Simple models for charge and salt effects in protein crystallisation

P. B. Warren
Unilever Research and Development Port Sunlight,
Bebington, Wirral, CH63 3JW, United Kingdom
(Dated: June 13, 2002 — Final version, accepted J. Phys. Cond. Mat.)

A simple extension of existing models for protein crystallisation is described, in which salt ions and charge neutrality are explicitly incorporated. This provides a straightforward explanation for the shape of protein crystallisation boundaries, the associated scaling properties seen for lysozyme, and can also explain much of the salt dependence of the second virial coefficient. The analysis has wider implications for the use of pair potentials to understand protein crystallisation.

I. INTRODUCTION

Protein crystallisation is of great practical importance, both as a purification method and since high quality crystals are needed for X-ray diffraction work. Under typical conditions where crystals are obtained, protein-protein interactions appear to be characterised by hard core repulsions with short-ranged attractions. For instance, Rosenbaum et al. successfully collapsed the crystallisation boundaries for a number of proteins onto the adhesive hard sphere (AHS) model by matching the second virial coefficient $B_2$. Whilst the AHS model is frequently used to understand protein crystallisation, it is also possible that crystallisation is an energetic ordering transition driven by highly directional interactions. A model of hard spheres with sticky patches was introduced by Sear to capture this important possibility.

In this paper I examine the properties of a simple model for protein crystallisation which takes into account the effects of charge in a very elementary way. Such a model suggests that some generic aspects of protein crystallisation can be explained as a straightforward consequence of charge neutrality, at least for lysozyme which readily undergoes crystallisation. These various aspects are: (i) the overall shape of the crystallisation boundary, (ii) a scaling collapse of the crystallisation boundary noticed by Poon et al. when plotted as a function of $c_p/Q^2$, where $c_p$ is the NaCl concentration and $Q$ the protein charge; (iii) a similar scaling of the lysozyme solubility data in Guo et al.; and (iv) a similar scaling for $B_2$ noticed by Egelhaaf and Poon.

The appearance of $c_p/Q^2$ in all these is very reminiscent of the Donnan membrane equilibrium, where one finds a contribution to $B_2$ equal to $Q^2/4c_p$ provided one is in the high-salt limit $c_p \gg Qc_p$ where $c_p$ is the protein concentration. The effect is well known in polyelectrolyte theory. This result is usually derived by supposing ideal solution behaviour and imposing electrical neutrality on the two sides of a semi-permeable membrane. In the present case, I argue there is an analogous Donnan equilibrium between protein crystals and a dilute protein solution, with the crystal-solution interface playing the role of the membrane. The appearance of $c_p/Q^2$ as a scaling variable reflects the fact that the solution (not necessarily the crystal) is in the high-salt limit, as will be explained below. Donnan’s theory appears to have little to do with the more conventional McMillan-Mayer approach in which effective potentials between proteins are invoked to explain crystallisation and the effects of added salt, often in the context of DLVO potentials where Debye-Hückel theory is used to account for the charge interactions. However, a careful study of the relationship between the two approaches made by Hill in the 1950s indicates both approaches should converge on the same results in the high salt limit.

To apply these ideas to the protein crystallisation problem, one constructs a free energy which enforces the charge neutrality constraint. Small ions can be treated as ideal solution species, but one must at least incorporate non-ideality of the protein solution, to allow it to form the ordered phase which represents the protein crystals. The first thing to do is to set up this general theory.

II. GENERAL THEORY

I suppose the proteins under consideration have a molecular volume $V_p$, for example lysozyme is a charged globular protein of approximate size $V_p = (\pi/6) \times 4.5 \times 3.0 \times 3.0 \text{ nm}^3 = 21.2 \text{ nm}^3$, or a molar volume $N_AV_p = 12.8 \text{ M}^{-1}$ where $N_A$ is Avogadro’s number. If the protein concentration is $c_p$, the effective volume fraction is $\phi = V_p c_p$. In the absence of charge effects, a baseline model which incorporates a freezing transition to correspond to protein crystallisation will be fully specified by a dimensionless free energy density $f^{(0)} = V_p \beta F^{(0)}/V$, where $F^{(0)}$ is the free energy of the protein solution or crystal, $V$ is the system volume, and $\beta = 1/kT$. There may be different branches of $f^{(0)}$ to correspond to the fluid and ordered (crystal) phases. Below I shall consider two possibilities for this baseline model: Sear’s model of hard spheres with sticky patches, and the AHS model. The full free energy is obtained from the baseline model by adding in contributions from the coions and counterions (which I suppose to be univalent) and imposing charge neutrality. If the added salt concentration is $c_s$, there will be coions at a concentration $c_s$ and counterions at a concentration $Qc_p + c_s$. Denoting the dimensionless
free energy density for the full model by \( f \), I write
\[
\begin{align*}
 f(\phi, \phi_s) &= f^{(0)}(\phi) + \phi_s[\log \phi_s - 1] \\
 &+ (Q\phi + \phi_s)[\log(Q\phi + \phi_s) - 1]
\end{align*} 
\] (1)

where the last two terms are the ideal mixing terms from the coions and counterions respectively, written using \( \phi_s \equiv V_p c_s \) for notational simplicity. Unimportant constants and terms linear in \( \phi \) or \( \phi_s \) have been dropped. Non-ideality of the small ions is neglected in the present treatment, although this is certainly a refinement which should be considered for more quantitative work.

This free energy is a function of two density variables: \( \phi \) and \( \phi_s \). A phase equilibrium such as between protein crystals and protein solution corresponds to equality of osmotic pressure and chemical potentials for both components. As a consequence there will in general be different values of \( \phi_s \) in coexisting phases, a salt repartitioning effect which is not usually taken into account. To solve for phase equilibria, it is useful to transform \( f \) into a semi-grand potential \( h(\phi, \phi_s^{(R)}) \) where \( \phi_s^{(R)} \equiv V_p c_s^{(R)} \) is a dimensionless salt reservoir concentration [2]. To make this transformation note that the salt chemical potential is
\[
\beta \mu_s = \frac{\partial f}{\partial \phi_s} = \log \phi_s + \log(Q\phi + \phi_s).
\] (2)

Now \( \phi_s \rightarrow \phi_s^{(R)} \) in the limit \( \phi \rightarrow 0 \), therefore \( \beta \mu_s = 2 \log \phi_s^{(R)} \). This gives
\[
(\phi_s^{(R)})^2 = \phi_s(Q\phi + \phi_s)
\] (3)

which is the usual Donnán equilibrium result that the product of the coion and counterion concentrations takes a constant value in all phases including the reservoir. Solving this gives
\[
\phi_s = [(Q^2\phi^2 + 4(\phi_s^{(R)})^2)^{1/2} - Q\phi]/2.
\] (4)

The semi-grand potential \( h = f - \beta \mu_s \phi_s \) and the first two derivatives with respect to \( \phi \) at constant \( \phi_s^{(R)} \) are, after a few lines of calculus,
\[
\begin{align*}
 h &= f^{(0)} + Q\phi \log(Q\phi + \phi_s) - (Q\phi + 2\phi_s), \\
 \partial h/\partial \phi &= \partial f^{(0)}/\partial \phi + Q \log(Q\phi + \phi_s), \\
 \partial^2 h/\partial \phi^2 &= \partial^2 f^{(0)}/\partial \phi^2 + Q^2/(Q\phi + 2\phi_s).
\end{align*} 
\] (5)

The advantage of this transformation is that \( h \) is effectively a one-component free energy and can be treated accordingly. To use these results, one should remember to substitute for \( \phi_s \) from Eq. (1). For example, the osmotic pressure follows from \( V_p \Pi = \phi \partial h/\partial \phi \) - \( h \), ie
\[
\Pi = \Pi^{(0)} + kT(Qc_p + 2c_s).
\] (8)

What is obvious from these results is that there is a cross-over in behaviour at \( Qc_p \sim c_s \) or salt concentrations of the order \( Q\phi/V_p \). The difference in protein volume fraction between the solution and crystal is often \( \Delta \phi \sim 0.5 \). Putting \( Q \sim 10 \), and \( V_p \sim 10^4 \text{M}^{-1} \), this corresponds to a cross-over salt concentration \( \sim 0.5 \text{M} \). If the salt concentration is much less than this, there will be a large osmotic pressure difference between the crystal and the solution due to the counterions, having the effect of narrowing the coexistence gap. This is the basic reason why the coexistence boundary occurs at salt concentrations of this order of magnitude in this model. It is a much larger cross-over salt concentration than intuition might have suggested based on experiences for colloidal systems (eg \( Q \sim 10^3 \), \( V_p \sim 10^6 \text{nm}^3 \) gives \( c_s \sim 10^{-3} \text{M} \)), but then globular proteins are much smaller than colloids.

On the other hand, typical protein solutions on the crystallisation boundary have \( \phi \sim 0.05 \), corresponding to a cross-over salt concentration \( \sim 50 \text{mM} \). For salt concentrations much larger than this, the effects of salt and charge are subsumed into a scaling variable \( \phi_s^{(R)}/Q^2 \). Since \( \phi_s^{(R)} \) is more or less the salt concentration in the solution if the protein concentration is small, this is a natural explanation for the scaling properties described in the introduction. In the next sections, I place these arguments on a firm footing, starting with discussions of the second virial coefficient and the high-salt scaling of solubility, since these are not dependent on any particular baseline model.

III. SECOND VIRIAL COEFFICIENT

Any theory for the free energy of a protein solution contains a prediction for the second virial coefficient. For the present theory, \( B_2 \) can be obtained from the osmotic pressure result Eq. (8) (recalling that Eq. (4) should be used for \( \phi_s \)), or directly by inspection from Eq. (5). Either way, as advertised in the introduction, \( B_2 = B_{2}^{(0)} + Q^2/4c_s \) where \( B_{2}^{(0)} \) is the contribution from the baseline model. One can prove the same result holds in light scattering determination since charge density fluctuations are suppressed in the long wavelength limit. We can hope that the baseline model is insensitive to the net charge and salt concentration, so that \( B_2^{(0)} \) is independent of \( Q \) and \( c_s \), but is there any evidence for this? For lysozyme, Egelhaaf and Poon have collected experimental data on \( B_2 \) from the available literature [3]. This data is used to construct Fig. 2(a) which shows \( B_2 - Q^2/4c_s \) normalised by the value expected if lysozyme proteins behaved as hard spheres, namely \( B_{HS}^{RS} = 4V_p \approx 85 \text{nm}^3 \) (this is close to \( B_{HS}^{RS} = 82.3 \text{nm}^3 \) used in Ref. [2]). Inspecting Fig. 2(a), it appears there is indeed a data collapse to an approximate plateau for \( c_s \gtrsim 0.25 \text{M} \) although a downwards trend of 40% or so can be detected at high salt and there may also be a problem if the protein charge is too small (the ‘rogue’
The predicted slope from Eq. (11) is \( \phi_s \approx 2V_p \). If we take \( \phi_s \approx 0.5 \) as a reasonable estimate of the protein volume fraction in the crystal, then \( \phi_s/2V_p \approx 0.02 \) M which around twice the measured slope. However there are a number of omitted effects in the present theory which could account for the discrepancy. Nevertheless the observation that the logarithm of the protein solubility is inversely proportional to the inverse salt concentration is apparently quite common [22]. Also note that solubility \( \sim c^2 \) indicates a very strong dependence on the protein charge. This is a natural explanation for the dramatic decrease in solubility around the isoelectric point described in Ref. [1].

V. TWO SPECIFIC BASELINE MODELS

For lower salt concentrations, the protein solubility is not small and the above treatment needs refinement. I
now consider the minimal extension to the basic model that takes into account protein non-ideality by specifying a particular baseline model. In this way, the advantage of an analytic transformation to the semi-grand potential $\pi n$ in Eqs. (3)−(4) is retained. The baseline model should encompass both the fluid and crystal phases, and have a second virial coefficient consistent with the analysis above. One such suitable model has been devised by Sear.

A. Sear’s model

Sear’s model comprises hard spheres (HS) with ‘sticky patches’ [9]. The HS diameter is $\sigma$, chosen such that $\pi \sigma^3/6 = V_p$ (ie $\sigma \approx 3.4$ nm for lysozyme). There are $n_s$ sticky patches per sphere which associate in pairs and are characterised by a range $r_c > \sigma$, an angular width $\theta_c$, and a depth $\epsilon$. The free energy of the fluid ($F$) phase (I reproduce only the essential details of the model here) is

$$f_F^{(0)}(\phi) = f_{HS}(\phi) + \phi n_s [\log p + (1 - p)/2]. \quad (13)$$

In this $f_{HS}$ is the HS fluid free energy and $p$ is the proportion of non-bonded sites, solving $(1 - p)/p^2 = k\phi g_{HS}$ where

$$k = 6(r_c/\sigma - 1)(1 - \cos \theta_c) e^{\beta \epsilon} \quad (14)$$

is a dimensionless bond association constant. The association equilibrium includes an enhancement factor, $g_{HS}$, for the HS pair correlation function at contact. The second virial coefficient in this model is given by

$$B_2^{(0)} = B_2^{HS} - (n_s/2) k V_p. \quad (15)$$

It is apparent that the fluid fluid properties are completely determined by $k$ and $n_s$. The model predicts fluid-fluid phase separation for sufficiently large values of $k$, and Table I gives the critical point for $n_s = 4−8$. Note that the second virial coefficient at the critical point in this model provides an marked counterexample to the observation of Vliegenthart and Lekkerkerker that $B_2^{(C)}/B_2^{HS} \approx -1.5$ for a wide variety of other models [19]. Whether fluid-fluid phase separation is metastable in this model depends on the actual values of $r_c$, $\theta_c$ and $\beta \epsilon$, and examples of both are given by Sear.

Sear provides a cell model for the protein chemical potential in an ordered cubic phase, arguing that osmotic pressure is not important. My approach here is slightly different. I use the same cell model to specify the free energy and take into account the osmotic pressure. The results are not essentially different from Sear’s results, although the analysis does point out an important property of Sear’s model (Fig. 2(a) below). Following Sear, the free energy per protein in the crystal ($X$) is

$$f_X^{(0)}(\phi) = C - 3 \log (\phi \sigma^3) - \beta \epsilon w \left[ (\phi - \phi_{min}) / \delta \phi \right] \quad (16)$$

where $a/\sigma = (6\phi/\pi)^{-1/3}$ is the unit cell size relative to the HS diameter. The constant in this is $C = \log (V_p/\sigma^3)$, provided that $f_{HS} \sim \phi (\log \phi - 1)$ in the fluid phase as $\phi \to 0$. This free energy is appropriate for $\phi \geq \phi_{min}$ where $\phi_{min} = \pi (\sigma/r_c)^3/6$ is the volume fraction around which the bonds in the crystal become dissociated. As $\phi$ decreases past $\phi_{min}$ the last term in Eq. (13) vanishes rapidly. To implement this I have introduced an ad hoc cut-off function $w((\phi - \phi_{min})/\delta \phi)$ in the last term, where $\delta \phi$ sets the rate at which the cut-off operates. For the present calculations I take $w(x) = 1/(1 + e^{-x})$ and $\delta \phi = 0.01$. This cut-off function represents the way in which the short-range attraction potential falls off with distance in the model. The actual details may shift the phase boundaries but are not important for the broad picture.

Although several values of the parameters in the model were examined, I only report in detail here on calculations for $n_s = 6$, $r_c = 1.05 \sigma$ and $\theta_c = 0.45$ (so the range of the attraction is $r_c - \sigma \approx 2$ Å and the angular width about $26^\circ$). These values were chosen to give quite good agreement with the crystallisation boundaries for lysozyme in the present model. Interestingly, in a separate analysis Curtis et al. [2] also conclude $n_s = 6−8$ is appropriate, and Oki et al. [23] identify three ‘macrobonds’ between lysozyme molecules from crystallographic data, again corresponding to $n_s = 6$ contacts per protein.

Fig. 2(a) shows the fluid and fluid free energies for $n_s = 6, r_c = 1.05 \sigma$ and $\theta_c = 0.45$, for two values of $\beta \epsilon$. It is clear that there is a certain minimum value $\beta \epsilon \approx 4.8$ below which the crystal is metastable with respect the fluid. This is an important contrast with the AHS model (next subsection) in which the ordered phase is present at $\beta \epsilon \to 0$. Crystallisation in Sear’s model is essentially an energetic transition to an ordered phase dominated by the directional interactions (compare ‘energetic fluid’ concept introduced by Louis [26]) and not a continuation of the entropic HS freezing transition. The position of the sticky patches controls the crystal structure, and as suggested by Sear, this may explain the relative ease or difficulty of crystallising various proteins.

Fig. 2(b) shows the phase behaviour for the model for the chosen parameter set. The solid lines in Fig. 2(b) are for the baseline model and include representative tie-lines. As $\beta \epsilon$ increases, the fluid-crystal phase transition widens (the re-entrant fluid phase expected at larger $\phi$ is not shown). There is a metastable fluid-fluid phase
FIG. 2: (a) Free energies for baseline model illustrating metastability of crystal branch (X) with respect to fluid branch (F) for $\beta \epsilon \lesssim 4.8$. (b) Phase behaviour in $(\phi, \beta \epsilon)$ plane for full model: solid lines are for the baseline model ($Q = c_s^{(R)} = 0$), dashed line is for $Q = 10$ and $c_s^{(R)} = 0$, and chained lines are for $Q = 10$ and $c_s^{(R)} = 0.2$ M. Other parameters are $n_s = 6$, $r_c = 1.05 \sigma$, $\theta_c = 0.45$.

Adding salt weakens and eventually destroys the effect. The salt concentration required to do this is $c_s \sim Q c_p$, as discussed already in section II. I now set $\beta \epsilon = 7.4$, marked by an arrow in Fig. 2(b), so that $B_2^{(0)}/B_2^{HS} = -2.7$ reproduces the value calculated in section III above [27]. Fig. 3(a) shows a typical phase diagram for the full model at this value of $\beta \epsilon$, as a function of added salt. Salt reservoir concentrations have been converted back to actual salt concentrations. The sloping tie-lines indicate a salt repartitioning effect in which the coion concentration is enhanced in the more dilute phases. In this simple model I have not taken into account excluded volume in the dense phases, so in reality the salt repartitioning would be more marked (see below for a further discussion on this). In this representation we can clearly see how adding salt broadens the fluid-crystal coexistence. Metastable fluid-fluid phase separation does not appear until salt concentrations $\gtrsim 2.3$ M for this value of $\beta \epsilon$. Note that the shape of the fluid-crystal separation for $\beta \epsilon \gtrsim 7.18$.

The effect of charges and added salt is obtained by inserting the baseline model into the general formalism in section II. The dashed line in Fig. 3(b) shows the fluid-crystal phase boundary at $Q = 10$ in the absence of salt. The transition has been markedly narrowed and the fluid-fluid transition moves to such a high value of $\beta \epsilon$ that it is no longer on the diagram. Repeating the calculation for $c_s^{(R)} = 0.2$ M obtains the chained lines in Fig. 3(b). The fluid-cystal phase boundary is intermediate between the zero salt limit and the baseline model, and the metastable fluid-fluid transition has reappeared in the diagram for $\beta \epsilon \gtrsim 9.14$. Finally as $c_s^{(R)} \to \infty$, the phase boundaries all move back to coincide with the baseline model. Thus the effect of charge in the model is to strongly suppress existing phase transitions in the absence of added salt.

FIG. 3: (a) Phase behaviour in $(\phi, c_s)$ plane for Sear’s model for $Q = 10$. (b) Fluid-crystal binodals for $Q = 8$ (triangles), 9.4 (diamonds) and 11.4 (circles). Other parameters are $n_s = 6$, $r_c = 1.05 \sigma$, $\theta_c = 0.45$, $\beta \epsilon = 7.4$. Other parameters are $n_s = 6$, $r_c = 1.05 \sigma$, $\theta_c = 0.45$.
binodal (which I interpret to be the same thing as the crystallisation boundary) is in agreement with common experience. This is a natural but non-trivial consequence of charge neutrality in the present model.

This calculation is now repeated for \( Q = 8.0, 9.4 \) and 11.4, which are the three values of lysozyme charge examined experimentally by Poon et al. Fig. 3(b) shows the crystallisation boundary as a function of added salt, for the three values of \( Q \). Clearly, the higher the charge, the more the phase transition is suppressed. Finally, these same phase boundaries are replotted in Fig. 4(a) as a function of the scaling variable \( c_s/Q^2 \). In this representation, the curves all collapse to lie on approximately the same quasi-universal crystallisation boundary. The scaling collapse is robust: if the calculations are repeated for different parameter values, the quasi-universal crystallisation curve moves up or down but a similar scaling collapse is always obtained.

The mean crystallisation boundary from Fig. 3(a) is shown in Fig. 3(b), which now includes the experimental data from Poon et al. There is reasonable agreement on the location of the quasi-universal crystallisation boundary, although it appears there is always some discrepancy between the shape of the theoretical and experimental boundaries at low \( \phi \) which recalls the discrepancy in the slope of the high-salt law in Fig. 3(b). It is worth re-emphasising that the present model is constrained to have the correct \( B_2 \), at least for \( c_s \gtrsim 0.25 \) M. The absence of a metastable fluid-fluid phase separation agrees with Poon et al who observe that no such phase transition occurs at the temperature of the experiments (22.5°C). However, a temperature decrease of just a few percent in the model, so that \( \beta \epsilon = 7.7 \) for example, is sufficient to bring fluid-fluid phase separation to accessible salt concentrations in the range 0.5–1.0 M. This is in accordance with the observations of Muschol and Rosenberger [28] (but additional caution is required since it is unlikely the only effect of temperature is through the value of \( \beta \epsilon \)). In the scaling representation of Fig. 3(a) such metastable fluid-fluid binodals also collapse to a quasi-universal curve—this is a prediction of the theory that would be interesting to test experimentally.

The scaling collapse in Fig. 3 occurs because the effects of charge and salt in fluid free energy are effectively combined into a single scaling variable \( c_s/Q^2 \), at salt concentrations in the vicinity of the crystallisation boundary. To show this is not just a feature of Sear’s model, I now turn briefly to the more commonly studied AHS model.

**B. Adhesive hard sphere model**

The adhesive hard sphere (AHS) model is also a suitable baseline model for the general theory in section II. In this model, hard spheres interact with a short range isotropic attractive potential. As noted by Rosenbaum et al [29], the phase behaviour is largely insensitive to the details of the potential provided the second virial coefficient

\[
B_2^{(0)} / B_2^{\text{HS}} = 1 - (4\tau)^{-1}.
\]  

(17)

The crystal phase is expected to be FCC since this has the greatest density of intersphere contacts. As the stickiness is switched off (\( \tau \to \infty \) or \( \beta \epsilon \to 0 \)) the fluid-crystal phase transition goes over into the usual HS freezing transition, which is a contrast to Sear’s model. The AHS ordered phase has always proved rather more difficult to
A link between the cell model for the crystal phase and the Baxter-Barboy treatment of the fluid phase can be made by matching $B_2$ of Eq. (17) with the exact expression

$$B_2^{(0)} / B_2^{HS} = 1 + 3 \int_1^\infty dx x^2 [1 - \exp(-\beta \xi x^{-n})]. \quad (18)$$

Fig. 5(a) shows the phase behaviour for the model. The solid line in this figure is the fluid-crystal phase boundary for the baseline model, and the dashed line is the corresponding boundary for $Q = 10$ with no salt. Similar to Fig. 5(b), the effect of charge is to narrow the fluid-crystal transition. As salt is added (not shown here), the boundary moves back towards that of the baseline model.

There is a complication that arises for Baxter’s solution for the AHS fluid phase free energy. Baxter’s theory involves the solution of a quadratic equation. If

$$\tau < \left( \frac{12\phi + 6\phi^2}{6(1-\phi)} \right)^{1/2} - 6\phi$$

then this quadratic equation has complex roots and Baxter’s theory becomes inadmissible. Normally this ‘danger zone’, shown shaded in Fig. 5(a), is happily hidden within the fluid-crystal two phase region. In the charged version though, it emerges into the single phase fluid region. For this reason one cannot choose $\tau > (2-\sqrt{2})/6 \approx 0.098$, the maximum of Eq. (19), and unfortunately this excludes $\tau = 0.067$ which would match $B_2^{(0)} / B_2^{HS} = -2.7$ from section II.

To demonstrate the scaling collapse for this model, I therefore choose $\tau = 0.1$ (equivalent to $B_2^{(0)} / B_2^{HS} = -1.5$ [23].) I repeat the calculations of the previous section to obtain the crystallisation boundaries (fluid-crystal binodal) as a function of added salt, for $Q = 8.0, 9.4$ and $11.4$ as used previously. Fig. 5(b) shows these boundaries plotted using the scaling variable $c_s/Q^2$. Again the curves collapse to a quasi-universal crystallisation boundary, similar to those in Fig. 5(a). This result confirms that the scaling collapse is a common feature for two different baseline models for the phase behaviour.

VI. SALT REPARTITIONING AND INTERFACE STRUCTURE

These ideas have some additional interesting physical implications for the equilibrium between crystals and solution which are explored in this section. The first implication concerns salt repartitioning. In the simple theory of section II, the product of coion and counterion concentrations is a constant as in Eq. (3). Each phase has to be electrically neutral, so that a phase enriched in protein is also enriched in counterions, and consequently depleted in coions (Donnan common ion effect). This explains the slope of the tie-lines in Fig. 4(a) for example. We can ask: is there any independent evidence for this phenomenon?
TABLE II: Lysozyme salt repartitioning data from Palmer et al [33]. Solution is 0.855 M NaCl / 0.2 M NaAc buffer (pH = 4.5). Ion concentrations in the third column are calculated by multiplying those in the second column by 1/(1 − ϕ), where ϕ = 0.64 is the crystal volume fraction. The charge unaccounted for in the crystal, assuming Q = 11.4, is 0.05 M or ≤ 3% of the total charge.

| conc / M | solution | crystal | crystal free volume |
|----------|----------|---------|------------------|
| Lysozyme<sup>Q−</sup> | 0.056 |        |                  |
| Ac<sup>−</sup> | 0.2 |        |                  |
| Na<sup>+</sup> | 1.055 | 0.19 | 0.53 |
| Cl<sup>−</sup> | 0.855 | 0.78 | 2.2 |
| Na<sup>+</sup> × Cl<sup>−</sup> | 0.9 |        | 1.2 |

Let me preface the answer to this by some cautionary remarks. Deviations from the simple Donnan equilibrium result may arise from three sources. Firstly excluded volume effects mean one should really consider the small ion concentrations in the available free volume. This will be an important consideration in the crystal phase where the volume fraction occupied by the protein is ~ 50%. Secondly, deviations from ideal solution behaviour can be expected at high salt concentrations—such deviations are usually absorbed into ‘activity coefficients’. Thirdly there may be significant specific ion effects, such as seen in the Hofmeister series.

Repartitioning of salt was studied in detail by Vekilov et al [34]. They found non-uniform salt concentrations inside protein crystals, but consider this is likely to be an effect of impurities or growth kinetics. The present theory only addresses the equilibrium repartitioning of salt in perfect lysozyme crystals though. Fortunately, some relevant equilibrium results for salt repartitioning can be found in recent experiments by Morozova et al [35] on cross-linked lysozyme crystals in contact with salt solutions. For example, Table II shows concentrations of Na<sup>+</sup> and Cl<sup>−</sup> in solution and in the crystal from the early work of Palmer et al [33], cited by Morozova et al. There is marked repartitioning, for example the coion concentration in the crystal free volume is about half that in the external solution. The product of the two ion concentrations is approximately constant though (final row in Table I), provided the protein excluded volume is taken into account. Detailed calculations by Morozova et al for their more recent data also take into account activity coefficients and give excellent agreement between theory and experimental results: they conclude “...for small ions capable of penetrating into the crystal channels [the] electrostatic (Donnan) potential controls the equilibrium internal concentration of ions in just the same way as in polyelectrolyte gels” [35]. By way of contrast, the same work also suggests Br<sup>−</sup> has a significant specific interaction with lysozyme.

The second implication concerns the electrical structure of the interface between crystal and solution phases.

In a Donnan membrane equilibrium [13, 14, 15], an electrostatic potential difference, the Donnan potential, develops between the two compartments on either side of the membrane. In the present theory for protein crystallisation, it is still true that a difference in the mean electrostatic potential develops between the crystals and the solution phase. Whether this should also be called a Donnan potential, a Galvani potential, or perhaps something else, can be debated [36, 37].

At any rate, the potential difference is readily calculated given the salt repartitioning. If c<sub>s</sub> and c<sub>c</sub> are the coion and counterion concentrations respectively, then c<sub>c</sub> = c<sub>s</sub><sup>R</sup> exp[±βκRφ], where φ is the mean electrostatic potential in the phase of interest measured relative to the salt reservoir (this result can also be used to recover Eq. (3)). The difference in mean potential between fluid (F) and crystal (X) phases, Δφ = φ<sub>F</sub> − φ<sub>X</sub>, is Δφ = (βe)<sup>−1</sup> log(c<sub>F</sub><sup>R</sup> / c<sub>X</sub><sup>R</sup>). For example, for the salt concentrations in Table II, c<sub>F</sub><sup>R</sup> = 1.055 M, c<sub>X</sub><sup>R</sup> = 0.53 M (in the free volume), and therefore Δφ ≈ 17 mV. The potential difference arises because an electrical double layer is formed at the crystal-solution interface and is intimately connected with salt repartitioning. The details of this will be discussed in a forthcoming paper [38].

Another consequence of the Donnan potential is that the pH in the crystal will be 0.434Δφ higher than in the solution (where 0.434 = log<sub>10</sub> e). If the solution pH is below the isoelectric point, as is often the case for lysozyme, one might expect this to lower the charge per protein in the crystal although it is unlikely that the charge can be determined with any accuracy from such a naïve calculation.

Pair-potential theories miss both the effects of salt repartitioning and the potential difference between the two phases, which are essentially many-body phenomena [24, 14]. Whilst this is not a problem at high salt where Hill’s mapping between the McMillan-Mayer / Debye-Hückel pair potential approach and Donnan’s method goes through [39], the effects can be significant at lower salt concentrations. The importance of the effects can be gauged by comparing the magnitude of Δφ to the thermal energy kT. If Δφ ≳ kT/e then there may be significant errors introduced by using a single effective pair potential (such as a DLVO potential) for both the crystal and the solution phases. On the other hand, if Δφ ≪ kT/e, the use of pair potentials cannot be criticised on these grounds. It can easily be shown that Δφ ≳ kT/e corresponds to the by now familiar c<sub>s</sub> ≳ Qe<sub>p</sub>, in other words salt concentrations smaller than the protein charge density in the crystal, which is typically 0.5 M.

VII. CONCLUSIONS

There are several conclusions from the present work. The first concerns the lysozyme / NaCl system. The
quasi-universal crystallisation boundary observed by Poon et al. [10] can be fit reasonable well by the present version of Sear’s model with ~ 6 sticky contacts per molecule, similar to the conclusions of other workers [24, 25].

The scaling collapse for \( B_2 \) noticed by Egelhaaf and Poon [2] is here attributed to a contribution of a ‘bare’ \( B_2^{(0)} \) which is approximately independent of salt concentration above about 0.25 M. In this system therefore, NaCl appears to be acting as an indifferent electrolyte in the sense that it does not seem to exhibit specific ion effects. This appears to be confirmed by the experiments of Morozova et al. [35] discussed in the preceeding section. Other salts, for example NaBr, may not behave the same way of course.

The other conclusions are general ones. Firstly, a simple extension of existing models to incorporate salt ions and charge neutrality provides a straightforward explanation for the shape of protein crystallisation boundaries and the associated scaling properties seen for lysozyme.

Even if the present theory proves inadequate to describe the experiments in quantitative detail, this is surely a robust observation. Secondly, there are a large number of effects which have been omitted from the present theory, but which could be incorporated with some additional numerical effort, such as excluded volume effects, non-ideality of the salt ions, and other electrostatic correlation effects. Finally, the twin phenomena of salt repartitioning and the concomitant appearance of a significant Donnan potential difference discussed in the previous section are essentially many-body effects, and are not captured in simple pair-potential theories.

For discussions and correspondence, I thank S. U. Egelhaaf, M. E. Fisher, H. N. W. Lekkerkerker, M. G. Noro, W. C. K. Poon and R. P. Sear. Finally, it is a great pleasure to dedicate this paper to Peter Pusey on the occasion of his 60th birthday.

---

[1] McPherson A 1999 Crystalisation of biological macromolecules (New York: Cold Spring Harbor).
[2] Piazza R 2000 Curr. Opinion Coll. Int. Sci. 5 38.
[3] Poon W C K 1997 Phys. Rev. E 55 3762.
[4] Lekkerkerker H N W 1997 Physica A 244 227.
[5] Rosenbaum D, Zamora P C and Zukoski C F 1996 Phys. Rev. Lett. 76 150.
[6] Marr D W and Gast A P 1993 J. Chem. Phys. 99 2024.
[7] Tejero C F and Baus M 1993 Phys. Rev. E 48 3793.
[8] Hagcn M H J and Frenkel D 1994 J. Chem. Phys. 101 4093.
[9] Sear R P 1999 J. Chem. Phys. 111 4800.
[10] Poon W C K, Egelhaaf S U, Beales P A, Salonen A and Sawyer L 2000 J. Phys. Cond. Mat. 12 L569.
[11] Guo B, Kao S, McDonald H, Asanov A, Combs L L and Wilson W W 1999 J. Cryst. Growth 196 424.
[12] Egelhaaf S U and Poon W C K private communications. Data for \( B_2 \) is collected from Guo et al. [1], Piazza R and Pieno M 2000 J. Phys. Cond. Mat. 12 A443. Rosenbaum et al. [2]. Skouri cited in Piazza and Pieno, and Velev O D, Kaler E W and Lenhoff A M 1998 Biophys. J. 75 2682.
[13] Donnan F G 1911 Z. Elektrochem. 17 572.
[14] Rice S A and Nagasawa M 1961 Polyelectrolyte solutions (London: Academic Press) ch 8.
[15] Morawetz H 1965 Macromolecules in solution (New York: Wiley Interscience) ch 7.
[16] McMillan W G and Mayer J E 1945 J. Chem. Phys. 13 276.
[17] Belloni L 2000 J. Phys. Cond. Mat. 12 R549.
[18] Hill T L 1956 Disc. Faraday Soc. 21 31.
[19] Brodie M L, Tomine T M and Saxowsky M D 1996 Phys. Rev. E 53 6325.
[20] Warren P B 1997 J. Phys. II (France) 7 343.
[21] Rosenbaum D F, Kulkarni A, Ramakrishnan S and Zukoski C F 1999 J. Chem. Phys. 111 9422.
[22] Ref. [3] suggests alternatively that the logarithm of the protein solubility should be proportional to ionic strength, but this is a poorer fit to the solubility data of Guo et al. [1] than Eq. [2].
[23] Vliegenhart G A and Lekkerkerker H N W 2000 J. Chem. Phys. 112 5364.
[24] Curtis R A, Blanch H W and Prausnitz J M 2001 J. Phys. Chem. B 105 2445.
[25] Zukoski, C F 1999 J. Chem. Phys. 112 9822.
[26] Louis A A 2001 Phil. Trans. Roy. Soc. A 359 939.
[27] Oki et al. suggest ‘macrobond’ energies of the order 50 kcal mol\(^{-1}\) \(\approx 80 kT \gg \beta c\) but this estimate is for the bare macrobond energy and does not include re-solvation of the protein surface.
[28] Muschol M and Rosenberger F 1997 J. Chem. Phys. 107 1953.
[29] Noro M G, Kern N and Frenkel D 1999 Europys. Lett. 48 332.
[30] Barboy B 1974 J. Chem. Phys. 61 3194. Barboy’s definition of \(\tau\) differs from the present use and Baxter’s original work by a factor 6.
[31] Baxter R J 1968 J. Chem. Phys. 49 2770.
[32] Daanoun A, Tejero C F, and Baus M 1994 Phys. Rev. E 50 2913.
[33] Palm K J, Ballantyne M and Galvin J A 1948 J. Am. Chem. Soc. 70 906.
[34] Vekilov P G, Monaco L A, Thomas B R, Stojanoff V and Rosenberger F 1996 Acta Cryst. D 52 785.
[35] Morozova T Ya, Kachalova G S, Lanina N F, Evtodienko O D, Kaler E W and Lenhoff A M 1998 Biophys. Chem. 75 114.