Anionic and cationic Hofmeister effects are non-additive for guanidinium salts†

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To understand specific ion effects on a molecular level we explore the effect of salts on the rotational mobility of a model amide using dielectric spectroscopy. Based on our previous studies on the effect of strong denaturing anions or cations, here we study the additivity of the anionic and cationic effect. Using salts consisting of denaturing spherical anions and spherical cations we find such salts to affect the amide according to what one expects based on the additive activity of the individual ions. The guanidinium (Gdm+) cation appears to be a notable exception, as our results suggest that GdmI (and accordingly GdmSCN) is less efficient in hindering the rotation of the amide than KI or GdmCl.

Understanding the fundamentals of specific ion effect on the stability of proteins is a longstanding enigma in biophysical chemistry.1–3 The first systematic study on the effect of ions on biomolecules dates back to 1888, when Hofmeister ordered different salts according to their efficiency to stabilize or denature proteins.4 From these experiments it was concluded that salts containing cations with a high surface charge density and/or anions with a low surface charge density tend to destabilize proteins in solution.5,6 Experimentally several other macroscopic properties, e.g. the stability of colloids in solution,7 the surface propensity of ions8 and the surface potential of charged surfaces in contact with a salt solution,9 have been shown to follow trends according to the Hofmeister series. This intriguing observation has inspired many physico-chemical studies to explore the molecular-level origins of these specific ion-effects,10–15 since specific ion effects are commonly used in biotechnological and pharmaceutical applications to reversibly un-fold and fold proteins.16 However, despite the seminal work of Hofmeister describing the effect of salts, the overall effect is often separated into individual contributions from the anion and cation, as anionic and cationic Hofmeisters series, though it has remained unexplored whether specific ion effects can be attributed to the properties to individual ions or are characteristic to a salt, i.e. an anion–cation pair.

In recent years increasing evidence has emerged that specific ion effects on biomolecules originate from direct interaction between ions and proteins.16–21 The most prominent interaction site for binding of ions to proteins is the amide backbone, the main structural motif common to all proteins. Various – predominantly spectroscopic studies – have shown that interaction of ions with amide groups follow the Hofmeister series13,19–21 and it has thus been concluded that ion–amide interaction explains part of the observed destabilization of proteins. We have recently shown that the rotational mobility of a model amide, N-methylacetamide (NMA) is a very sensitive reporter of the interaction of ions with amide groups.19–21 Importantly, the rotational mobility can probe the interaction of the amide with both anions and cations – as opposed to other techniques that probe either cationic or anionic interaction sites.21 Consequently, using the sensitivity to both ions allows us to address the question if the interaction of the anion and the cation with NMA is additive. In the present study we explore the effect of salts, for which both the anion and the cation tend to denature proteins, on the rotational mobility of NMA. Comparison to our earlier results focusing on either the cationic or anionic effects (by choosing a non- or weakly interacting counter-ion) allows determining whether Hofmeister effects are specific ion effects or specific salt effects.

To investigate additivity of anionic and cationic specific ion effects, we use aqueous mixtures with a constant concentration of NMA (2 mol L⁻¹) with an increasing concentration of salt, c_salt = 0 to 2 mol L⁻¹. Here we chose a series of salts that consist of ions with both anion and cation reducing the rotational mobility of NMA: the monovalent salts NaBr, NaI, bivalent MgI₂ together with two guanidinium (Gdm⁺) salts GdmI and GdmSCN and compare these to our earlier results where we used the weakly interacting K⁺ and Cl⁻ as counter-ions.19–24

To quantify the interaction of the salts with NMA we use dielectric relaxation spectroscopy (DRS) at frequencies ranging from 0.8 to 1600 GHz21 (for experimental details see ESI†). DRS
probes the polarization of a sample as a response to an oscillating electric field with field frequency $\nu$. At the frequencies of this study the polarization of liquid samples predominantly stems from the rotation of dipolar molecules in the externally applied field. The polarization is measured in terms of the complex permittivity $\varepsilon(\nu) = \varepsilon'(\nu) - i\varepsilon''(\nu)$, with the real part $\varepsilon'$ being the dielectric permittivity and the imaginary part $\varepsilon''$ the dielectric loss. In Fig. 1a we show the dielectric spectra (corrected for the conductivity of the solutions) for a series of ternary solutions consisting of water + NMA + GdmI. Spectra for other salts are shown in the ESI† Fig. S1. All spectra exhibit a broad dispersion in $\varepsilon'$ and a corresponding peak in $\varepsilon''$ at ~10 GHz, which is due to the rotation of both dipolar NMA and water molecules. With increasing salt concentration the static permittivity of the samples (the limiting value of $\varepsilon''$ at low frequencies) decreases and simultaneously the amplitude of the dielectric loss peak (the overall dielectric strength) decreases: addition of salt leads to depolarization of the sample.

As discussed in our earlier studies,¹⁹-²¹ this depolarization has three different molecular-level origins: (i) reduction of the molar concentration of water, (ii) formation of rigid solvation shells around ions, and (iii) kinetic depolarization (KD). While (i) can be readily accounted for by using the reduced water concentration in our analysis and (ii) has only significant contributions for Na⁺ and Mg²⁺ ions (see below and details of the analysis in the ESI†), KD (iii) is relevant to all samples and dominates the depolarization for the present samples. KD results from the coupling between the translation of an ion with the rotation of a dipole in the external electric field. According to the first description of KD by Hubbard and Onsager,²² the magnitude of the depolarization scales with the medium’s conductivity (ion mobility) and the dipolar relaxation time. More recent studies on KD have indicated that at high concentrations of salt the magnitude of KD may deviate from this simple scaling due to screening of the ionic charges.²³ In addition to these purely electrostatic considerations for kinetic depolarization, we have recently demonstrated that for aqueous solutions of NMA the depolarization is also ion-specific.²⁹-³¹ The reduction of the static permittivity not only depends on the concentration of the salt and the mobility of the ions (conductivity) but also depends on the chemical identity of the ions: in line with the Hofmeister series, ions with an increasing protein denaturation tendency also increasingly depolarize aqueous NMA solutions. This ion specificity can be explained from the increasing proximity of these ions to NMA, as KD of NMA results in a larger depolarization as compared to KD of water.²⁹

To explore whether these specific effects of anions and cations on the depolarization – and thus the interaction between anions and cations with NMA – are additive, we quantify the depolarization using a dielectric relaxation model to describe the experimental spectra. In-line with our previous studies,¹⁹-²¹ we fit a combination of three relaxation modes to the experimental spectra. The rotational diffusion of NMA centred at ~4 GHz is modelled with a Debye relaxation,²⁴ the orientational relaxation of the hydrogen-bonded water network at ~20 GHz is modelled with a symmetrically broadened (as compared to a Debye relaxation) Cole–Cole relaxation,²⁵ and the small amplitude, fast relaxation mode at ~200 Hz attributed to a small angular motion of water²⁶ is described with a Debye type relaxation mode:

\[
\varepsilon(\nu) = \frac{S_{\text{NMA}}}{1 + \left(2\pi i\nu/T_{\text{NMA}}\right)^{1+\kappa}} + \frac{S_{\text{water}}}{1 + \left(2\pi i\nu/T_{\text{water}}\right)^{1-\kappa_\text{CC}}} + \frac{S_{\text{fast}}}{1 + \left(2\pi i\nu/T_{\text{fast}}\right)} + \frac{e_{\infty}}{\epsilon_0} + \frac{\kappa}{2\pi i\nu/\epsilon_0}
\]  

In eqn (1), $S_j$ and $T_j$ are the relaxation amplitudes and the relaxation times, respectively. $S_{\text{CC}}$ accounts for the broadening of the water relaxation mode. The limiting permittivity at high frequencies, $e_{\infty}$, subsumes all polarizations (e.g. due to molecular vibrations, or electronic polarization) that occur at frequencies higher (on time scales that are faster) than the frequencies covered in the present study. The last term of eqn (1) accounts for Ohmic loss contributions, which we approximate using a purely real frequency independent conductivity, that is the dc conductivity, $\kappa$. $\epsilon_0$ is the permittivity of free space.

In order to disentangle the closely overlapping relaxations due to NMA (~4 GHz) and water (~20 GHz), we reduce the number of parameters in fitting eqn (1) to the spectra. As water is the prevailing molecular species in all samples, we fix $S_{\text{water}}$ to what would be expected for an ideal solution (accounting for the reduced water concentration in the samples upon addition of salt, the KD for random distribution of the ions in the solution, and full hydration of Na⁺ and Mg²⁺; for details see ESI†).²⁷,²⁸ This constrained model describes the experimental spectra very well (solid lines in Fig. 1a and Fig. S1, ESI†) and the contributions of the NMA relaxation, the water relaxation, and the fast water relaxation to the dielectric loss are indicated as shaded areas in Fig. 1a.

Both, the parameters describing the fast water relaxation and the relaxation time of water vary with salt concentration as reported previously: $\tau_{\text{water}}$ decreases with increasing salt concentration.
concentration (Fig. S5a, ESI†) – a common observation for aqueous electrolytes,19–21,25 while the relaxation time of the fast water mode is largely unaffected by addition of salt (Fig. S5c, ESI†). The amplitude of the fast water mode somewhat increases with increasing c_{salt} as seen in Fig. S4c (ESI†). This increase in S_{max} is a common observation upon addition of salt and can be related to an increasing angular mobility of water in the hydration shells of ions.19–21 More relevant to the ion-specific effects discussed above are the parameters of the NMA relaxation (first term of eqn (1)). As can be seen from Fig. 1b and c, the contribution of NMA to the dielectric loss (the imaginary part of the first term of eqn (1)) strongly depends on the nature of the added salt: both the amplitude and the position vary with varying salt. The somewhat different center positions (i.e. values of t_{NMA}) of the peaks have been shown to correlate with the solution viscosity,19 evidencing that the rotational relaxation of NMA is consistent with diffusive rotation of NMA. Additionally, we observe a very pronounced variation of the relaxation amplitude (peak intensity) of the NMA relaxation (Fig. 1b and c) when changing the chemical identity of the salt. Comparing this relaxation strength for variation of an individual cation/anion with keeping the counterion (anion/cation) the same, the observed variation follows the Hofmeister series: for instance the NMA relaxation amplitude is lower for NaI compared to NaCl or KI (Fig. 1b). Hence, ions with increasing protein denaturation tendency increasingly reduce the relaxation strength. In other words: our results indicate that denaturing ions (e.g. KI) tend to destabilize proteins in contrast to Cl⁻, which gives a first indication of non-additivity of the individual ion-properties.

To quantify the reduction of the rotational mobility due to the different salts, we relate the relaxation strength of NMA (S_{NMA}) to the concentration of free NMA – the molar concentration of NMA that is not affected by the salt, c_{NMA,free}. Since for diffusive rotation as the underlying molecular mechanism of the observed relaxation, the relaxation amplitude is proportional to the molar concentration of the dipolar species and their squared effective electrical dipole moment (see eqn (S2), ESI†), we extract c_{NMA,free} by assuming the effective dipole moment of NMA to be constant.‡ In turn, (2 mol L⁻¹·c_{NMA,free}) is a measure for the effective concentration of NMA molecules that are rotationally hindered. The thus obtained values of (2 mol L⁻¹·c_{NMA,free}), are displayed in Fig. 2 for the investigated salts. In general, for all studied salts the number of affected NMA molecules is increasing with increasing salt concentration, with MgI₂ showing the largest effect. The trends follow the Hofmeister series and – in line with our previous study – variation of the cation has a more pronounced effect on c_{NMA,free} than substitution of the anion.

To explore whether the effects of the studied salts on c_{NMA,free} can be explained by the additive effect of the anion and of the cation, we compare the present values of (2 mol L⁻¹·c_{NMA,free}) to the value that would be expected from the cation and the anion individually. The effect of the ‘individual’ ions we take from our earlier studies on the K⁺ salt and the Cl⁻ salts, as c_{NMA,free} has remained virtually constant at c_{NMA,free} = 2 mol L⁻¹ upon addition of KCl.19,20 As can be seen from Fig. 2a–c the values of (2 mol L⁻¹·c_{NMA,free}) observed for NaBr, NaI, and MgI₂ agree within the experimental uncertainty well with the sum of the affected NMA concentration of the corresponding K⁺ and Cl⁻ salt: within the error bars the red symbols and the black solid lines in Fig. 2a–c coincide. Hence, our results suggest that the observed reduction of the rotational mobility of NMA is the average, additive effect of both the anion and the cation. This notion however does not hold for the salts containing the Gdm⁺ cation: in Fig. 2d and e we show the same comparison for GdmSCN/GdmI, where the observed reduction of the rotational mobility of NMA is lower than the sum of GdmCl and KSCN/KI, respectively. Hence, our results provide evidence for non-additivity for guanidinium salts.

To provide further evidence for the non-additivity for Gdm⁺ salts, we perform two additional series of experiments at c_{salt} = 1.5 mol L⁻¹. In one series we vary the molar ratio of the salt from neat NaCl to neat KI. In line with the additivity of the individual effects of Na⁺ and Cl⁻ described above, we find that (2 mol L⁻¹·c_{NMA,free}) varies linearly with the salt composition (Fig. 3a). Conversely, when varying the molar ratio of the salt from neat GdmCl to neat KI, the variation of (2 mol L⁻¹·c_{NMA,free}) cannot be described well by a linear relation (dashed blue line in Fig. 3b). Rather, our results indicate that the number of affected NMA molecules shows a minimum for the equimolar GdmCl/KI mixture: In the presence of 0.75 mol L⁻¹ GdmCl and 0.75 mol L⁻¹ KI the number of NMA molecules with reduced rotational mobility is lower than for solutions containing 1.5 mol L⁻¹ GdmCl or 1.5 mol L⁻¹ KI. This nonlinear behavior for GdmCl/KI mixtures thus further supports the notion that the effect of Gdm cannot be described by the
independent interaction of the Gdm\textsuperscript{+} cation and the I\textsuperscript{−} anion with NMA.

Thus our results suggest that the additivity of the ion-amide interaction depends on the identity of the ions. For the simple spherical cations K\textsuperscript{+}, Na\textsuperscript{+}, and Mg\textsuperscript{2+} and the corresponding spherical halide anions Cl\textsuperscript{−}, Br\textsuperscript{−}, and I\textsuperscript{−} our results suggest that their ability to reduce the rotational mobility of NMA can be explained from the sum of the individual interaction of the anion and the cation with NMA. Hence, the cation and the anion interact (on average) independently with the amide and – despite the long-range electrostatic interactions – there is no need to invoke any cooperativity. In contrast, our experiments provide evidence that such additivity does not hold for salts containing the guanidinium cation. For the guanidinium salts GdmI and GdmSCN we find less NMA molecules to be affected in their rotational mobility than what would be expected based on the individual effect of Gdm\textsuperscript{+} and the anions I\textsuperscript{−} or SCN\textsuperscript{−}. It may be significant that the minimum in Fig. 3b is located close to the equimolar composition, which points towards a competitive interaction for Gdm\textsuperscript{+} and I\textsuperscript{−}.

In general there are two plausible explanations for such competitive interaction: (i) the Gdm\textsuperscript{+} cation binds preferentially to the I\textsuperscript{−} or SCN\textsuperscript{−} anion and thus the cation–anion interaction is competing with the ion–NMA interaction. However, such a scenario appears unlikely, since the tendency of Gdm\textsuperscript{+} to associate with anions in solution (i.e. to form ion-pairs) is rather low and comparable to the ion-pair association of the corresponding sodium salts.\textsuperscript{29,30} Additionally, the dielectric spectra of the aqueous solutions of GdmI and GdmSCN (see ESI\textsuperscript{†} Fig. S2) do not exhibit any indications for contributions due to long-lived ion-pairs. As ion-pairs can in principle be readily detected with DRS via a separate ion-pair relaxation due to their intrinsically large electrical dipole moment,\textsuperscript{31} the absence of a contribution due to long-lived ion-pairs renders scenario (i) rather unlikely. Alternatively, (ii) the origins of the competitive behavior may be a result of competitive interaction with NMA: if Gdm\textsuperscript{+} and I\textsuperscript{−}/SCN\textsuperscript{−} reduce the rotational mobility of NMA via association to the same binding site at NMA, a similar dependency of the affected NMA molecules as shown in Fig. 3b would be expected, as at the equimolar composition the anions and the cations compete with a limited number of NMA binding sites. This notion might in fact be crucial for the high denaturation efficiency of the Gdm\textsuperscript{+} cation, as in general the disruption of proteins by anions – i.e. binding to anionic binding sites – more effectively denatures proteins, as compared to spherical cations.\textsuperscript{18}

Conclusions

In the present work we study for a range of salts the additivity of anionic and cationic interaction with the model amide NMA in aqueous solutions. We determine the reduction of the rotational mobility of NMA due to the presence of these salts via the rotational relaxation of NMA that contributes to the dielectric spectra of the samples. For the spherical, monovalent salts NaI, NaBr, and the bivalent MgI\textsubscript{2}, we find that the reduction of rotationally mobile NMA molecules is consistent with the additive effect of the individual anions and cations. Salts containing the strongly denaturing guanidinium cation deviate from additivity and we find that GdmI (and similarly GdmSCN) interacts to a lesser extent with NMA than the sum of the interaction of its individual ions (Gdm\textsuperscript{+} and I\textsuperscript{−}). This non-additive behaviour is likely related to competitive interaction for the Gdm\textsuperscript{+} cation and the I\textsuperscript{−} or SCN\textsuperscript{−} anion. Overall, our results indicate that separating the Hofmeister series into separate series for anions and cation is largely justified, with the notable exception of guanidinium salts.

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Notes and references

\textsuperscript{†} Note that the used effective dipole moment values may contain medium effects (e.g. molecular polarizability) and dipolar correlations. Given the similar static permittivities and ranges of ionic strengths for each studied salt (except Mg\textsuperscript{2+} salts), screening effects and thus dipolar contributions are expected to be similar as only the nature of the cation is changed.


