent with pressure due to increased hydrogen-bond breaking. Increasing ion association in high-pressure aqueous solutions implies that, at a given salinity, high-density water in deep, cold planetary oceans and pore waters will possess lower ionic strength and electric conductivity than previously thought. This behaviour will also lead to higher ocean salinity in the interiors of Pluto and the largest icy moons of Jupiter and Saturn, Ganymede, Callisto, and Titan, or in exoplanet water-worlds, through enhancement of submarine silicate weathering.

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Introduction

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The physical properties of liquid water at ambient temperature consistently display anomalous behaviour upon compression to densities of ~1.1 g cm⁻³. These include *e.g.*, radial distribution functions from X-ray scattering (Okhulkov *et al.*, 1994) and neutron diffraction (Soper and Ricci, 2000), a distinct anomaly in the high-frequency sound velocity from inelastic X-ray scattering experiments (Krisch *et al.*, 2002), the Raman shift of the O–H stretching vibration (Kawamoto *et al.*, 2004), the sound velocity from Brillouin spectroscopy (Li *et al.*, 2005),

and the isothermal compressibility (Mirwald, 2005). The experiments indicate a transition from an open structure in low-density water (LDW) to a structure with a collapsed second coordination shell in high-density water (HDW) (Soper and Ricci, 2000). Based on femtosecond infrared pump-probe spectroscopy on a D_2O-H_2O mixture, the LDW-HDW transition is at \sim 0.25 GPa at 273 K and \sim 0.2 GPa at 298 K, but not observed at 363 K (Fanetti *et al.*, 2014).

Dissolved salts also disrupt the hydrogen-bonded dynamic H_2O structure due to formation of hydration shells around the ions. It is not clear if, and by how much, the structure outside the first hydration shell is affected (Marcus, 2009), but there is considerable evidence that the LDW-HDW transition persists in saline solutions even at fairly high solute concentrations (Mirwald, 2005; Schmidt, 2009; Valenti *et al.*, 2012). However, the effects of the transition on solute-solute interactions have not been investigated. Here, we studied the pressure-dependent variation in contact ion pairing in MgSO₄ solutions at 21 °C over a large range in pressure across the LDW-HDW transition until the sample solidified upon compression. We used MgSO₄ solutions because they are thought to be important components of planetary oceans (Zolotov and Shock, 2001) and because Raman scattering from the symmetric stretching vibration of sulphate (v_1 -SO₄²⁻) provides direct information about the fraction of contact ion pairs.

Compression-Enhanced Contact Ion Pairing

At all conditions, the Raman band shape in the v_1 -SO $_4^{2-}$ region was accurately described using two Gaussian + Lorentzian components (Fig. S-1a). The higher wavenumber component is assigned to contact ion pairs Mg^2 +SO $_4^{2-}$ (aq) (CIP), and the lower wavenumber component, here designated SO $_4^{*-}$, includes Ramanindistinguishable contributions from unassociated SO $_4^{2-}$ (aq) ions, solvent-shared ion pairs Mg^2 +(OH $_2$)SO $_4^{2-}$ (aq) (SIP), and double solvent-separated ion pairs Mg^2 +(OH $_2$)SO $_4^{2-}$ (aq) (2SIP) (Rudolph *et al.*, 2003). Spectra in the O–H stretching region were recorded for the 2.0 molal solution. A three component model yielded good fits with internally consistent results for all spectra (Fig. S-1b). We assign the components C2 and C3 at ~3450 cm $^{-1}$ and ~3580 cm $^{-1}$ to v_1 and v_3 of H $_2$ O monomers (*e.g.*, Walrafen, 1962). The component C1 at ~3280 cm $^{-1}$ is assigned to O–H stretching vibrations of strongly hydrogen-bonded, polymerised water species because the relative intensity of this component correlates with the dependence of polymerisation on pressure and temperature (*e.g.*, Sahle *et al.*, 2013) and decreases with addition of salt (*e.g.*, Walrafen, 1962).

In our experiments at $21 \pm 1\,^{\circ}\text{C}$, we obtained data for the v_1 -SO₄²⁻ Raman shift of the SO₄* and CIP components (Figs. 1a and 1b), the abundance of contact ion pairs, as calculated from the relative integrated intensities of the CIP and SO₄* components (Fig. 1c), and the Raman shifts of the three band components in the O–H stretching region (Fig. 1d). At each studied concentration of 0.5, 1.7 and 2.0 molal MgSO₄, increasing pressure produces linear changes in the v_1 -SO₄²⁻ Raman shift of SO₄* of ~6 cm⁻¹GPa⁻¹ for the data to 0.17 GPa (Fig. 1a).



Deutsches GeoForschungsZentrum (GFZ), Section 4.3 Chemistry and Physics of Earth Materials, Telegrafenberg, 14473 Potsdam, Germany

^{*} Corresponding author (email: Christian.Schmidt@gfz-potsdam.de)

Department of Earth, Planetary, and Space Sciences, University of California, Los Angeles, CA 90025-1567. USA

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Above 0.7 GPa, the pressure dependence of that Raman shift is also linear and independent of composition, but the slope is shallower (\sim 3 cm $^{-1}$ GPa $^{-1}$). All data show systematic negative departures from linearity beginning at 0.17 GPa, which corresponds to the LDW-HDW transition (Fig. 1a) and agrees with a previous study on NaSO₄ solutions (Schmidt, 2009). Departures from linearity continue to \sim 0.6 GPa. Similar behaviour is observed in the Raman shift of the CIP component in the same pressure interval (Fig. 1b).

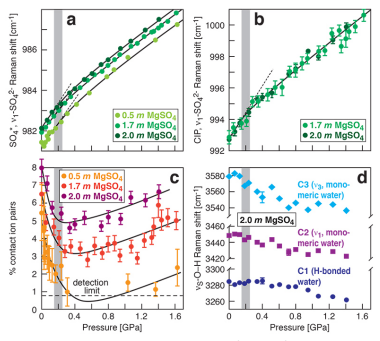


Figure 1 Raman spectroscopic results at 21 \pm 1°C as a function of pressure. The shadowed band is the pressure range of the LDW-HDW transition at about 20°C (Fanetti *et al.*, 2014). (a-b) Raman shift ω of the v_1 -SO $_4$ ²⁻ mode of the SO $_4$ * component (a) and of the CIP component (b). Dashed lines denote the $(\partial \omega/\partial P)_{T=21^{\circ}C}$ slopes in cm $^{-1}$ GPa $^{-1}$ from linear fits of data at <0.17 GPa. (c) Percentage of contact ion pairs as a function of pressure at 21°C. (d) Raman shift of fitted band components in the v_5 -O-H region of the 2.0 molal MgSO $_4$ solution.

At 21 °C, the percentage of ${\rm Mg^{2+}SO_4^{2-}}(aq)$ CIP initially decreases with pressure along the isotherms at each concentration (Fig. 1c). Above ~0.4 GPa, however, it increases nearly linearly with rising pressure in 1.7 and 2.0 molal solutions. The 0.5 molal solution is consistent with this behaviour, although errors are larger and the abundance of CIPs is below the detection limit at about 0.4–0.9 GPa. This was not observed at 50–150 °C, where the percentage of CIP in 0.75 and 2.25 molal MgSO₄ solutions generally decreased with pressure along all

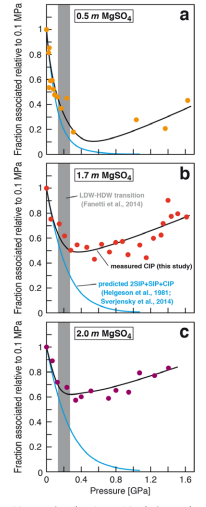


Figure 2 Fraction of MgSO₄ associated at 21 \pm 1°C relative to that at ambient pressure, as determined by Raman spectroscopy and thermodynamic modelling, for 0.5 (a), 1.7 (b), and 2 molal solutions (c). Thick black lines are trend lines for the fraction of CIP from Raman spectroscopic data fit by eye. Blue lines denote the fraction of the sum of the CIP, SIP and 2SIP ion pairs calculated from a thermodynamic model based on the DEW model (Sverjensky et al., 2014) and the Helgeson-Kirkham-Flowers ion activity model (Helgeson et al., 1981). The comparison assumes that the proportionality between equilibrium constants associated with expelling the H₂O molecules between ions is constant and independent of pressure. This assumption yields a minimum difference in predicted and observed CIP. The shadowed band is the pressure range of the LDW-HDW transition at about 20°C (Fanetti et al., 2014).



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isotherms (Fig. S-2). The data for the pressure dependence of the v_S -O–H Raman shift for C1, C2 and C3 in the 2.0 molal solution generally suggest declining frequency with rising pressure (Fig. 1d), consistent with lengthening of the O–H-bonds and thus shorter hydrogen bonds between the water molecules. The scatter in these data is fairly large because the v_S -O–H components are very broad and overlap (Fig. S-1b).

The finding of compression-enhanced contact ion pairing above ~0.2 GPa at 21 °C (Fig. 1c) is contrary to the predicted trend of increasing salt dissociation with pressure. Previous Raman spectroscopic and electrical conductivity studies (Ritzert and Franck, 1968; Chatterjee et al., 1974) at less than about 0.5 GPa and near ambient temperatures indicated increasing dissociation of MgSO₄ in aqueous solution upon compression. Thermodynamic modelling also yields a continually increasing extent of ion pair dissociation with rising pressure, unaffected by the LDW-HDW transition (Fig. 2), but data used in such models are based on correlations and extrapolations typically derived from conductivity data at very different pressures and temperatures. The static dielectric constant of H₂O is a key parameter governing the extent of ion association in aqueous solutions; however, it does not decrease with increasing pressure in the HDW field (Fernandez et al., 1995). Based on Coulomb's law, the observed increased contact ion pairing must therefore be caused by a decrease in the distance between cation and anion. An increasing tendency to form ion pairs with pressure can thus be related to increased hydrogen-bond breaking with pressure resulting in a collapsed second coordination shell in HDW (Soper and Ricci, 2000; Krisch et al., 2002; Fanetti et al., 2014). Therefore, the inferred relationship between water structure and ion pairing implies that significant interaction between ions and water exists at distances larger than the first shell, which has long been a matter of debate (c.f. Marcus, 2009).

Implications for Planetary Oceans and Interiors

Generally, the increase in ion pairing with pressure affects properties such as solubility, viscosity, sound absorption and electrical conductance. For a given salinity, isothermal salt re-association will yield lower ionic strength (and thus ion activity) than predicted from extrapolation of low-pressure behaviour, and also lower conductivity of deep ocean waters, which is relevant for the interpretation of spacecraft-based magnetometric data (e.g., Hand and Chyba, 2007).

Our results show that high pressures and fairly low temperatures are required for pressure-induced ion pairing in salty water. In nature, it will therefore only occur in relatively large, cool planetary bodies with thick liquid water shells. Extrasolar ice planets may well meet these requirements, as well as some ocean planets such as Kepler-62e and -62f (Kaltenegger *et al.*, 2013). So too would cold, deep, aqueous planetary settings in our solar system, which include the interiors of Pluto and the largest icy moons of Jupiter and Saturn. There is considerable evidence for a subsurface ocean in Europa, but plausible basal ocean pressure is too low (Spohn and Schubert, 2003; Hand and Chyba, 2007) for salt

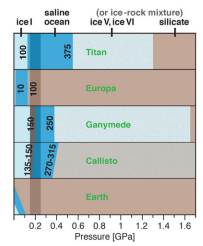


Figure 3 Comparison of models of the interiors of Titan (Tobie *et al.*, 2012), Europa (Spohn and Schubert, 2003; Hand and Chyba, 2007; Valenti *et al.*, 2012), Ganymede (Vance *et al.*, 2014; Saur *et al.*, 2015), and Callisto (Kuskov and Kronrod, 2005) with approximate depths of the subsurface ocean boundaries in km. The shadowed band is the pressure range of the LDW-HDW transition at about 20°C (Fanetti *et al.*, 2014).

re-association (Fig. 3). However, Ganymede (Vance *et al.*, 2014; Saur *et al.*, 2015) and Saturn's moon Titan (Grasset and Pargamin, 2005; Tobie *et al.*, 2012) likely possess deep subsurface oceans with substantial portions of the liquid water column in the LDW-HDW transition region (Fig. 3). Deep ocean waters of Pluto, and an at least partly differentiated Callisto, would also exceed 0.2 GPa (Spohn and Schubert, 2003; Kuskov and Kronrod, 2005; Hammond *et al.*, 2016). As in Europa, the existence of a thick saline subsurface ocean in Callisto is indicated by a conductive layer at a depth of less than about 300 km inferred from magnetic field perturbations observed by the Galileo spacecraft (Zimmer *et al.*, 2000).

In this context, our finding of increasing ion association with pressure is important for understanding the coupled evolution of ocean salinity and the silicate interiors of the icy satellites. The salts of planetary oceans are acquired chiefly from sub-seafloor weathering reactions (Zolotov and Shock, 2001). About 75 % of the salts produced by low-temperature aqueous alteration of chondrite under oxidising conditions are magnesium sulphate hydrates (Hogenboom *et al.*, 1995). Thus, $H_2O + MgSO_4$ is one of the most relevant model systems for planetary oceans. Consumption of ions by ion pair formation via

$$Mg^{2+} + SO_4^{2-} = MgSO_4^0$$
 Eq. 1

requires commensurate forward progress of any magnesium silicate weathering reactions, such as olivine:

$$Mg_2SiO_4 + 2H^+ = Mg^{2+} + 2SiO_2 + 2H_2O$$
 Eq. 2



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Therefore, the greater extent of salt association in HDW pore waters beneath the ocean floors of large icy satellites will enhance the progress of dissolution reactions relative to expectations based on LDW. This will generate a more voluminous planetary reservoir of altered rock, and – when the pore fluids decompress upon delivery to the oceans – higher ocean salinities. A thick layer of high-pressure ices and hydrates on the floor of a deep cold subsurface ocean (Fig. 3) could decrease this effect, but dissolved salts would diminish such a layer. Ganymede might have a stack of several ocean layers separated by different phases of ice, with the lowest liquid layer adjacent to the rocky mantle below (Vance *et al.*, 2014).

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Additional Information

Supplementary Information accompanies this letter at www.geochemicalperspectivesletters.org/article1707

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C. Schmidt¹*, C.E. Manning²

Supplementary Information

The Supplementary Information includes:

- ➤ Methods
- Discussion of Validity of Results
- ➤ Discussion of Results at 50 to 150 °C
- Supplementary Information References

Methods

An externally heated Bassett-type HDAC (Bassett *et al.*, 1993) equipped with type Ia ultra-low fluorescence and ultra-low birefringence diamond anvils (culet diameter 0.9 mm) was used for all experiments. The temperature in the sample chamber of the cell was measured using K-type thermocouples attached to the diamonds. Temperature was calibrated by measurement of two phase-transition temperatures, water ice melting in presence of liquid and vapour (triple point at 0.01 °C and 0.6 kPa) and the transformation from α -quartz to β -quartz (574 °C at 0.1 MPa). Accuracy and reproducibility of the temperature measurements were about ± 0.1 °C between -5 and 0 °C (for measurement of the vapour-saturated liquidus temperature of the MgSO₄ solution in the sample chamber) and about ± 0.5 °C for experiments at 50 to 150 °C. The power input to the resistive heaters was controlled using Eurotherm® 2408 temperature controllers, which held the set temperature within ± 0.2 °C during acquisition of Raman spectra. A set of MgSO₄ solutions of varying composition was prepared from chromatographic grade H₂O (LiChrosolv®, Merck) and analytic grade magnesium sulphate



Deutsches GeoForschungsZentrum (GFZ), Section 4.3 Chemistry and Physics of Earth Materials, Telegrafenberg, 14473 Potsdam, Germany

^{*} Corresponding author (email: Christian.Schmidt@gfz-potsdam.de)

Department of Earth, Planetary, and Space Sciences, University of California, Los Angeles, CA 90025-1567, USA

heptahydrate (Merck). For a given experimental series, a piece of natural quartz was loaded with a drop of solution into the HDAC sample chamber formed by a hole with a diameter of 300 or 400 µm in a rhenium gasket separating the anvils. To minimise background fluorescence, care was taken during preparation and loading to avoid contamination by organic materials, particularly by residue made by fingers touching a surface. The procedure for loading and sealing the sample chamber led to minor but unavoidable H₂O evaporation, so the MgSO₄ concentration in the solution composition needed to be re-determined. This was done by measurement of the vapour-saturated liquidus temperature (cryometry). First, prior to sealing, an air bubble was allowed to grow to about 10 % of the liquid volume by controlled leaking of the chamber. The sample chamber was then sealed by slight compression of the gasket between the anvils, and the sample was frozen. The ice or MgSO₄ hydrate melting temperature in the presence of vapour was then determined, giving the MgSO₄ concentration from comparison to literature data (Wolf, 1966; Hogenboom et al., 1995; Marion and Farren, 1999). Magnesium sulphate concentrations determined by cryometry were checked by comparison with concentrations calculated from the full width at half maximum of the fitted SO₄* component of the v₁-SO₄²- Raman band (Fig. S-1a), calibrated using MgSO₄ standard solutions in glass containers. The results from both techniques were in very good agreement. The error in the reported MgSO₄ concentrations is ±0.05 molal. For MgSO₄ concentrations of 0.5, 1.7, and 2.0 molal, the experiments were performed along the 21 °C isotherm with stepwise increase of the pressure by compression of the sample chamber until solids formed. At each pressure step, Raman spectra of the fluid were collected in the region of the v_1 -SO₄²⁻ mode and additionally in the region of the O–H stretching vibrations of water in the 2.0 molal MgSO₄ experiments. In the runs along the 21°C isotherm, the pressure was determined by recording Raman spectra of quartz before and after those of the aqueous solution and using the calibrated frequency shift of the 206 cm⁻¹ Raman line of quartz and equation 5 in Schmidt and Ziemann (2000). Two additional experimental series were performed on 0.75 and 2.25 molal MgSO₄ solutions along several isochoric or quasi-isochoric paths at 50, 75, 100, 125, and 150 °C. In these runs, pressures above the vapour pressures of the solutions were obtained from the calibrated frequency shift of the 464 cm⁻¹ Raman line of quartz using the equations 2 and 3 in Schmidt and Ziemann (2000). The random error in the obtained pressures was about 10 MPa if the 206 cm⁻¹ Raman line of quartz was used, or about 25 MPa in the case of the 464 cm $^{-1}$ line. The experiments at elevated temperatures were restricted to T \leq 150 °C to avoid precipitation of kieserite (MgSO₄·H₂O) (Hogenboom *et al.*, 1995; Jahn and Schmidt, 2010) and liquid-liquid phase separation (Wang et al., 2013), and terminated upon gasket failure. The unpolarised Raman spectra of solutions and quartz were acquired using a HORIBA Jobin Yvon LabRAM HR800 UV-Vis Raman microprobe (gratings 1800 lines/mm, focal length 800 mm) equipped with a thermoelectrically cooled CCD-detector (1024 × 256 pixel). The spectra were obtained in backscattering geometry with a Nikon MPlan SLWD 40× objective (numerical aperture 0.4), a confocal pinhole aperture of 100 µm, and using the 488.0 nm argon ion laser line for excitation. The laser interference filter was

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removed to record plasma lines from the Ar+ laser and the Raman lines in the same spectrum. The frequency shift of the quartz lines was calibrated using the plasma lines at 221.54, 351.60, and 520.30 cm⁻¹ and the frequencies of the v_1 -SO₄²⁻ band components by monitoring the plasma lines at 737.28 and 1056.94 cm⁻¹. The laser power was set to 275 mW at the source, which was low enough to avoid laser heating of the sample in the HDAC (Schmidt, 2009). Raman spectra of the solutions were collected with 20 accumulations of 5 s each to minimise spectral noise and to attain low and linear backgrounds of similar intensity. Before acquisition of these spectra, the nominal focal point position was always set to 20 µm below upper sample chamber surface to obtain comparable integrated intensities of the Raman bands (Everall, 2000; Schmidt and Chou, 2012). Spectral fitting was done using the software package PeakFit v4.11 from SYSTAT Software Inc., a linear baseline, and the Gaussian-Lorentzian sum (area) function with the same shape factor (i.e. the same degree of Lorentzian character) for all peaks in an individual fit of the two v_1 -SO₄²⁻ components (Rudolph *et al.*, 2003) and the three components in the O-H stretching region of water (Brubach et al., 2005). The Pearson IV model was used for the Raman lines of quartz and the plasma lines to allow for peak asymmetry (Schmidt and Ziemann, 2000; Schmidt, 2009). This procedure resulted in a very good description of the shape of the recorded Raman bands, i.e. non-statistical residuals were insignificant in all cases. The signal-to-noise ratio in the Raman spectra did not permit reliable fitting of additional components in the O–H stretching region with consistent results.

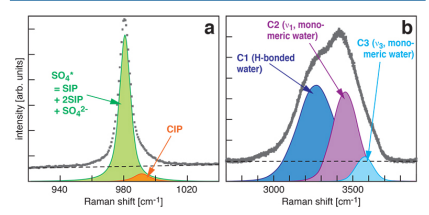


Figure S-1 Raman spectra of a 2.0 molal MgSO₄ solution at 21 °C, 0.1 MPa. The fitted baselines are shown as dashed lines. (a) v₁-SO₄² region, with two fitted band components assigned to CIP (contact ion pairs) and SO₄* (including solvent-separated ion pairs (SIP, 2SIP) and unassociated SO₄²⁻ ions). (b) O-H stretching region, with three fitted band components, C1 (mostly from hydrogen-bonded water), C2 (assigned to the symmetric stretching vibration, v₁, of H₂O monomers), and C3 (assigned to the antisymmetric stretching vibration, v₃, of H₂O monomers).



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Discussion of Validity of Results

Our observations point to a previously unrecognised tendency for progressively increasing contact ion pairing with rising pressure in high-density water at ambient temperature. Alternative explanations are not viable. The internal consistency of all collected data at three concentrations argues strongly against an artefact from hypothetical water loss during the experiments and against any non-isoplethic behaviour. Dissolved silica or rhenium can likewise be excluded as causes because the solubilities of Re and quartz (Manning, 1994; Xiong and Wood, 1999) are many orders of magnitude lower than the magnesium sulphate concentration at all pressure-temperature conditions of this study, and no rhenium or other sulphate was observed to form in the sample chamber. An artefact due to variable and nonlinear background can be excluded because all spectra were fitted using the same procedure and had low and very near linear backgrounds in the frequency region of the v_1 -SO₄²⁻ Raman mode. The fluorescence contribution from the solution or the diamond anvils was insignificant. Fitting of spectra in which the measured background of the diamond anvil was subtracted from the spectrum of the solution yielded results that were almost identical. The enhanced contact ion-pairing shown in Figure 1 could have been the result of an increase in temperature by about 10 to 15 °C during isothermal compression, but monitored temperature changes were <1 °C. Laser heating could have increased the percentage of CIP, but the determined fractions of CIP in the solutions in the HDAC at 0.1 MPa are even slightly lower than literature data (Rudolph et al., 2003) from Raman spectra acquired at much higher laser power at the sample. Finally, significantly different pressure dependencies of the v_1 -SO₄²⁻ Raman scattering cross sections of CIP and SO₄* are implausible because they stem from the same vibrational mode of SO_4^{2-} . Furthermore, this would be in conflict with the fact that equal Raman scattering coefficients of CIP and SO₄* at ~25 °C, 0.1 MPa are verified by the close agreement of Raman spectroscopic speciation data for MgSO₄ solutions with data from electrical conductivity measurements and dielectric relaxation and ultrasound absorption spectroscopy (Fisher, 1962; Davis and Oliver, 1973; Rull et al., 1994; Tomšič et al., 2002; Rudolph et al., 2003; Buchner et al., 2004; Akilan et al., 2006).

Discussion of Results at 50 to 150°C

For 0.75 and 2.25 molal MgSO₄ solutions, we determined the variation in the percentage of Mg²⁺SO₄²⁻(aq) contact ion pairs with pressure at 50, 75, 100, 125, and $150\,^{\circ}$ C (Fig. S-2). The percentage of CIP generally decreased with pressure along these isotherms. Although the slopes become considerably less negative with pressure, there is no clear evidence for increasing ion association for any isotherm. If our observation of pressure-induced salt association is related to the LDW-HDW transition, then the high temperature data imply that HDW stability either disappears with increasing temperature or moves rapidly to high pressure. Raman, infrared, and Brillouin spectroscopic studies show that the LDW-HDW transition becomes increasingly difficult to detect above ambient temperature,

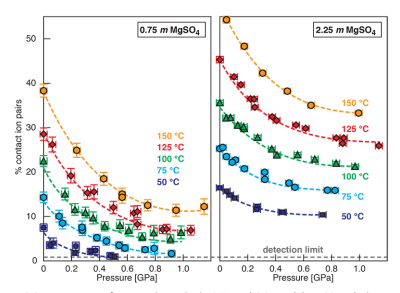


Figure S-2 Percentage of contact ion pairs in 0.75 and 2.25 molal MgSO₄ solutions as a function of pressure along isotherms (dark blue, 50°C; cyan, 75°C; green, 100°C; red, 125°C; orange, 150°C). Vertical error bars, 2 standard errors; horizontal error bars, uncertainty in the pressure obtained from the shift in the wavenumber of the 464 cm⁻¹ Raman line of quartz.

even at 50°C (Li et al., 2005; Okada et al., 2005; Schmidt, 2009; Fanetti et al., 2014b). There is no consensus on the $\partial P/\partial T$ slope of the LDW-HDW transition. The available experimental constraints variously suggest slopes that are strongly positive $(0.4 \pm 0.1 \text{ GPa at } 25 \,^{\circ}\text{C}, 1.0 \pm 0.1 \text{ GPa at } 100 \,^{\circ}\text{C}, \text{ and } 1.3 \pm 0.1 \text{ GPa at}$ 300 °C) (Kawamoto et al., 2004), to slightly negative (0.29 GPa at 293 K, 0.21 GPa at 316 K, and 0.19 GPa at 353 K) (Li et al., 2005), to negative (~0.2 GPa at 273.7 K, ~0.16 GPa at 298.0 K, ~0.11 GPa at 318.0 K, and ~0.08 GPa at 339.0 K) with the interpretation that LDW ceases to exist at ~363 K (Fanetti et al., 2014b). Although the most recent data of Fanetti et al., (2014b) appear to be of high quality, it should be noted that the implied change in the solvent structure at ~363 K would affect volume and entropy of reactions. However, to our knowledge, there are no anomalies in vapour pressure or aqueous solubility curves at about this temperature. At lower temperatures, anomalies in reaction curves exist at pressures near the LDW transition, e.g., a change from a negative to a positive $\partial P/\partial T$ slope of the hydrohalite liquidus at -7.5 °C and 140 ± 40 MPa (Valenti et al., 2013) and an inflection in the dehydration curve of NaCl-2H₂O at 12 °C, 0.27 GPa (Mirwald, 2005). Moreover, the minimum in the isothermal compressibility of water is at 46.5 °C. Above this temperature, water behaves like a regular liquid, which would be incompatible with a transition from more compressible LDW to less compressible HDW (Li et al., 2005) at about 90 °C. Furthermore, for the v_1 -SO₄²⁻ Raman band of a 1.54 molal Na₂SO₄ solution, the slopes of the wavenumber with



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pressure at 100, 200, 300, 400, and $500\,^{\circ}\text{C}$ are similar to that of LDW at $21\,^{\circ}\text{C}$, and the slopes of the line width with pressure at 100, 200, 300, 400, and $500\,^{\circ}\text{C}$ are much closer to that of LDW than that of HDW at $21\,^{\circ}\text{C}$ (Schmidt, 2009). In summary, the considerations above are consistent with our interpretation that a relationship exists between ion paring behaviour and the LDW-HDW transition, and that this behaviour is a low-temperature phenomenon.

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