The behavior of ions in water is controlled by their water affinity

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Abstract

The strong, long-range electrostatic forces described by Coulomb’s law disappear for ions in water, and the behavior of these ions is instead controlled by their water affinity – a weak, short-range force which arises from their charge density. This was established experimentally in the mid-1980s by size-exclusion chromatography on carefully calibrated Sephadex® G-10 (which measures the effective volume and thus the water affinity of an ion) and by neutron diffraction with isotopic substitution (which measures the density and orientation of water molecules near the diffracting ion and thus its water affinity). These conclusions have been confirmed more recently by molecular dynamics simulations, which explicitly model each individual water molecule. This surprising change in force regime occurs because the oppositely charged ions in aqueous salt solutions exist functionally as ion pairs (separated by 0, 1 or 2 water molecules) as has now been shown by dielectric relaxation spectroscopy; this cancels out the strong long-range electrostatic forces and allows the weak, short-range water affinity effects to come to the fore. This microscopic structure of aqueous salt solutions is not captured by models utilizing a macroscopic dielectric constant. Additionally, the Law of Matching Water Affinity, first described in 1997 and 2004, establishes that contact ion pair formation is controlled by water affinity and is a major determinant of the solubility of charged species since only a net neutral species can change phases.

Introduction

While purifying Escherichia coli dihydroorotase (an enzyme in the biosynthetic pathway of the pyrimidines) to study its catalytic mechanism of action, we discovered that the enzyme exhibited instability of two kinds. The first was instability to oxygen which covalently modified the enzyme and lowered its catalytic activity by about a third; this could be avoided by carefully removing adventitious copper and iron from all solutions and minimizing the exposure of the enzyme to oxygen (Washabaugh and Collins, 1984; Brown and Collins, 1991). The second kind of instability was the tendency of the enzyme to completely inactivate by unfolding (denaturing) at concentrations below about 100 µg ml⁻¹ (Figs 1 and 2) (Washabaugh and Collins, 1986a, 1986b). The actual mechanism of denaturation was the dissociation of a dimer to a monomer which subsequently unfolded irreversibly, increasing the solvent exposed surface area of the polypeptide chain (Washabaugh and Collins, 1986a, 1986b). We tested the ability of many small molecules, mostly salts, to stabilize low concentrations of the enzyme in its native, folded, catalytically active conformation (Fig. 3). It was the anionic part of these salts which had the largest effect, probably because the anions are larger than the cations. When arranged from most stabilizing on the left to most destabilizing on the right with Cl⁻ near the center (Fig. 4), these anions formed a Hofmeister series, so-called because Franz Hofmeister formed a similar series in 1888 based upon their precipitating effects on hen egg white protein (Hofmeister, 1888; Collins and Washabaugh, 1985; Hofmeister, 2004; Collins, 2012). [This sequence is actually the order in which the ions elute from a Sephadex® G-10 column (Washabaugh and Collins, 1986a, 1986b).] The series that we initially produced had neutral, negatively charged, and positively charged species – so it was not obvious what property of these small molecules made them protein-stabilizing or destabilizing. Our studies determining the molecular basis of Hofmeister effects (Collins and Washabaugh, 1985; Washabaugh and Collins, 1986a, 1986b; Collins, 1995, 1997, 2004, 2006, 2012; Kiriukhin and Collins, 2002), the work of Neilson and Enderby using neutron diffraction with isotopic substitution (NDIS) to characterize ion hydration (Skipper and Neilson, 1989; Enderby, 1995), and advances in molecular dynamics simulations of aqueous salt solutions (Vrbka et al., 2006; Fennell et al., 2009; Stirnemann et al., 2013; Shi and Beck, 2017), all of which we review in this paper, clarify the forces in aqueous solution that give rise to biological structure and allow us for the first time to manipulate biological macromolecules in a systematic way and to control the stability and solubility of proteins.

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The molecular basis of the Hofmeister effect

Strong and weak hydration

For simplicity we decided to examine the anions by size-exclusion chromatography using Sephadex® G-10, which is the most highly epichlorohydrin-cross-linked dextran in beaded form commercially available. Sephadex® G-10 may be thought of as neutral beads with a nonpolar surface (Holmberg, 1983) (Fig. 5) containing uniform size pores which can separate small molecules below a molecular weight of about 700 by a 'size-exclusion' mechanism: small molecules penetrate the beads and have a longer pathlength through a packed column (slow elution), whereas larger molecules are excluded from the interior of the beads and have a shorter pathlength through a packed column (fast elution) (Fig. 6). This separation mechanism as described involves no direct interaction of the solute with the neutral Sephadex® G-10 beads. Solutes which have an apparent molecular weight larger than their anhydrous molecular weight are binding some water molecules with a lifetime of at least several picoseconds and effectively carry these water molecules with them through the column (Kiriukhin and Collins, 2002); solutes which have an apparent molecular weight smaller than their anhydrous molecular weight are adsorbed to the neutral surface of the beads. To ensure that no adventitious charges on the Sephadex® G-10 beads could affect
A “Hofmeister Series”
Hofmeister (1888)

\[
\text{SO}_4^{2-} \sim \text{HPO}_4^{2-} \succ F^- \succ I^- > Cl^- \succ Br^- > I^- \sim \text{ClO}_4^- \succ SCN^-
\]

(stabilizing)

(destabilizing)

Fig. 5. A typical structure isolated from Sephadex G-25 Holmberg (1983), which is less highly cross-linked than Sephadex G-10. The epichlorohydrid-cross-linked dextrans contain no aromatic residues. This research was originally published in the Proceedings of the National Academy of Sciences. Collins (1995) © 1995 National Academy of Sciences.

our results, all eluting solutions contain 0.1 M NaCl; leaving the NaCl out of the eluting solutions causes no detectable difference in the chromatographic behavior of ions on Sephadex G-10. The included volume of the column (the longest pathlength attainable by gel sieving) was determined by chromatographing tritiated water (THO \[^{3}\text{HOH}\]), or with slightly better accuracy, \[^{18}\text{O}\]-water (Marsden, 1971; Kiriukhin and Collins, 2002). Ions eluting later than isotopically labeled water must be adsorbed to the neutral poly-ether surface of the Sephadex G-10. The excluded volume of the column (the shortest pathlength through the column) was determined by chromatographing macromolecular dextran of average molecular weight of 40 000, or, with slightly better accuracy, by chromatographing the strongly hydrated negatively charged polyglutamate of average molecular weight of 40 000, or, with slightly better accuracy, by chromatographing the strongly hydrated negatively charged polyglutamate of molecular weight of 13 700 at pH 7 (Washabaugh and Collins, 1986a; 1986b; Kiriukhin and Collins, 2002). The specific calibration of the column allows correlating an elution position with a molecular weight is achieved with \[^{18}\text{O}\]-water and polymers of glycine (the simplest amino acid) containing from two to six subunits (Kiriukhin and Collins, 2002). The apparent molecular weight of a salt on Sephadex G-10 is assigned to the sum of the weight of the anion plus the weight of the cation plus the weight of any additional water molecules bound tightly enough to either ion to have a lifetime of at least several picoseconds at room temperature (30 °C) (Kiriukhin and Collins, 2002). Apportioning the bound water to a specific ion is achieved by judicious selection of the counterion, yielding an ‘appropriate dynamic hydration number’ (Kiriukhin and Collins 2002). For example, we know that Cl\(^{-}\) is weakly hydrated (relative to the strength of water–water interactions) by its slightly negative Jones–Dole viscosity B coefficient (see below), and thus its apparent dynamic hydration number is 0. And since it does not interact with Sephadex G-10 (see below), it passes through the column by gel sieving and thus is the ideal counterion for determining the dynamic hydration number of cations. Sephadex G-10 itself contains no charged groups; but to make sure that ions studied using Sephadex G-10 experience no ion-exchange effects, the eluent always contains 0.1 M NaCl.

The experimental setup is extremely simple, and the results are both surprising and very informative. A 4-l side arm flask containing 0.1 M NaCl which has been de-gassed while stirring with a water aspirator and filtered through a 0.45 µm filter is set atop a cabinet about 7 ft tall and run by gravity at 1.2 ml min\(^{-1}\) through a 1 m × 1.6 cm diameter column of Sephadex G-10 jacketed at 30 °C and then into a drop counting fraction collector sitting on the floor. Each experiment is begun by layering 0.6 ml of a 0.1 M salt solution onto the top of the Sephadex G-10 bed, and eluent fractions are collected in glass tubes. At the end of the experiment, which takes between 2.5 and 12 h, the 0.6 ml fractions are assayed manually. For ease of detection we used radioactive ions whenever possible (Washabaugh and Collins, 1986a, 1986b).

When NaF, NaCl, NaBr and NaI (the anhydrous ion radii of the halide anions increase as one goes down the periodic table – see Table 1) (Sharpe, 1992) are chromatographed separately or together on Sephadex G-10 the data obtained are shown in Fig. 7 (Washabaugh and Collins, 1986a, 1986b). The first thing that one notices is that the ions elute from the smallest (F\(^{-}\)) firstly to the largest (I\(^{-}\)) last, the opposite order of solute elution normally seen with gel sieving chromatography; therefore something other than the anhydrous ionic radius is controlling the behavior of the anions. The second thing that one notices is that Br\(^{-}\) and I\(^{-}\) elute after the gel sieving region (bracketed by THO \[^{3}\text{HOH}\] and dextran) indicating that they are being pushed onto the neutral surface of the Sephadex G-10 beads by the strength of water–water actions. This is confirmed by examining the concentration (from 0.025 to 0.6 M, see Fig. 8) and temperature (from 4 to 50 °C, see Fig. 9) chromatographic dependence of all four anions: F\(^{-}\) and Cl\(^{-}\) chromatographic behavior is temperature and concentration independent (consistent with gel sieving) whereas Br\(^{-}\) and I\(^{-}\) chromatographic behavior is temperature and concentration dependent (indicating interaction with the neutral surface of the Sephadex G-10 beads). In these figures the elution position is represented as the \(K_d = ([V_i - V_0]/(V_i - V_0))\), which is a linear fraction that varies from 0 for the elution position of the macromolecular excluded volume (\(V_0\)) (the earliest position for size exclusion) to 1 for the elution position of the \[^{18}\text{O}\]-water included volume (\(V_i\)) (the latest position for size exclusion); a \(K_d > 1\) indicates slow elution caused by interaction with the surface of the column. Since the surface of the Sephadex G-10 beads is mostly polyethers and is thus not very polarizable, we do not think that there are significant attractive forces between the larger, more polarizable anions Br\(^{-}\) and I\(^{-}\) and the surface.
of the neutral beads (more on this below); rather the weakly hydrated \( \text{Br}^- \) and the even more weakly hydrated \( \text{I}^- \) are pushed onto the surface of the column by strong water–water interactions. This interpretation is supported by comparison with several other techniques as discussed below. As will also be explained below, carefully calibrating the column with polymers of glycine allows us to conclude that \( \text{F}^- \) diffuses through the column with 5 water molecules attached whereas \( \text{Cl}^- \) appears to be smaller than \( \text{F}^- \) because \( \text{Cl}^- \) diffuses through the column with no water molecules attached. The \( \text{Cl}^- \) ion has remarkable properties: it is slightly weakly hydrated as shown, for example, by its negative Jones–Dole viscosity \( B \) coefficient (see below); yet it does not interact with the surface of the Sephadex® G-10 beads but it does orient the adjacent water molecules (as shown by NDIS, Fig. 10) (Powell et al., 1993; Enderby, 1995) via a nearly linear hydrogen bond (rather than via a dipolar interaction), showing that the strong, long-range electrostatic forces cancel out, allowing chemistry to come to the fore. [X-ray absorption spectroscopy of aqueous \( \text{Cl}^- \) detects two solvation shells of 7 water molecules each, the first at 3.15 Å and the second at 4.14 Å (Antalek et al., 2016)].

Another way to present the Sephadex® G-10 chromatographic behavior of \( \text{F}^- \), \( \text{Cl}^- \), \( \text{Br}^- \) and \( \text{I}^- \) as their Na\(^+\) salts is as the elution position of the salt on the vertical axis (measured as its \( K_d \)) versus the log\(_{10}\) of its molecular weight on the horizontal axis (Fig. 11); while the neutral calibration standards are very well behaved and fall on a straight line, the experimental ions show non-ideality with a vengeance. They form a line essentially perpendicular to that of the calibration standards, demonstrating that something other than anhydrous molecular weight is controlling their behavior. In this plot, a point that falls below the calibration line indicates a functional molecular weight larger than the anhydrous molecular weight caused by bound water whereas a point that falls above the line indicates a functional molecular weight smaller than the anhydrous molecular weight, which is caused by adsorbing to the surface of the Sephadex® G-10 beads. [Guadinium and thiocyanate, both of which adsorb to the surface of the Sephadex® G-10 beads in Figs 8 and 11, have been shown by NDIS to be very weakly hydrated (Mason et al., 2003).] We see that \( \text{F}^- \), \( \text{HPO}_4^{2-} \) and \( \text{SO}_4^{2-} \), the ions which bind water tightly and thus look larger than their anhydrous molecular weight, are the anions which stabilize proteins. This suggests that the stabilizing anions are competing for interfacial water at the surface of the protein, making the solution a poorer solvent and encouraging the protein to minimize its solvent-exposed surface area by acquiring a more compact form. This, in fact, is correct and

Table 1. Ionic radii (Å) for six-fold coordination (Sharpe, 1992)

| Ion  | Radius (Å) |
|------|------------|
| \( \text{Li}^+ \) | 0.74 |
| \( \text{Na}^+ \) | 1.02 |
| \( \text{F}^- \) | 1.33 |
| \( \text{K}^+ \) | 1.38 |
| \( \text{Cl}^- \) | 1.81 |
| \( \text{Br}^- \) | 1.96 |
| \( \text{Cs}^+ \) | 1.70 |
| \( \text{I}^- \) | 2.20 |

Fig. 6. Size exclusion chromatography. Large solutes which are unable to enter the beads have a shorter path through the column and elute early; small solutes which are able to enter the beads have a longer path through the column and elute late.

Table 1. Ionic radii (Å) for six-fold coordination (Sharpe, 1992)
was Hofmeister’s conclusion in 1888 although he had no way to prove it (Hofmeister, 1888; Hofmeister, 2004).

Chromatographing the group IA cations as the chloride salts shown in Fig. 12 gives a pattern analogous to that of the anions, with the small Li\(^+\) and Na\(^+\) ions having positive dynamic hydration numbers (see below) and not interacting with the column (although the position of Na\(^+\) is slightly offset in Fig. 12 because the 40,000 MW macromolecular dextran used to calibrate the column in these experiments interacts weakly with the surface of Sephadex\(^\text{\textregistered}\) G-10) whereas the larger K\(^+,\) Rb\(^+\) and Cs\(^+\) are pushed onto the neutral surface of the Sephadex\(^\text{\textregistered}\) G-10 beads by strong water–water interactions. The simplest way to think about this is shown in Fig. 13. For simplicity we shall consider an ion to be a sphere with a point charge at its center; as one goes down the periodic table, the water molecules at the surface of the ion become further from the point charge at the center such that the ion–water interactions become weaker than the bulk water–water interactions for K\(^+\) and larger cations and for Cl\(^-\) and larger anions. [In this picture, water is modeled as an embedded spheres zwitterion with a negative portion of radius 1.78 Å and a positive portion of 1.06 Å (Collins, 1997).] When these large monovalent ions adsorb to a surface, they release weakly bound water to become more strongly interacting bulk water. Our ion experiments on Sephadex\(^\text{\textregistered}\) G-10 and the solution experiments of Neilson and Enderby utilizing NDIS clearly show the transition from strong hydration for Li\(^+\) and Na\(^+\) to weak hydration for K\(^+\) (Fig. 14) (Enderby, 1995); Neilson and Enderby were also able to show that D\(_2\)O was bound to Li\(^+\) and Na\(^+\) preferentially through the oxygen atom whereas K\(^+\) did not orient the adjacent D\(_2\)O (Enderby, 1995) (data not shown in this paper). Their NDIS data indicate that the effects of these ions in water is short range, not extending beyond the adjacent single water layer, and that the dynamic hydration numbers described below measure only tightly bound, oriented water molecules. These data can be explained only if water affinity is the controlling force on ions in water, and, they are consistent with many other physical techniques. For example, proton nuclear magnetic resonance (NMR) longitudinal \(T_1\) relaxation rates show that this weakly bound water adjacent to large monovalent ions (K\(^+\) and larger) is actually tumbling faster than bulk water (see below) (Endom \textit{et al.}, 1967). This picture of short-range ion effects on water is supported by molecular dynamics simulations (Stirnemann \textit{et al.}, 2013).

\section*{Salting in and salting out effects}

Sephadex\(^\text{\textregistered}\) G-10 size exclusion chromatography also proves to be a powerful tool for examining salting in and salting out effects (Washabaugh and Collins, 1986a, 1986b). These experiments involve the use of \(10^{-4}\) to \(10^{-5}\) M of a radioactive ion [SO\(_4\)\(^{2-}\) (strongly hydrated), Cl\(^-\) (slightly weakly hydrated but almost neutral) or SCN\(^-\) (very weakly hydrated)] in the presence of 50–600 mM cold ion of the same or different hydration type.

When sulfate is the test (radioactive) solute (Fig. 15), adding large amounts of cold disodium sulfate decreases the hydrodynamic radius of the radioactive sulfate, making it smaller and causing it to elute later; when large amounts of cold sodium chloride are added there is no effect on the hydrodynamic radius of radioactive sulfate; when large amounts of cold sodium thiocyanate are added, the hydrodynamic radius of the radioactive sulfate becomes larger, causing it to elute earlier. Typically, weakly hydrated solutes act both indirectly in a water-mediated manner, and directly, by adsorbing to nonpolar surfaces; if the thiocyanate were partially clogging the pores of the Sephadex\(^\text{\textregistered}\) G-10, it would also cause the radioactive sulfate to elute earlier.

When chloride is the test (radioactive) solute (Fig. 16), adding large amounts of cold disodium sulfate salts out the radioactive chloride onto the surface of the Sephadex\(^\text{\textregistered}\) G-10 as shown by its elution position with a \(K_d\) well above 1. Adding large amounts of cold sodium chloride has very little effect on the elution position of the radioactive chloride. Adding large amounts of cold sodium thiocyanate increases the hydrodynamic radius of radioactive chloride, causing it to elute earlier; again, if the thiocyanate were partially clogging the pores of the Sephadex\(^\text{\textregistered}\) G-10, it would also cause the radioactive chloride to elute earlier.
When thiocyanate is the test (radioactive) solute (Fig. 17), adding large amounts of cold disodium sulfate salts out the radioactive thiocyanate onto the surface of Sephadex® G-10 even more strongly, increasing the $K_d$ from $\approx 7$ to $\approx 14$. Adding large amounts of cold sodium chloride has very little effect on the elution position of the radioactive thiocyanate. Adding large amounts of cold sodium thiocyanate increases the hydrodynamic radius of radioactive thiocyanate, causing it to elute earlier; it also competes for binding to the surface of Sephadex® G-10, displacing the adsorbed radioactive thiocyanate and causing it to elute earlier; it may also clog the pores of Sephadex® G-10, causing the radioactive thiocyanate to elute earlier. In all circumstances the radioactive thiocyanate shows absorption to the surface of Sephadex® G-10.
The disappearance of the strong, long-range electrostatic forces described by Coulomb’s law \( F_e = \frac{1}{4\pi\varepsilon_0}\frac{|q_1 \cdot q_2|}{r^2} \) appears to be the formation of close +/− ion pairs separated by 0, 1 or 2 water molecules as determined by dielectric relaxation spectroscopy (Buchner and Hefter, 2009) (Fig. 18). This cancels out the strong electrostatic forces and allows water affinity to come to the fore as the controlling force.

A simple general model for the hydration of most ions (and other solutes) in water

We can produce a simple general model for the hydration of most ions (and other solutes) in water. First, we know that the magnitude of the surface tension increment at an air/0.1 M salt solution interface indicates a separation of the ions from the interface of two water molecules (Randles, 1957). Second, gas phase studies of K⁺ solvation indicate that water is unique in that the second solvent layer makes a substantial contribution to the free energy of interaction of the solvent with test solutes (Sunner and Kebarle, 1984). In combination with several other kinds of data that we have summarized elsewhere (Collins and Washabaugh, 1985; Collins, 2004), we conclude that almost all of the favorable energy of interaction of water with ions occurs in the first two one-water molecule-thick layers. We shall postulate a two one-water molecule-thick layer hydration model for test solutes (test solutes are protein molecules in the Hofmeister effect) (Fig. 19). The inner layer is designated the solvation layer; the test solute controls the behavior of the solvation layer, which is not the same as saying that the test solute binds the solvation layer tightly. The second water layer is designated the transition layer. Beyond that is the bulk surface. Although estimates vary, we shall assume that the solvation layer constitutes about two-thirds of the solvent energy of interaction with the solute and the transition layer about one-third. In this model, the small molecule Hofmeister effect on a protein test solute is mediated by two intervening water molecules. [We have concluded from the temperature dependence of relative viscosity that [solute·H₂O·H₂O·solute] is a preferred configuration in aqueous solution, especially as the temperature is lowered toward 0 °C (Collins and Washabaugh, 1985)]. The test solute transition layer is the solvation layer of the Hofmeister ions, and its availability to the test solute is increased by weakly hydrated Hofmeister ions (making the solution a better solvent) and decreased by strongly hydrated Hofmeister ions (making the solution a poorer solvent), the latter causing the protein to minimize its solvent exposed surface area, thus stabilizing the protein and lowering its solubility. Weakly hydrated ions also adsorb directly onto the nonpolar portions of proteins.
relative to the strength of water ions with a negative viscosity change corresponds to the strength of water.

The line our best estimate of the region of normal behavior for solutes on Sephadex® G-10. Symbols are defined in the legend to Fig. 8. This research was originally published in the Journal of Biological Chemistry (Washabaugh and Collins, 1986b). © 1986 the American Society for Biochemistry and Molecular Biology.

**Jones–Dole viscosity \( B \) coefficient**

There is another method to study ions in water that is so simple and informative and complements the Sephadex® G-10 experiments so well that we shall introduce it now. The Jones–Dole viscosity \( B \)-coefficient, introduced in 1929, is a single ion-specific parameter that is defined by the equation

\[
\eta / \eta_0 = 1 + Ac^{1/2} + Bc
\]

which is valid at concentrations (c) up to about 0.1 M for binary strong electrolytes (Jones and Dole, 1929). \( \eta \) is the viscosity of an aqueous salt solution; \( \eta_0 \) is the viscosity of water at the same temperature; \( A \) is an electrostatic term that can be neglected at moderate concentrations; and \( B \) is a measure of the strength of ion–water interactions normalized to the strength of water–water interactions as verified by Sephadex® G-10 chromatography and also by NDIS. Viscosity \( B \) coefficients for relevant ions are given in Table 2 (Robinson et al., 1981; Krestov, 1991; Collins, 1997). Ions with a positive viscosity \( B \) coefficient are strongly hydrated relative to the strength of water–water interactions; ions with a negative viscosity \( B \) coefficient are weakly hydrated relative to the strength of water–water interactions; and the sign change corresponds to the strength of water–water interactions.

**Apparent dynamic hydration number**

This simple technique of aqueous Sephadex® G-10 size exclusion chromatography, which uses inexpensive equipment, effectively allowed us to measure the number of water molecules that move with the ion as long as the water–ion lifetime was at least a few picoseconds (Table 4) (Kiriukhin and Collins, 2002). We were even able to determine the apparent dynamic hydration number of the proton and of the hydroxide ion, and may have been the first (in 2002) to establish the ‘Zundel proton’ (the di-hydrate) as the correct form of the proton at 0.1 M (Kiriukhin and Collins, 2002; Reed, 2013). The four calibration points available to us establish that the ‘apparent dynamic hydration number’ is also the true ‘dynamic hydration number.’ The most informative points of comparison between gel sieving and informative and complements the Sephadex® G-10 experiments so well that we shall introduce it now. The Jones–Dole viscosity \( B \)-coefficient, introduced in 1929, is a single ion-specific parameter that is defined by the equation

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chromatography and other techniques are where natural discontinuities occur: the change from weak to strong hydration between K⁺ and Na⁺ as determined by neutron and X-ray diffraction with isotopic substitution (Enderby, 1995) and by Jones–Dole viscosity B-coefficients (Collins, 1997); the change from weak to strong hydration between Cl⁻ and F⁻ as determined by Jones–Dole viscosity B-coefficients (Collins, 1997); the change from weak to strong second-shell hydration between Mg²⁺ [as shown by solution X-ray diffraction (Skipper et al., 1989)] and Be²⁺ [as shown by ab initio molecular orbital calculations (Bock and Glusker, 1993)] and solution neutron diffraction studies (Mason et al., 2008) or Cr³⁺ [as shown by solution neutron (Broadbent et al., 1992) and X-ray diffraction (Munoz-Paez et al., 1995)] and the

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**Fig. 13.** How to think about ions in water Collins (1997). Division of group IA cations and the halide anions into small, strongly hydrated ions and large, weakly hydrated ions. The virtual water molecule on the right is drawn as a zwitterion of radius 1.78 Å for the anionic portion and 1.06 Å for the cationic portion.

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**Fig. 14.** The radial distribution functions g₁(r) for Li⁺, Na⁺, water and K⁺ in liquid water (Skipper and Neilson, 1989; Enderby, 1995). These curves measure the density of the solution as a function of the distance from the isotopically substituted ion, and effectively measure the distance from the monovalent cation to the nearest solvent oxygen. The curve labeled 'H₂O' measures the oxygen–oxygen distance in liquid water. Both neutron and X-ray diffractions were used to generate these data. The radial distribution for Li⁺ is drawn assuming a coordination number of six; subsequent experiments suggest a number closer to four, with no major effect on the results presented here. This research was originally published by Skipper and Neilson (1989). © 1989 Institute of Physics Publishing Ltd.

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**Fig. 15.** Uniform experiment with sulfate as test solute on Sephadex® G-10 (Washabaugh and Collins, 1986a, 1986b). 0.5 ml samples containing 9 × 10⁻⁵ M Na₂[³⁵S]SO₄ in 0–0.60 M uniform solute plus 0.10 M NaCl, THO [³HOH], and 0.5% dextran were chromatographed on a Sephadex® G-10 column (1.5 × 34.5 cm) at 30 °C and a flow rate of 1.5 ml min⁻¹. The eluent was 0.060 M uniform solute as indicated on the horizontal axis plus 0.10 M NaCl. Thus the concentration of NaCl as the uniform solute varied from 0.1 to 0.7 M. 0.65 ml fractions were collected. The uniform solute was Na₂SO₄, NaCl or NaSCN. With NaSCN as the uniform solute, the Kᵯ of Na₂[³⁵S]SO₄ was determined twice at each of the five NaSCN concentrations. The test solute elution profile was determined as described earlier. The tritiated water elution position (Kᵯ = 1) is off-scale at the top of the figure. This research was originally published in the Journal of Biological Chemistry Washabaugh and Collins (1986). © 1986 the American Society for Biochemistry and Molecular Biology.
change from an inner sphere coordination number of six for Mg$^{2+}$ (Skipper et al., 1989) to four for Be$^{2+}$ (Bock and Glusker, 1993; Mason et al., 2008). The apparent dynamic hydration numbers determined by gel sieving chromatography on Sephadex® G-10 are in excellent agreement with these calibration points from other techniques.

When the Sephadex® G-10 is properly calibrated with 18O-water and negatively charged polyglutamate of molecular weight 13 700 at pH 7 (as it was for the dynamic hydration number determinations), the behavior of ions on the column correlates well with the sign of their Jones–Dole viscosity $B$-coefficients; this implies that the polyglycine polymers used to calibrate the column are ideal solutes – i.e. that they bind water as strongly as water binds itself. This means that the hydration/dehydration of the peptide backbone associated with the unfolding/folding of the globular protein plays a neutral role, not favoring either conformational state.

**The Law of Matching Water Affinity**

*Ions of opposite charge form contact ion pairs in solution when they have matching water affinities*

The Law of Matching Water Affinity states that ions of opposite charge tend to form contact ion pairs in solution when they have matching water affinities (Fig. 20) (Morris, 1969; Collins, 1997, 2004). Since water affinity is a strong function of ion size [small ions of high charge density bind water strongly whereas large monovalent ions of low charge density (K$^+$ and larger for cations; Cl$^-$ and larger for anions) bind water weakly], small ions tend to form contact ion pairs with each other and large ions tend to form contact ion pairs with each other, but large-small contact ion pairs tend not to form (Collins 2004). This
same pattern is manifested in the solubility of the alkali halides (see below) (Collins, 1997). The small ions of opposite charge form contact ion pairs because of electrostatic attraction; the large ions of opposite charge form contact ion pairs because this releases weakly hydrated water which becomes strongly interacting water in bulk solution.

In 1969, D.F.C. Morris generated ‘volcano plots’ for the alkali halides based on either the absolute free energy of hydration of the constituent ions or the absolute enthalpy of hydration of the constituent ions (Morris, 1969). In 1997, when we first concluded that these plots indicated small-small and large-large contact ion pair formation, they were discussed in terms of the absolute free energies of hydration of the constituent ions (Collins, 1997). In 2004, we concluded that the absolute enthalpies of hydration of the constituent ions gave a better fit to the data and that they therefore gave rise to the Law of Matching Water Affinity (Collins, 2004).

The heat of solution of the alkali halides plotted as a function of the difference in the water affinity of the constituent ions (a volcano plot) supports the Law of Matching Water Affinities

We shall interpret Fig. 20 to indicate that oppositely charged ions with equal water affinity tend to stay together in solution as contact ion pairs whereas oppositely charged ions with differing water affinities tend to separate. We shall attribute the release of heat to the formation of strong bonds and the uptake of heat to the breaking of strong bonds, and shall assume that the strongest interactions in the system will tend to dominate the behavior of the system. In aqueous salt solutions of small ions of high charge density which are strongly hydrated and large ions of low charge density which are weakly hydrated, the interactions in order of decreasing strength are as follows: small–small > small–water > water–water > large–water > large–large (Collins, 1997). Figure 20 shows the relationship between the standard heat of solution of a crystalline alkali halide at infinite dilution (on the vertical axis; this is an experimental quantity) and the difference between the absolute heats of hydration of the constituent gaseous anion and cation [on the horizontal axis; this is a calculated quantity (Fig. 21)]. (In this context, ‘absolute’ refers to the conceptual experiment of transferring an isolated ion from the gas phase to the solution.) Figure 20 illustrates that a necessary but not sufficient condition for the standard heat of solution of a crystalline alkali halide to be negative (exothermic) is that one of the ions be large and the other ion to be small, suggesting that small plus large neutral salts dissociate extensively upon dissolution, and that the small ion of this salt acquires stronger interactions with water in solution than it has with large ions in the crystal, thus tending to release heat (Fig. 22). In contrast, when crystalline small–small alkali halides dissolve in water, the strongly interacting small–small ion pairs will tend to stay together, and moderately strong water–water interactions are replaced by weaker water–ion pair interactions, thus tending to take up heat. When crystalline large–large alkali halides dissolve in water, relatively strong water–water interactions will keep the large ion pairs together, and moderately strong water–water interactions are replaced by weaker water–ion pair interactions, thus tending to take up heat. These patterns suggest that oppositely charged

| Cations     | B  | Anions     | B  |
|-------------|----|------------|----|
| Mg$^{2+}$  | 0.385 | PO$_4^{3-}$ | 0.590 |
| Ca$^{2+}$  | 0.285 | CH$_3$CO$_2^-$ | 0.250 |
| Ba$^{2+}$  | 0.22  | SO$_4^{2-}$ | 0.208 |
| Li$^+$     | 0.150 | F$^-$       | 0.10 |
| Na$^+$     | 0.086 | HCO$_3^-$   | 0.052 |
| K$^+$      | −0.007 | Cl$^-$      | −0.007 |
| NH$_4^+$   | −0.007 | Br$^-$      | −0.032 |
| Rb$^+$     | −0.030 | NO$_3^-$    | −0.046 |
| Cs$^+$     | −0.045 | ClO$_4^-$   | −0.061 |
| I$^-$      | −0.068 | SCN$^-$     | −0.103 |

Source: Phosphate, formate, and perchlorate from Krestov (1991); all others from Robinson et al. (1981).
ions with equal water affinities will tend to form contact ion pairs in solution, whereas those with differing water affinities will tend to separate (Collins, 1997). In simplest possible terms, (strongly hydrated) small ions pair with (weakly hydrated) large ions pair with (weakly hydrated) large ions (Fig. 23). Since all of the salts in the volcano plot are monovalent, the long-range electric fields generated by each salt must be very similar, and the dramatic differences in their behavior must be due to the differences in the strength of their short range (electro-chemical) interactions with water. Forming a contact ion pair requires a partial dehydration of both the positive and negative ion, which occurs most readily when both ions have the same water affinity. A simple model can be used to show that the relative affinity of a monovalent ion for water closely correlates with its relative affinity for monovalent ions of opposite charge (Collins, 1997). Therefore, when one ion is more strongly hydrated than its oppositely charged partner, dehydrating the more strongly hydrated ion costs more in energy than is gained by forming a contact ion pair with the more weakly hydrated ion, and thus these ions tend to stay apart. The issue of charge density-dependent microscopic hydration–dehydration is not included in the electrostatic calculation using the macroscopic dielectric constant, but energetically it actually dominates and controls the process of contact ion pair formation (Collins et al., 2007). Evidence for contact ion pairing in water comes from protein X-ray crystallography (Dauter et al., 1999; Vaney et al., 2001; Pokhrel et al., 2011; Benas et al., 2014; Fox et al., 2015) and dielectric spectroscopy (Buchner and Hefter, 2009) in addition to those techniques discussed in the context of the Law of Matching Water Affinity above and below. The Law of Matching Water Affinity is supported by molecular dynamics simulations of the alkali halides (Fennell et al., 2009; Shi and Beck, 2017) and by activity coefficients (Kunz, 2010) as well as by other techniques discussed below.

The solubility of the alkali halide salts correlates with the charge density and thus the water affinity of their constituent ions

Although lattice enthalpies play a role (Perkyns and Pettitt, 1994; Shriver et al., 1994), the solubility of simple salts appears to be controlled largely by the tendency of the constituent ions to form contact ion pairs in solution as shown by the pattern of their solubility: salts composed of ions with similar water affinities have lower solubility. For example, both Li+ and F− are strongly hydrated, and thus tend to form contact ion pairs (the first step in the process of coming out of solution), whereas Cs+ is weakly hydrated (Ramos et al., 2005), and will tend to stay away from F−. The solubility of LiF in water is only 0.1 M at 18 °C. In contrast, the solubility of CsF at 18 °C is 24.2 M or 48.4 M in ions, and since pure water is about 55.5 M, a saturated solution of CsF contains only about one water molecule per ion’ (Collins et al., 2007). It is worth noting that The Law of Matching Water Affinity provides a reasonable explanation for the solubility of the alkali halides (Collins, 1997) whereas continuum electrostatics (lattice enthalpy) models do not (Perkyns and Pettitt, 1994).

The interaction of cations with proteins

In 1996, Wolff et al. (1996) showed that alkali metal cations increase the rate of polymerization of pure tubulin driven by either taxol or dimethyl sulfoxide according to the sequence Na+ > K+ > Li+ > Cs+, with an optimum concentration for Na+ at ≈160 mM. Because both the α and β monomers of nearly all tubulins carry considerable excess anionic charge, particularly at the C termini, added cations allow polymerization by binding
to protein carboxylates and eliminating repulsive anionic interactions in the monomer. Figure 24 shows that the most rapid tubulin polymerization correlates linearly with the smallest difference in water affinity between the added cation and the protein carboxylate as measured by their Jones–Dole viscosity $B$-coefficients; that is to say, these data support the prediction of The Law of Matching Water Affinities that $\text{Na}^+$ is the best monovalent cation binder to the carboxylate (Collins 2006). This result is confirmed by molecular dynamics simulations (Vrbka et al., 2006). [Anions have also been shown to bind the active site $\text{Zn}^{2+}$ of human carbonic anhydrase II according to the Law of Matching Water Affinities (Fox et al., 2015).]

It was shown in 1956 (Bello et al., 1956), 1962 (Von Hippel and Wong, 1962) and 1964 (von Hippel and Wong, 1964) that $\text{MgCl}_2$ affects the helix–coil transition of the fibrous protein collagen. However Hofmeister found that $\text{MgCl}_2$ by his procedure, did not affect the solubility of globular egg white protein (Hofmeister, 1888; Hofmeister, 2004), and Hofmeister effects are typically studied on globular proteins. It was shown in 1984 (Arakawa and Timasheff, 1984) that $\text{Mg}^{2+}$ does bind to surface carboxylates of globular proteins. It was shown in 1965 (Robinson and Jencks, 1965), 1966 (Bello et al., 1966) and 1969 (von Hippel and Schleich, 1969) that $\text{Ca}^{2+}$ seems to act on globular proteins mostly by complexing with the amide moiety.

In 2014 (Benas et al., 2014), the adsorption of $\text{Rb}^+$, $\text{Cs}^+$, $\text{Mn}^{2+}$, $\text{Co}^{2+}$ and $\text{Yb}^{3+}$ onto the positively charged hen egg white lysozyme was studied by X-ray crystallography and electrospray ionization mass spectrometry; $\text{Rb}^+$ and $\text{Cs}^+$ bound preferentially to carbonyl groups whereas the multivalent $\text{Mn}^{2+}$, $\text{Co}^{2+}$ and $\text{Yb}^{3+}$ interacted with carboxylate groups. A summary of specific ion effects on proteins was published in 2017 (Okur et al., 2017).

**The roles of charge groups in protein structure and function**

**Supercharged green fluorescent protein**

Green fluorescent protein (GFP) is an intensely studied protein which is 238 amino acids long. A ‘superfolder’ variant of GFP...
which had been highly optimized for folding robustness and resistance to denaturants had a net charge of $-7$. Its surface residues were mutagenized to produce either $+36$ or $-30$ ‘supercharged’ variants that were only slightly less stable than the starting version (Fig. 25) (Lawrence et al., 2007). When expressed in E. coli, these supercharged variants folded normally (bringing the charges closer together) showing that the like-charged side chains were not experiencing strong repulsive forces between each other. This indicates that there must have been an oppositely charged counterion near each of the surface charges thus neutralizing any electrostatic interactions.

**Increased negative surface charge correlates with increased solubility on seven different proteins**

Kramer et al. (2012) in 2012 determined the solubility of seven different globular proteins as a function of the fraction of positive or negative charges on the protein accessible surface area; while
the positive protein surface charges (which are weakly hydrated) had no effect on protein solubility, the negative surface charges (which are strongly hydrated) produced a large increase in protein solubility (Fig. 26).

Normal fibrin polymerization (blood clotting) requires the binding of physiological \( \text{Cl}^- \)

Fibrinopeptides must polymerize into a fine net to trap platelets to the site of injury and initiate processes that stop the bleeding and promote wound repair and healing. For this reason blood clotting requires the binding of (weakly hydrated) \( \text{Cl}^- \) to protein (weakly hydrated) positive charges to increase the \( pK_a \) of a basic group on the peptide and prevent the growth of thicker, stiffer and straighter fibers which make a poor net. (Strongly hydrated) \( \text{F}^- \) is inert in this assay as predicted by the Law of Matching Water Affinity which predicts that it should not bind to peptide (weakly hydrated) positive charges; thus \( \text{F}^- \) allows the unphysiological thicker, stiffer and straighter fibers to form (Fig. 27) (Di Stasio et al., 1998). Figure 28 shows that strongly hydrated acetate, like \( \text{F}^- \), also does not bind to the weakly hydrated positive charges on proteins and thus produces a thick fiber abnormal clot, whereas the weakly hydrated \( \text{Cl}^- \), \( \text{Br}^- \) and \( \text{I}^- \) do bind to the weakly hydrated positive charges on proteins and thus produce the thin, normal fibrin net which leads to blood clotting. The \( K_d \) for \( \text{Cl}^- \) binding to protein positive charges is from 133 (Overman and Lohman, 1994) to 150 mM (Makhatadze et al., 1998); the physiological concentration of NaCl is 145 mM (Vindigni and Di Cera, 1996), indicating that about half of the positive charges on blood proteins are binding \( \text{Cl}^- \).

Protein crystallization consists of two parts: (1) binding the necessary ions to produce a neutral complex and (2) dehydrating the protein surface, including the carboxylates, to allow protein–protein contacts

The H1 collagenase has a pI of 4.1 and is a model for acidic proteins. At pH 7.2 (and 18 °C) it has about 26 negative charges (carboxylates) and 13 positive charges (lysines and arginines). Figure 29 (Riès-Kautt and Ducruix, 1997) shows that H1 collagenase is crystallized most effectively by the strongly hydrated di- and tri-valent anions phosphate, sulfate and citrate (as the ammonium salts), which at high concentrations are able to dehydrate the surface carboxylates. That is, dehydrating the surface carboxylates is the energetically difficult step for crystallization of acidic proteins at neutral pH.

Hen egg white lysozyme has a pI of 11.1 and is a model for basic proteins. At a pH of 7 it has about 18 positive charges (lysines and arginines) and nine negative charges (carboxylates). At pH 4.5 (and 18 °C), most of the carboxylates are protonated, so there are few carboxylates that need to be dehydrated for crystallization. The energetically difficult step in crystallizing basic proteins is to produce a net neutral complex by binding (weakly hydrated) anions to the proteins’ (weakly hydrated) cationic groups. Thus the very weakly hydrated thiocyanate is the most effective crystallizing anion (Fig. 30) (Riès-Kautt and Ducruix, 1997). [In 1998, thiocyanate was also shown to interact with the amide backbone of Arg42 in bovine pancreatic trypsin inhibitor by using NMR (Jolivalt et al., 1998).] The strongly hydrated anions acetate, phosphate and citrate bind only very weakly to the (weakly hydrated) positive charges on the protein and thus are poor crystallizing salts.

Fig. 24. The maximal rate of tubulin polymerization versus the difference in Jones-Dole viscosity \( B \) coefficients between added monovalent cations and the tubulin carboxylates (Collins, 2006). The Jones-Dole viscosity \( B \) coefficients for the carboxylate \((0.052), \text{Na}^+ (0.086), \text{K}^+ (0.007), \text{Cs}^+ (0.045) \) and \( \text{Li}^+ (0.150) \) are measures of the water affinity of these ions. The Law of Matching Water Affinities states that those ions with matching water affinity are those which will most readily form inner sphere ion pairs. The difference in Jones-Dole viscosity \( B \) coefficients of two ions is a measure of their mismatch in water affinity. Therefore \( \text{Na}^+ \) has the greatest tendency to bind to carboxylates, followed by \( \text{K}^+ \), followed by \( \text{Li}^+ \), followed by \( \text{Cs}^+ \). © 2005 Elsevier B.V. All rights reserved.

Fig. 25. Supercharged variants of GFP (Lawrence et al., 2007). Red indicates negatively charged carboxylate side chains. Blue indicates positively charged arginine and lysine side chains. From Lawrence et al. (2007) [https://pubs.acs.org/doi/10.1021/ja071641y]. Further permissions should be directed to the American Chemical Society.
A note on osmolytes

(Net neutral) osmolytes have very similar hydration properties to those of ions. Stabilizing osmolytes with a strongly hydrated portion such as trimethylamine N-oxide and betaine act primarily by competing for the water solvating the transiently exposed peptide backbone, driving the latter back into the interior of the protein (Bolen and Baskakov, 2001; Auton et al., 2011); presumably sodium malonate, the most effective salt for crystallizing proteins, does much the same (McPherson, 2001). (Weakly hydrated) urea, in contrast, denatures proteins by increasing the availability of water to solvate the peptide backbone and also by adsorbing to the surfaces of buried nonpolar amino acid side chains, pulling them into solution (Moeser and Horinek, 2014).

Ions in enzyme active sites

The forces stabilizing proteins have recently been summarized (Pace et al., 2014). Enzymes typically have loops which close

Fig. 26. Globular protein solubility increases with the number of (strongly hydrated) negatively charged surface carboxylates, but not with the number of (weakly hydrated) positively charged surface amino acid side chains (Kramer et al., 2012). ASA = accessible surface area. Protein solubility was measured by (NH₄)₂SO₄ precipitation (on the right) and polyethylene glycol precipitation (on the left). From Kramer et al. (2012). © 2012 Biophysical Society. Published by Elsevier Inc.

Fig. 27. Scanning electron micrographs of fibrin clots grown in the presence of NaCl or NaF (Di Stasio et al., 1998). (A) Abnormal thicker, stiffer, straighter fibrin clots grown in 150 mM NaF plus 50 mM NaCl (F⁻ is inert in this system; some Cl⁻ is required to keep the fibrin clots in solution). (B) Normal fibrin clots grown in 200 mM NaCl. From Di Stasio et al. (1998). © 1998 The Biophysical Society. Published by Elsevier Inc.

Fig. 28. Turbidity of fibrin clots developed in the presence of different salts, at I = 200 mM kept constant with NaF (Di Stasio et al., 1998). Turbidity is measured as A₅₅₀. The strongly hydrated acetate is inert in this system, whereas the weakly hydrated Cl⁻, Br⁻, and I⁻ are active and bind to the weakly hydrated positive charges on the protein, inhibiting aggregation and leading to the physiologically active thinner fibers. From Di Stasio et al. (1998). © 1998 The Biophysical Society. Published by Elsevier Inc.
Over the active site, producing a nearly anhydrous protein box (Zhai et al., 2014) in which electrostatic interactions can be strong. Charged substrate binding groups such as phosphate mono-esters can grip the protein strongly and pull it into its rigid, tightly packed transition state binding conformation (Richard, 2019) where catalysis is facilitated by strong local electric fields (Fried and Boxer, 2017) such as in low barrier hydrogen bonds (Graham et al., 1992) in which electrostatic interactions can be strong.

**Future prospects**

Until recently, work on the basic physics of aqueous solutions and on the physical biochemistry of biological macromolecules has been undertaken largely to facilitate our ability to study pre-existing structures such as enzymes which are the targets of small molecule drugs. We are now entering a new era in which we are creating new structures or structures with new activities (Howes, 2019). For example, one of the fastest growing classes of pharmaceuticals is monoclonal antibodies targeted to human regulatory proteins which control fundamental processes such as cell replication or immune function (Chiu and Gilliland, 2016; Lonberg and Korman, 2017). These drugs are given by injection and require very concentrated solutions so that a monthly injection can achieve clinical activity. A large financial incentive therefore exists for developing the ability to produce highly soluble antibodies, which can be done with the principles described here.

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