Charge pattern matching as a ‘fuzzy’ mode of molecular recognition for the functional phase separations of intrinsically disordered proteins

Yi-Hsuan Lin¹, Jacob P Brady², Julie D Forman-Kay³ and Hue Sun Chan⁴

¹ Department of Biochemistry, University of Toronto, and Molecular Medicine, Hospital for Sick Children, Toronto, Ontario, Canada
² Departments of Molecular Genetics, Biochemistry, and Chemistry, University of Toronto, Toronto, Ontario, Canada
³ Molecular Medicine, Hospital for Sick Children, and Department of Biochemistry, University of Toronto, Toronto, Ontario, Canada
⁴ Departments of Biochemistry and Molecular Genetics, University of Toronto, Toronto, Ontario, Canada

E-mail: chan@arrhenius.med.toronto.edu

Keywords: random phase approximation polymer theory, liquid–liquid phase separation, membraneless organelles, effective medium approximations

Abstract

Biologically functional liquid–liquid phase separation of intrinsically disordered proteins (IDPs) is driven by interactions encoded by their amino acid sequences. Little is currently known about the molecular recognition mechanisms for distributing different IDP sequences into various cellular membraneless compartments. Pertinent physics was addressed recently by applying random-phase-approximation (RPA) polymer theory to electrostatics, which is a major energetic component governing IDP phase properties. RPA accounts for charge patterns and thus has advantages over Flory–Huggins (FH) and Overbeek–Voorn mean-field theories. To make progress toward deciphering the phase behaviors of multiple IDP sequences, the RPA formulation for one IDP species plus solvent is hereby extended to treat polyampholyte solutions containing two IDP species plus solvent. The new formulation generally allows for binary coexistence of two phases, each containing a different set of volume fractions, for the two different IDP sequences. The asymmetry between the two predicted coexisting phases with regard to their ratios for the two sequences increases with increasing mismatch between their charge patterns. This finding points to a multivalent, stochastic, ‘fuzzy’ mode of molecular recognition that helps populate various IDP sequences differentially into separate phase compartments. An intuitive illustration of this trend is provided by FH models, whereby a hypothetical case of ternary coexistence is also explored. Augmentations of the present RPA theory with a relative permittivity that depends on IDP volume fraction lead to higher propensities to phase separate, in line with the case with one IDP species we studied previously. Notably, the cooperative, phase-separation-enhancing effects predicted by the prescriptions for we deem physically plausible are much more prominent than that entailed by common effective medium approximations based on Maxwell Garnett and Bruggeman mixing formulas. Ramifications of our findings on further theoretical development for IDP phase separation are discussed.

1. Introduction

Nearly two decades of increasingly intensive research established that intrinsically disordered proteins/protein regions (abbreviated collectively as IDPs here) serve many important biological functions, and are especially critical for signaling and regulation in multicellular organisms [1–12]. Recently, it was discovered that IDPs function not only at the level of individual molecules. An expanding repertoire of IDPs have been seen to undergo liquid–liquid phase separation in vitro, intriguingly paralleling the formation of many types of condensed liquid/gel-like bodies/organizations in living organisms, including extracellular materials, transcription complexes, nucleating sites of intermediate filament organization, and various membraneless...
organelles. It is apparent from these experimental observations that IDP condensation constitutes a major physical underpinning of these condensed bodies, which serve as hubs for specifically regulated sets of biomolecules to interact. As such, IDP phase separation is one of Nature’s means to achieve the spatial and temporal compartmentalization necessary for the organization of vital processes [13–32]. Although much detail remains to be ascertained, examples of membraneless organelles and an IDP species whose phase separation has been found to be a likely contributor to their assembly include chromatoid bodies, nuage or germ granules in mammalian male germ cells [33] and the DEAD-box RNA helicase Ddx4 [20], the Caenorhabditis elegans germline-specific perinuclear RNA granules known as P-granules [34] and the Ddx3 RNA helicase LAF-1 [21], as well as the stress granules triggered by integrated stress response [35] and the RNA-binding heterogeneous nuclear ribonucleoprotein A1 (hnRNPA1) [22]. Because of the importance of these bodies to biological regulation, malfunctioning of the corresponding IDP phase separation processes can lead to deregulation and diseases, including cancer due to loss of regulation of stress granules [36], protein fibrillization and thus amyloid diseases [22], and various forms of neurological disorder [37–39].

1.1. Seeking ‘sequence-phase’ relationships

With the advent of IDPs, the molecular biology paradigm of seeking ‘sequence-structure’ relationships for globular proteins has to be expanded to encompass ‘sequence-ensemble’ relationships for individual IDPs [40–42]. Now, the additional question we need to ask is: How do the phase behaviors of IDPs depend on their amino acid sequences? In other words, what are the ‘sequence-phase’ relationships? [43] Although computational study of IDPs is still in its infancy, much insight into the conformational properties and binding energetics of individual IDPs has been gained by explicit-chain simulations [8, 9, 44–49]. In contrast, because IDP phase separation is a multiple-chain property, computationally it is extremely costly to simulate using fully atomic explicit-chain models [30], notwithstanding promising progress made by coarse-grained approaches that treat groups of amino acid residues of IDPs as interaction modules in continuum space [51] or on lattices [27] and simulation algorithms developed recently [52, 53] for phase separations of globular proteins [54]. In this context, analytical theories of IDP phase separation are valuable not only because of their tractability, but also—and more importantly—for the conceptual framework they offer for understanding a highly complex phenomenon.

1.2. Mean-field and random-phase-approximation (RPA) theories of phase coexistence

Developed mainly for synthetic polymers at its inception, the basic statistical mechanical framework of Flory–Huggins (FH) theory [55, 56] is useful for describing phase separation in the biomolecular context [17, 20, 57, 58]. FH assumes that the interactions among the monomers (residues) of the polymers have short spatial ranges. In view of the fact that not only short-range interactions such as solvent-mediated hydrophobic effects but long-range Coulomb forces are important for driving the phase separation of certain IDPs, it has been suggested [58] that the Overbeek–Voorn (OV) theory [59, 60] should be more appropriate in those cases. Inasmuch as sequence dependence is concerned, however, both FH and OV are mean-field theories that account only for composition but not sequence information. In these theories, all residues belonging to any given set of chain sequences are allowed to interact on equal footing irrespective of the correlation dictated by chain connectivity. Therefore, to address sequence specificity of IDP phase separation, one needs to go beyond FH and OV. Accordingly, we recently put forth a RPA theory that approximately accounts for the effects of arraying different charge patterns along the IDP chain sequence [61–63]. The term ‘RPA’ was first introduced by Bohm and Pines in their quantum mechanical collective description of electron interactions [64]. The analogy of this approach with the approximate polymer theory that considers terms up to quadratic in particle density was recognized by de Gennes [65]. As detailed elsewhere [62], the RPA formulation we developed [61], which follows largely that of Olvera de la Cruz [66, 67], is successful in providing a physical rationalization [61, 62] for the experimental salt dependence of Ddx4 phase separation as well as the difference in phase behavior between the wildtype and a charge-scrambled variant of Ddx4 [20]. Our RPA theory suggests further that the tendency for a collection of IDP chains with a given charge sequence to phase separate is strongly—but negatively—correlated with the conformational dimensions of individual IDP molecules of the sequence [63], and that both of these properties are well correlated with the charge pattern parameter $\kappa$ of Das and Pappu [68] and the ‘sequence charge decoration’ parameter SCD of Sawle and Ghosh [69]. In principle, these predictions are now testable using experimental techniques similar to those employed to study ‘IDP polymers’ [70].

1.3. Sequence-dependent multiple-component IDP phase separation

Membraneless organelles are complex functional units consisting of many protein and nucleic acid components [18]. Different types of such units are enriched with different varieties of proteins and nucleic acids. Some individual membraneless organelles have mesoscopic substructures with different degrees of fluidity, as in the
case of stress granules [71]. In a similar vein, immiscible liquid phases have been shown to contribute to subcompartmentalization of the nucleolus [27]. Clearly, a viable spatial organization of cellular processes necessitates a heterogeneous distribution of different biomolecular components into different membraneless organelles and their substructures, rather than having all IDP species condensing into a big gemisch. How is this achieved physically?

One aspect of this question was addressed recently using a simple cubic lattice model of multicomponent mixtures confined to a \(6 \times 6 \times 6\) box, with each component represented by a bead on the lattice [72]. By considering hypothetical intercomponent interaction strengths (contact energies), the authors found that with sufficient heterogeneity in contact energies, demixed domains are likely to segregate. This and other results of this big-picture study suggest that phase separation into cellular compartments with different compositions is a robust consequence of interaction heterogeneity among biomolecules [72, 73].

With this in mind, the logical next step in our pursuit of sequence-phase relationships is to ascertain how interaction heterogeneity is encoded genetically. For globular protein folding, the fact that the amino acid alphabet is finite [74] implies that there are physical limits to interaction heterogeneity and structural encodability, as has been illustrated by simple exact models [75, 76]. Similarly, physical limits should exist in the ability of different IDP sequences to demix. Taking a step toward deciphering what is physically achievable, here we present an RPA formulation for the phase behavior of two charged sequences as models for two IDP species. Consistent with physical intuition, we found that the tendency for the two IDP species to demix in two coexisting phases increases with increasing difference in their charge patterns. This phenomenon represents a statistical, multivalent mode of molecular recognition for cellular organization that differs from the structurally highly specific form of recognition among folded proteins but share similarities with the ‘fuzzy complexes’ [77–79] involving individual IDP molecules [80–83].

Delving deeper into the role of electrostatics in IDP phase separation, we have also extended the two-sequence RPA formulation to address how a relative permittivity \(\epsilon_r(\phi)\) that depends on IDP volume fraction \(\phi\) may affect IDP phase properties. Several common effective medium approximations [84] posit a gradual decrease in \(\epsilon_r\) from the \(\epsilon_r(\phi = 0) \approx 80\) value for pure water with increasing \(\phi\). But physical consideration [62] and experimental volumetric measurements suggest a much sharper decrease, with \(\epsilon_r(\phi = 0.2) \approx 20\), leading to large cooperative effects that enhance phase separation significantly. These findings and their ramifications are detailed below.

2. Methods

2.1. Theoretical development of the RPA formulation

The development of RPA theory for a pair of charged sequences constitutes the bulk of the results presented in subsequent sections of this article. This effort is based on an extension of the RPA formulation for a single sequence that we put forth recently [61, 62].

2.2. Experimental determination of dissolved protein volumes

To address the effect of volume of dissolved proteins on the relative permittivity of the resulting aqueous solution, nuclear magnetic resonance (NMR) and absorbance measurements were performed on two folded globular proteins bovine serum albumin (BSA) and hen egg white lysozyme (HEWL) as well as two IDPs Ddx414FtoA and Ddx4cond, which are, respectively, a mutant of Ddx4 in which all 14 phenylalanines are mutated to alanines and the concentrated phase of phase-separated wildtype Ddx4.

2.3. Measurement of water content of protein samples

NMR spectra were recorded using a Bruker Ascend III spectrometer at 14.0 T equipped with a cryogenically cooled triple resonance gradient probe. Spectra were processed using NMRPipe [85]. 1D \(^1\)H spectra were recorded on protein samples over a range of concentrations (5–400 mg ml\(^{-1}\)) and the integrated water signals were compared with the corresponding integrals obtained from a spectrum recorded of buffer (the same buffer composition as used for each protein sample). BSA was purchased from Sigma and samples were prepared in 20 mM sodium phosphate (NaPi), 100 mM sodium chloride (NaCl), 10% \(\text{H}_2\text{O}/90% \text{H}_2\text{O}, \text{pH} 6.5\). HEWL was purchased from BioBasic and samples were dissolved in 20 mM sodium citrate, 100 mM NaCl, 10% \(\text{H}_2\text{O}/90% \text{H}_2\text{O}, \text{pH} 5\) (lower pH was used due to limited solubility of HEWL in NaPi at pH 6.5). Ddx414FtoA samples were prepared according to [86] and dialysed against 20 mM NaPi, 100 mM NaCl, 5 mM tris(2-carboxyethyl) phosphine (TCEP), 10% \(\text{H}_2\text{O}/90% \text{H}_2\text{O}, \text{pH} 6.5\). For Ddx4cond the same buffer was used but the NaCl concentration was varied between 100 and 400 mM in order to generate samples with protein concentrations between 200 and 400 mg ml\(^{-1}\). For phase-separated samples, it was ensured that the entirety of the probe coil was occupied by the condensed phase, thus avoiding contaminating signals from the more hydrated dilute...
The present study is restricted to IDP sequences with zero net charge. Counterions are not considered. Possible phase behaviors of a three-component liquid system are outlined in figure 1. In general, the system can be a homogeneous solution (figure 1(b)), or it can separate into two coexisting phases (binary coexistence; figures 1(c)–(e)), or separate into three coexisting phases (ternary coexistence; figure 1(f)). Fundamentally, phase behavior is governed by the intra- and inter-component interactions as well as environmental conditions such as temperature and pressure. Our theories below provide a rudimentary physical account of how interactions among IDP chains with two different amino acid sequences affect the conditions under which binary and ternary coexistence emerge. The theories presented here are for solution systems with an infinite effective volume. As such, our theories account for the differences among scenarios typified by the leftmost drawings in figures 1(c)–(f) but they are not equipped to address details such as droplet size and geometry. In other words, they provide no discrimination among different droplet geometries along a given horizontal row in figure 1. Accounting for the latter would require additional modeling of the interfacial tensions between different solution phases [27].

4. RPA theory for two charged sequences

Based on our previous RPA formulation for a single sequence [61, 62], the approach is now extended to consider two model polypeptide (IDP) sequences $s_1$ and $s_2$ in a salt-free aqueous solution. Using notation similar to before [62], the electric charges along the sequences are written as $\{\sigma_i\} \equiv \{\sigma_i^{(1)}$, $\sigma_i^{(2)}$, $\sigma_i^{(3)}$, ..., $\sigma_i^{(N)}\}$ and $\{\sigma_j\} \equiv \{\sigma_j^{(1)}$, $\sigma_j^{(2)}$, $\sigma_j^{(3)}$, ..., $\sigma_j^{(N)}\}$, where $N_1$ and $N_2$ are the numbers of residues in $s_1$ and $s_2$, respectively. The corresponding volume fractions of the IDPs in solution are denoted as $\phi_1$ and $\phi_2$. The present study is restricted to IDP sequences with zero net charge. Counterions are not considered.

4.1. Free energy as a function of two IDP volume fractions

Following the FH lattice argument [55], we partition the spatial volume $V$ of the solution system into lattice units, $a^3$, that corresponds to the volume of a solvent molecule. Accordingly, the RPA free energy per unit volume in units of the product of Boltzmann constant $k_B$ and absolute temperature $T$ is cast as the per-lattice-site quantity

$$f (\phi_1, \phi_2) = \frac{F_{\text{RPA}} a^3}{V k_B T} = -s(\phi_1, \phi_2) + f_{\text{el}}(\phi_1, \phi_2), \quad (1)$$

where the negative entropy $-s$ is the entropic contribution to free energy in units of $k_B T$. This term is given by the standard FH entropy of mixing for a system comprising of $s_1, s_2$, and solvent:

$$-s(\phi_1, \phi_2) = \frac{\phi_1}{N_1} \ln \phi_1 + \frac{\phi_2}{N_2} \ln \phi_2 + \ln (1 - \phi_1 - \phi_2) \ln (1 - \phi_1 - \phi_2), \quad (2)$$

where $1 - \phi_1 - \phi_2$ is the volume fraction of solvent. The electrostatic contribution $f_{\text{el}}$ is calculated by RPA [61, 62], viz.,

$$f_{\text{el}}(\phi_1, \phi_2) = \int_0^{\infty} \frac{\mathrm{d}k k^2}{4 \pi^2} \left[ \ln [1 + G(k)] - G(k) \right], \quad (3)$$

where $k$ is the reduced wave number that absorbs the virtual bond length $b \simeq a$ of the polypeptide backbone by re-defining $kb = \tilde{k}$ in [62] as $k$ (i.e., $kb \to k$), such that

$$G(k) = \frac{4 \pi}{k^2 (1 + k^2) T^*} \langle q | \hat{G}_0 | q \rangle, \quad (4)$$

where $4 \pi/[k^2 (1 + k^2)]$ is from the Fourier transformation of Coulomb interaction with a short-range cutoff [61, 67] in units of $k_B T$. Spectra were recorded using both small flip angle ($\theta < 10^\circ$) and $\theta = 90^\circ$ pulses with very similar results in both cases.

2.4. Measurement of protein concentration

Protein concentrations were determined by absorbance at 280 nm ($A_{280}$) after dilution into 6 M guanidinium HCl, 20 mM NaPi, pH 6.5 using the Beer–Lambert law with extinction coefficients of 23 950, 23 950, 36 000, and 44 309 $M^{-1} \text{cm}^{-1}$ for wildtype Ddx4, Ddx414FtoA, HEWL, and BSA, respectively [87].

3. Overview of three-component phase behaviors

Accounting for the latter would require additional modeling of the interfacial tensions between different solution phases [27].
\[ U_\text{23}(r) = \frac{e^2}{4\pi \varepsilon_0 \varepsilon_r k_B T} \frac{1 - e^{-r/b}}{r} , \]

\( \varepsilon_0 \) is vacuum permittivity and \( \varepsilon_r \) is relative permittivity, \( T^* \equiv b/l_b \) is the reduced temperature defined by Bjerrum length \( l_b = e^2/(4\pi \varepsilon_0 \varepsilon_r k_B T) \). Here \( |q| \) is the \( (N_1 + N_2) \)-dimensional column vector representing the two charge sequences, namely \( q_i = \sigma_i^{(1)} \) for \( 1 \leq i \leq N_1 \) and \( q_i = \sigma_i^{(2) - \text{tot}} \) for \( N_1 + 1 \leq i \leq N_1 + N_2 \), \( |q| \) is the transposed row vector, \( \langle q| \tilde{G}_k |q \rangle = \sum_{ij} q_i (\tilde{G}_k)_{ij} q_j \) with \( (\tilde{G}_k)_{ij} \) being the \( i, j \) element of the \textit{bare} two-body correlation matrix \( \tilde{G}_k \) of all possible sequence-sequence correlations [62],

\[ \tilde{G}_k = \begin{pmatrix} \tilde{G}_{11}(k) & \tilde{G}_{12}(k) \\ \tilde{G}_{21}(k) & \tilde{G}_{22}(k) \end{pmatrix}, \]

and \( \tilde{G}_{12}(k) = \tilde{G}_{21}(k) \). As in our previous studies [61, 62], we consider a simple formulation in which all IDPs are modeled as Gaussian chains without excluded volume within the RPA formalism [66]. In this approximation, there is no correlation between the positions of different chains; hence \( \tilde{G}_{12}(k) = \tilde{G}_{21}(k) = 0 \) and

Figure 1. Schematics of three-component phase separation scenarios. (a) The three pure components considered in our analysis—water solvent, IDP sequence 1 (seq1), and IDP sequence 2 (seq2)—in a totally demixed configuration. (b) A homogeneous solution of all three components. In this and subsequent drawings in this figure, the colors of various solution phases are resultant colors of mixing the pure components (white, red, and green) in proportions consistent with the given scenario. (c)–(e) Binary coexistence. (c) One of the coexisting phases is dilute in both seq1 and seq2 (top), the other is concentrated in both seq1 and seq2 (bottom). (d) The concentration of seq2 is constant throughout, whereas seq1 is dilute in one phase (top) and concentrated in the other (bottom). (e) One phase is concentrated in seq2 but dilute in seq1 (top), the other phase is concentrated in seq1 but dilute in seq2 (bottom). (f) Ternary coexistence. In this example, the system separates into a dilute phase for both seq1 and seq2 (top), a phase concentrated in seq2 but dilute in seq1 (bottom), with water present in all three phases. Circle(s)-in-square drawings to the right of (c)–(f) depict possible configurations of phase-separated droplets in the given scenario. In each case, the rightmost schematic phase diagram illustrates the manner in which a given set of bulk concentrations (bulk volume fractions for seq1 and seq2) of \( (\phi_1, \phi_2) \) represented by the central dot with the color of (b) is separated (arrows) into two or three phases with different \( (\phi_1, \phi_2) \)'s. The volume fraction of water is equal to \( 1 - \phi_1 - \phi_2 \).
\[
\hat{G}_{11}(k)_{ij} = \frac{\phi_i}{N_1} \exp \left( -\frac{1}{6} k^2 |i - j| \right)
\]
\[
\hat{G}_{22}(k)_{ij} = \frac{\phi_j}{N_2} \exp \left( -\frac{1}{6} k^2 |i - j| \right)
\]

(7)

follow from the average of \(\exp(i \mathbf{k} \cdot \mathbf{R}_{ij})\) over a Gaussian chain ensemble wherein \(\mathbf{R}_{ij}\) is the vector between chain positions \(i, j\) and \(k^2 = \mathbf{k} \cdot \mathbf{k}\) (equation (IX.59) of [65]). Equation (4) then becomes

\[
\hat{G}(k) = \frac{4\pi}{k^2(1 + k^2)} T^\ast \left[ \langle \sigma_i \rangle \hat{G}_{11}(k) |\sigma_i\rangle + \langle \sigma_2 \rangle \hat{G}_{22}(k) |\sigma_2\rangle \right],
\]

(8)

where

\[
\langle \sigma_i \rangle \hat{G}_{11}(k) |\sigma_i\rangle = \frac{\phi_i}{N_1} \sum_{i,j=1}^N \sigma_i^{(i)} \sigma_i^{(j)} \exp \left( -\frac{1}{6} k^2 |i - j| \right),
\]

(9a)

\[
\langle \sigma_2 \rangle \hat{G}_{22}(k) |\sigma_2\rangle = \frac{\phi_j}{N_2} \sum_{i,j=1}^N \sigma_2^{(i)} \sigma_2^{(j)} \exp \left( -\frac{1}{6} k^2 |i - j| \right).
\]

(9b)

4.2. Free energy landscape and spinodal instability

As an example, we first apply this formulation to two \(N_1 = N_2 = 50\) charged IDP sequences corresponding to sv28 and sv24 in Das and Pappu [68]. The sequences are labeled here as seq1 and seq2 respectively. The RPA free energy function for the two sequences (figure 2) consists of regions of different curvatures: parts of the landscape are convex downward (i.e., has a convex downward curvature) whereas some other parts are convex upward (concave downward).

If a given IDP solution has overall (bulk-state) IDP volume fractions \((\phi_1, \phi_2)\) situated in a convex-downward region, the state is thermodynamically stable and thus the bulk-state volume fractions are maintained. In contrast, if the bulk state is in a concave-downward region, the state is thermodynamically unstable because a phase-separated state allows for a lower free energy. Accordingly, the system undergoes phase separation to multiple coexisting phases with different protein volume fractions [62].

The boundary between the convex and concave regions is determined by the saddle point condition [62, 88]

\[
\det \hat{\mathbf{F}} \equiv \begin{vmatrix}
\frac{\partial \hat{F}}{\partial \phi_1} & \frac{\partial \hat{F}}{\partial \phi_2} \\
\frac{\partial \hat{F}}{\partial \phi_2} & \frac{\partial \hat{F}}{\partial \phi_1}
\end{vmatrix} = 0.
\]

(10)

This boundary defines a spinodal instability region in which \(\det \hat{\mathbf{F}} < 0\). Because the determinant of a matrix is equal to the product of all its eigenvalues, this instability condition indicates that one, but not both, of the eigenvalues of \(\hat{\mathbf{F}}\) is negative. This means that second-order perturbations of free energy with respect to \(\phi_1, \phi_2\) along the direction of the corresponding eigenvector diverge, signaling that the system cannot maintain a homogeneous phase.

Note that although the opposite of the \(\det \hat{\mathbf{F}} < 0\) condition, viz., \(\det \hat{\mathbf{F}} > 0\), does not by itself exclude the possibility that both eigenvalues of \(\hat{\mathbf{F}}\) are negative and thus the system is thermodynamically unstable despite not satisfying \(\det \hat{\mathbf{F}} < 0\) (e.g., at a local maximum), exhaustive numerical searches \((\phi_1, \phi_2 = 0.001, 0.002, \ldots, 0.999)\) did not find any such instance for all RPA and FH systems studied in this work. Hence, for these systems, equation (10) is the valid condition for spinodal boundaries, examples of which are shown as dashed curves in figure 2 for the (seq1, seq2) system.

In two-component systems such as those consisting of a single IDP species and solvent molecules, spinodal instability necessarily leads to binary coexistence [62]. In systems with more than two components, more coexisting phases are allowed. According to the Gibbs phase rule, the maximum number of coexisting phases under a given set of environmental conditions is equal to the number of components in the system [62, 89] but, depending on system specifics, the actual number of coexisting phases can be smaller. Spinodal instability always implies that the system existing as a single phase is thermodynamically untenable and thus phase separation must occur. However, when the number of components is larger than two, the precise number of coexisting phases is governed by how total free energy varies with changes in the component volume fractions in different combination of phases or, equivalently, by the balance of chemical potentials for each component across different phases [62, 88]. As examples, three instances of binary coexistence in the (seq1, seq2) system are depicted in figure 2 as pairs of dots connected by solid tie lines. The general procedure for determining the conditions for such coexistence is as follows.
4.3. Binary coexistence in three-component systems: applications to IDPs with two different sequences plus solvent

As mentioned, we use two methods to determine phase equilibrium: by ascertaining the minimum free energy among single- and multiple-phase states \[90\], and by balancing the chemical potentials for each of the components across different phases \[56, 59, 91\]. The two approaches are mathematically equivalent. They can be applied simultaneously to yield more accurate numerical results. Here we describe in detail how the two approaches are applied to study three-component systems that undergo binary phase separation.

A system is dictated by thermodynamics to seek its lowest free energy state. Whether a system phase separates can be ascertained by comparing its overall free energy with and without phase separation. For a system with bulk IDP volume fractions \(\phi_1, \phi_2\), the free energy \(f_{\text{bulk}}\) without phase separation and the free energy \(f_{\text{sep}}\) for separating into two phases (labeled as \(\alpha\) and \(\beta\)) that take up fractional volumes \(v_\alpha = \nu (0 \leq \nu \leq 1)\) and \(v_\beta = 1 - \nu\) of the total system volume and with IDP volume fractions \(\phi_1^{\alpha}, \phi_2^{\alpha}\) and \(\phi_1^{\beta}, \phi_2^{\beta}\), respectively, are given by

\[
 f_{\text{bulk}} = f (\phi_1^0, \phi_2^0),
\]

\[
 f_{\text{sep}} = f (\phi_1^{\alpha}, \phi_2^{\alpha}),
\]

\[
 f_{\text{sep}} = f (\phi_1^{\beta}, \phi_2^{\beta}),
\]

Figure 2. RPA theory of phase separation for a pair of sequences. The sequences, shown along the axes in (a) and (c), wherein positive and negative charges are depicted as red and blue beads respectively, correspond to sv28 and sv24 in Das and Pappu \[68\]. \(\phi_1\) and \(\phi_2\) are their volume fractions, respectively, as the sequences are re-labeled as seq1 and seq2 in this work (table 1). Results in this figure are for \(T^* = 4\). (a) Free energy landscape. The plotted quantity is the free energy \(f (\phi_1, \phi_2)\) in equation (1) minus a linear function \(a_1 \phi_1 + a_2 \phi_2\) of \(\phi_1, \phi_2\), where the coefficients are chosen to be \(a_1 = -1.1447\) and \(a_2 = -1.1453\). The sole purpose of subtracting this linear term is to graphically highlight the changes in curvature on the landscape. The subtraction has no effect on the determination of phase coexistence (section 4.3). As examples of phase coexistence, three pairs of phase-separated states are marked by gray dots, whereby each pair is connected by its own tie line (gray solid line). (b) View of the landscape in (a) from an elevated vantage. The two dashed curves are spinodal phase boundaries defined by equation (10) (one of the boundaries is very close to the origin with \(\phi_1, \phi_2\) intercepts \(\approx 0.003\), see zoom-in view in (d)); the region between these two boundaries satisfies \(\det \tilde{\mathbf{F}} < 0\). The three pairs of dots marking coexisting phases connected by tie lines are the same as those in (a). Note that subtracting a linear function of \(\phi_1, \phi_2\)'s from \(f (\phi_1, \phi_2)\) does not alter the spinodal boundaries determined by the matrix \(\tilde{\mathbf{F}}\) that contains only second derivatives of \(\phi_1, \phi_2\)'s. (c) Heat map representation (scale on the right) of the function \(10^4 \times [f (\phi_1, \phi_2) - a_1 \phi_1 - a_2 \phi_2]\) in (a). Pairs of coexisting (\(\phi_1, \phi_2\)) connected by tie lines and spinodal boundaries are the same as those in (b). (d) Zoom-in plot of the \(0 \leq \phi_1, \phi_2 \leq 0.005\) region in (c) to provide a clearer view of the spinodal phase boundary that is very near to the origin (dashed line) and the dilute phases of the three coexisting pairs.
\[ f_{\text{sep}} = \nu f(\phi_1^0, \phi_2^0) + (1 - \nu)f(\phi_1^2, \phi_2^2), \]  

(11b)

wherein conservation of volume of each of the components implies that

\[ \nu \phi_1^0 + (1 - \nu) \phi_2^0 = \phi_1^0, \]  

(12a)

\[ \nu \phi_1^2 + (1 - \nu) \phi_2^2 = \phi_2^0. \]  

(12b)

By rewriting equation (12) to express IDP volume fractions in \( \beta \) as functions of \( \nu \) with \( 0 < \nu < 1 \) and volume fractions in \( \alpha \):

\[ \phi_1^0 = \frac{\phi_1^0 - \nu \phi_1^0}{1 - \nu}, \]  

(13a)

\[ \phi_2^0 = \frac{\phi_2^0 - \nu \phi_2^0}{1 - \nu}. \]  

(13b)

\( f_{\text{sep}} \) is seen as a function of \( \phi_1^0, \phi_2^0, \) and \( \nu \),

\[ f_{\text{sep}}(\nu, \phi_1^0, \phi_2^0) = \nu f(\phi_1^0, \phi_2^0) + (1 - \nu)f\left(\frac{\phi_1^0 - \nu \phi_1^0}{1 - \nu}, \frac{\phi_2^0 - \nu \phi_2^0}{1 - \nu}\right). \]  

(14)

Note that \( \lim_{\nu \to 0} f_{\text{sep}}(\nu, \phi_1^0, \phi_2^0) = \lim_{\nu \to 1} f_{\text{sep}}(\nu, \phi_1^0, \phi_2^0) = f(\phi_1^2, \phi_2^2) \), where \( (\phi_1^0, \phi_2^0) \to (\phi_1^0, \phi_2^0) \) for \( \nu \to 0 \) and \( (\phi_1^0, \phi_2^0) \to (\phi_0^0, \phi_0^0) \) for \( \nu \to 1 \).

To find the set of variables that yields the global minimum of \( f_{\text{sep}} \), we numerically search the three-dimensional space of \( (\nu, \phi_1^0, \phi_2^0) \) by implementing the sequential least squares programming algorithm [92] using the `scipy.optimize.minimize` function in Scipy, a Python-based numerical package for scientific computation [93]. If a given \( (\nu, \phi_1^0, \phi_2^0) \) is found to yield a minimum of \( f_{\text{sep}} \) among computed \( f_{\text{sep}} \) values and also satisfies \( f_{\text{sep}}(\nu, \phi_1^0, \phi_2^0) < f_{\text{bulk}}(\phi_1^0, \phi_2^0) \), the system is judged to be in a state of binary phase separation to the two phases \( \alpha \) and \( \beta \). In contrast, if all \( f_{\text{sep}} \) for a given \( (\phi_1^0, \phi_2^0) \) are larger than \( f_{\text{bulk}} \), the bulk state \( (\phi_1^0, \phi_2^0) \) is thermodynamically stable and the system does not phase separate. Unlike in the two-component case in which the two separated phases are unique and independent of the bulk IDP concentration/volume fraction insofar as it is in the phase-separated regime, in three-component systems different bulk IDP concentrations/volume fractions can result in different \( \alpha \), \( \beta \) phases. Thus a complete binary phase diagram is generated by considering all possible \( (\phi_1^0, \phi_2^0) \) combinations.

For two-component systems (one IDP sequence plus solvent), we have shown that linear terms of \( \phi \) in the system free energy do not affect the determination of phase equilibrium [62]. In the same vein, here we demonstrate that the same principle applies also to three-component (two IDP sequences plus solvent) systems. As described above, binary coexistence is governed by the free energy difference

\[ \Delta f \equiv f_{\text{sep}} - f_{\text{bulk}} = \nu f(\phi_1^0, \phi_2^0) + (1 - \nu)f(\phi_1^2, \phi_2^2). \]  

(15)

Consider a modified free energy \( g(\phi_1, \phi_2) \) with an additional arbitrary linear function \( a_0 + a_1 \phi_1 + a_2 \phi_2 \) of \( \phi \)’s where \( a_0, a_1, \) and \( a_2 \) are constants:

\[ g(\phi_1, \phi_2) = f(\phi_1, \phi_2) + a_0 + a_1 \phi_1 + a_2 \phi_2. \]  

(16)

Now, the free energy difference between phase-separated and bulk phases becomes

\[ \Delta g \equiv g_{\text{sep}} - g_{\text{bulk}} = \Delta f + a_1[\nu \phi_1^0 + (1 - \nu) \phi_2^0 - \phi_1^0] + a_2[\nu \phi_1^2 + (1 - \nu) \phi_2^2 - \phi_2^0] \]  

(17)

because \( a_0 \) in \( g_{\text{sep}} \) and \( g_{\text{bulk}} \) cancel. The two bracketed terms in equation (17) are identically zero because of equation (12). Hence \( \Delta g = \Delta f \), meaning that any linear function of \( \phi \)’s added to \( f \), such as the one utilized in figure 2 for graphical clarity, has no impact on phase separation.

We apply the above-described minimization procedure for three-component (two IDP sequences plus solvent) systems to determine \( (\alpha, \beta) \) for selected pairs of sequences in table 1. To minimize possible numerical errors, every set of \( (\phi_1^0, \phi_2^0, \phi_1^1, \phi_2^1) \) obtained by minimizing \( f_{\text{sep}} \) is subject to further testing by comparing the chemical potentials in \( \alpha \) and \( \beta \). As described in equation (A.5) of [62], phase equilibrium implies the following equalities,

\[ f_1^{\alpha} = f_1^{\beta}, \]  

(18a)

\[ f_2^{\alpha} = f_2^{\beta}, \]  

(18b)

\[ \mu_\omega^{\alpha} = \mu_\omega^{\beta}. \]  

(18c)
and equation

where

$$f^y \equiv f (\phi_1^y, \phi_2^y),$$

$$f^x_y \equiv \frac{\partial f (\phi_x, \phi_y)}{\partial \phi_x} \bigg|_{\phi_x=\phi_x, \phi_y=\phi_y},$$

$$x = 1, 2; \ y = \alpha, \beta,$$

and

$$\mu^y_x \equiv f^y - \phi_1^y f^x_1 - \phi_2^y f^x_2$$

is the chemical potential of water [62]. Making use of the volume conservation conditions in equation (18) and substituting equation (13a) for $\phi_1^x$ and equation (13b) for $\phi_2^x$, equation (18) becomes three equalities for three variables $\phi_1^x, \phi_2^x$, and $v$. It follows that a unique determination of the phase-separated volume fractions $\phi_1^x, \phi_2^x$, $\phi_1^x$, and $\phi_2^x$ is afforded by equation (18).

It is straightforward to show that the set of phase-separated volume fractions $\{\phi_1^x, \phi_2^x, \phi_1^y, \phi_2^y\}$ determined by equation (18) are identical to that obtained by minimizing $f_{sep}$ in equation (14). A necessary condition for the minimization of $f_{sep}$ is that its Jacobian vector $J_{sep}$ of first-order partial derivatives of independent variables vanishes:

$$J_{sep}(v, \phi_1^x, \phi_2^x) \equiv \left( \begin{array} {c} \frac{\partial f_{sep}}{\partial \phi_1^1} \\ \frac{\partial f_{sep}}{\partial \phi_1^2} \\ \frac{\partial f_{sep}}{\partial \phi_2^1} \\ \frac{\partial f_{sep}}{\partial \phi_2^2} \\ \frac{\partial f_{sep}}{\partial v} \end{array} \right) = 0.$$  \hspace{1cm} (21)

In other words,

$$\frac{\partial f_{sep}}{\partial \phi_1^1} = v (f_1^\alpha - f_1^\beta) = 0,$$  \hspace{1cm} (22a)

$$\frac{\partial f_{sep}}{\partial \phi_2^1} = v (f_2^\alpha - f_2^\beta) = 0,$$  \hspace{1cm} (22b)

$$\frac{\partial f_{sep}}{\partial v} = f^\alpha - f^\beta + (\phi_2^\alpha - \phi_2^\beta) f_1^\beta + (\phi_1^\alpha - \phi_1^\beta) f_2^\beta = 0,$$  \hspace{1cm} (22c)

wherein we have utilized equation (13a) for $\phi_1^x$ and equation (13b) for $\phi_2^x$. Clearly, equations (22a) and (22b) are equivalent to equations (18a) and (18b), respectively, and equation (22c) is equivalent to equation (18c) by virtue of equation (20).

Q.E.D.

Starting with $(\alpha, \beta)$ obtained by minimizing $f_{sep}$ in equation (14), only those that deviate less than 0.1% from the chemical-potential-balancing equalities in equation (18) are accepted as valid binary pairs in our analysis. For the sequence pairs (seq1, seq5) and (seq1, seq6), a smaller threshold of 0.01% is used to ensure accuracy of the computed phase-separated $\phi$’s because for these sequence pairs the chemical potential balancing conditions are quite insensitive to variations of the $\phi$’s.

---

**Table 1.** Sequences studied are identified as seq1–7 in this article. They correspond to seven of the thirty 50-residue charged sequences with zero net charge in Das and Pappu [68]. The sequences’ sv labels, the values of their charge pattern parameter $\kappa$ and simulated single-chain radius of gyration $R_g$ are those in the same [68]. The charge pattern parameter SC is that of Sawle and Ghosh [69]. Values of SC and the RPA-predicted critical temperature $T_m^c$ are from Lin and Chan [63].

| Sequence | sv label | $-\text{SCD}$ | $\kappa$ | $R_g/$Å | $T_m^c/$K |
|----------|----------|----------------|---------|----------|------------|
| seq1     | sv28     | 15.99          | 0.7666  | 17.9     | 5.177      |
| seq2     | sv24     | 17.00          | 0.4456  | 17.6     | 5.160      |
| seq3     | sv25     | 12.77          | 0.5283  | 19.6     | 4.144      |
| seq4     | sv20     | 7.37           | 0.2721  | 19.6     | 2.275      |
| seq5     | sv15     | 4.35           | 0.1354  | 20.4     | 1.282      |
| seq6     | sv10     | 2.10           | 0.0834  | 25.5     | 0.611      |
| seq7     | sv1      | 0.41           | 0.0009  | 29.9     | 0.089      |

---
4.4. Binary coexistence of two charged sequences

Using RPA, we investigated previously how the phase separation behaviors of charged IDP sequences are affected by their charge patterns [61–63]. In particular, for the set of thirty KE sequences of Das and Pappu [68] with zero net charge but an equal number of 25 positively charged lysine (K) and 25 negatively charged aspartic acids (E) in different permutations, the critical temperature $T^*_e$ of phase separation was found [63] to be correlated with charge pattern parameters $\kappa$ [68] and

$$SCD \equiv \frac{1}{N} \sum_{i=1}^{N} \sum_{j=i+1}^{N} \sigma_i \sigma_j \sqrt{j - i},$$

where $i, j$ label the residues with charges $\sigma_i, \sigma_j$ along a chain of length $N$ [69]. The $\kappa$ and SCD parameters exhibit similar correlations with single-chain radius of gyration $R_g$ [68, 69]. The correlation of $T^*_e$ and $R_g$ with SCD is stronger than that with $\kappa$. A likely reason is that SCD accounts for non-local effects between charges far apart along the chain sequence whereas $\kappa$ does not [63]. Although we use only SCD in our analysis below, an equivalent analysis using $\kappa$ is expected to produce a similar trend.

How does the phase behavior of an IDP solution with two sequences depend on the sequences’ difference in charge patterns? Intuitively, when two sequences with different SCD values are present together, their different propensities to phase separate are expected to interfere. Indeed, such an effect of inter-sequence interference is seen clearly in the Taylor expansion of the integrand of the RPA expression $f_{\text{el}}$ for the electrostatic contribution to free energy in equation (3),

$$\ln[1 + \mathcal{G}(k)] - \mathcal{G}(k) = -\frac{1}{2} \mathcal{G}(k)^2 + \frac{1}{3} \mathcal{G}(k)^3 + ...$$

$$= -\frac{1}{2} \langle \sigma_i \mathcal{G}_{11}(k) \sigma_i \rangle^2 - \frac{1}{2} \langle \sigma_i \mathcal{G}_{12}(k) \sigma_j \rangle^2$$

$$- \langle \sigma_i \mathcal{G}_{12}(k) \sigma_i \rangle \langle \sigma_j \mathcal{G}_{12}(k) \sigma_j \rangle$$

$$+ O(\mathcal{G}(k)^4),$$

where $\mathcal{G}_{11}(k)$ and $\mathcal{G}_{12}(k)$ are the product of $4\pi/[k^3 \{1 + k^2 T^*\}]$ with, respectively, the $\mathcal{G}_{11}(k)$ and $\mathcal{G}_{12}(k)$ in equation (8). The first two terms after the second equality in equation (24) are self-interactions of the two sequences, identical to those in one-sequence RPA theory (see, e.g. equation (1) in [63]). The third term represents the interference effect in RPA. Since it is the product of square roots of the two self-interaction terms, its strength is intermediate between them, suggesting that phase behaviors of two-sequence systems are sensitive to the similarity/dissimilarity in charge pattern between the two sequences.

To investigate this sensitivity, we use the sequences in table 1 to compute the phase diagrams of six pairs of sequences, namely seq1 with each of the six other sequences. The pairs are selected to represent a broad range of similarity/dissimilarity in charge pattern as quantified by the difference in SCD values: from the (seq1, seq2) pair with SCD $= (-15.99, -17.00)$ to (seq1, seq7) with SCD $= (-15.99, -0.41)$. To compare the phase behaviors of the six sequence pairs on an equal footing, all phase diagrams in figure 3 are computed at the same reduced temperature $T^* = 4$. Noting that condensed-phase volume fractions tend to decrease with increasing $T^*$, this temperature is chosen because it falls in the mid-range of the broad span of $T^*$ for the sequences in table 1. $T^* = 4$ is much higher than the $T^* = 0.55$ equivalent of room temperature ($T = 300$ K) when an aqueous $\epsilon_t = 80$ is assumed [63]. This seemingly unphysical condition in our calculation has little impact, however, on the present goal of ascertaining general principles and behavioral trends. Although we do not aim for direct, detailed comparison with experiment here, $T^* = 4$ is experimentally relevant, for example, to IDPs with charge patterns similar to those considered here but with their electrostatic interaction strength significantly scaled down for various physical reasons such as screening or a more sparse charge distribution along the IDP sequence.

For each of the six phase diagrams in figure 3, the area surrounded by blue dots and ‘shaded’ by black dashed lines is the region of binary coexistence. In other words, bulk-state volume fractions falling within this region will phase separate into two coexisting liquid phases (as in figures 1(c)–(e)), whereas bulk-state volume fractions residing outside this region will be stable as a single liquid phase (as in figure 1(b)). Every black dashed line is the tie line connecting a pair of blue dots representing separated phases (a, b) for any bulk-state volume fractions lying on the given tie line (corresponding to the arrows in the schematic phase diagrams for figures 1(c)–(e)).

The average slope of the tie lines changes from positive for similarly patterned sequences (figures 3(a), (b)) to negative for very differently patterned sequences (figures 3(e), (f)). This trend may be understood as follows. When both sequences of a given pair can undergo phase separation individually ($T^*_e > 4$ for both), the tie lines near the $\phi_1$ and $\phi_2$ axes must be close to being parallel to the axes because when either $\phi_1 \to 0$ or $\phi_2 \to 0$, the two-sequence system reduces to the corresponding single-sequence system that phase separates. This situation applies to (seq1, seq2) and (seq1, seq3), resulting in positive tie-line slopes, indicating that the populations of the two sequences in each pair are well mixed even when they undergo phase separation. They prefer to stay together.
Figure 3. RPA-predicted binary phase diagrams of six charged sequence pairs at $T^\ast = 4$. In each panel, the pair of sequences considered and their charge pattern parameters ($-SCD$) are shown at the top, with positively and negatively charged residues depicted as red and blue circles, respectively. Further properties of the sequences are provided in table 1. Horizontal axes ($\phi_a$) refer to the volume fraction of seq1, vertical axes ($\phi_b$) are for the volume fraction of the other sequence in each of the pairs. The charge patterns of the sequence pairs vary from being very similar (a) to very dissimilar (f). This trend is quantified by the difference in $-SCD$ values between the two sequences in each of the six pairs. Each dashed line is a tie line connecting a pair of blue dots that represent coexisting phases $\alpha = (\phi_a, \phi_b)$ and $\beta = (\phi'_a, \phi'_b)$. All bulk volume fractions $(\phi_a, \phi_b)$ lying on one tie line undergo phase separation to the same $(\alpha, \beta)$; and, following equation (12), $\nu/(1 - \nu) = (\phi_a - \phi'_a)/\phi'_a - (\phi_b - \phi'_b)/\phi'_b$, which is equal to the length ratio of the two segments of the tie line from the bulk volume fractions $(\phi_a, \phi_b)$ on the phase diagram to the phase boundary marked by the blue dots. The black inclined solid line in (f), $\phi_1 + \phi_2 = 1$, delimits the region $\phi_1 + \phi_2 \leq 1$ within which IDP volume fractions may vary. Insets in (a) and (b) are zoom-in plots of a part of the phase diagram with extremely low IDP volume fractions. Inset in (e) and (f) are zoom-in plots of the gray-shaded regions of the respective phase diagrams. To provide a scale for comparison, the size of the gray-shaded regions $\phi_1, \phi_2 \in [0, 0.06]$ in (e) and (f) is chosen to be equal to the plotted regions for the other four sequence pairs.

after they phase separate, with similar population ratios for the two sequences in the ‘both-dilute’ (small $\phi'$)s as well as the ‘both-condensed’ (large $\phi'$)s phases (as in figure 1(c)). Because $T^\ast \chi < 4$ for the other four sequences (seq4–seq7), they do not phase separate by themselves individually and therefore tie lines near the $\phi_2$-axis need not be approximately parallel to it. Nonetheless, tie lines close to the $\phi_1$-axis are still required to essentially line up with the axis. For tie lines that possess large $\phi_2$ values, the volume conservation condition $\phi_1 + \phi_2 = 1$ enforces negative tie-line slopes. The combined effect of these constraints lead to tie-line slopes that gradually change from $\approx 0$ near the $\phi_1$-axis to $\approx -1$ near the $\phi_1 + \phi_2 = 1$ boundary, as exemplified by the case of (seq1, seq7) in figure 3(f). As shown in figures 3(d) and (e) for (seq1, seq5) and (seq1, seq6), this trend is apparent even when the phase-separated regime does not extend all the way to the $\phi_1 + \phi_2 = 1$ boundary. Negative tie-line slopes imply various degrees of demixing of the populations of the two sequences: the phase-separated state ($\alpha, \beta$) now comprises one $\phi_1$-enriched ($\phi_a' > \phi_b'$) phase coexisting with one $\phi_2$-enriched ($\phi_a' < \phi'_b$) phase. The degree of population demixing depends on how dissimilar are the charge patterns of the two sequences in the pair. For large differences in SCD as in figure 3(f), one of the coexisting phases can have a very low population of seq1 but a substantial seq7 volume fraction, whereas the other phase has a relatively low population of seq7 but a substantial seq1 volume fraction (as in figure 1(e)).

The (seq1, seq4) pair in figure 3(c) is at the crossover between the well-mixed and demixed extremes. Tie-line slopes in this case are all $\approx 0$, indicating that although increasing seq4 volume fraction decreases the phase separation tendency of seq1, even in the phase-separated regime the concentration of seq4 is essentially identical in the two coexisting phases, i.e., $\phi_a'$ $\approx$ $\phi_a''$ $\approx$ $\phi_b''$ (as in figure 1(d)).
The difference in the ratio of component populations in coexisting phases \((\alpha, \beta)\) may be quantified by comparing the volume ratio of the two sequences in the two phases. We first consider a rather intuitive measure

\[
\Delta_{\alpha,\beta}(\phi_1/\phi_2) \equiv \left\langle \left( \frac{\phi_1^{\alpha}}{\phi_2^{\alpha}} - \frac{\phi_1^{\beta}}{\phi_2^{\beta}} \right) \right\rangle
\]  

(25)

of compositional asymmetry between coexisting phases, where the absolute value ensures that \(\Delta_{\alpha,\beta}(\phi_1/\phi_2)\) is \(\alpha \leftrightarrow \beta\) symmetric, and the bracket \(\langle \rangle\) denotes averaging over all \((\alpha, \beta)\) pairs of coexisting phases. One disadvantage of this measure, however, is that the average is strongly dominated by those pairs of \((\alpha, \beta)\) with large \(\phi_1\) but small \(\phi_2\), i.e., coexisting pairs that are close to \(\phi_1\)-axis in figure 3. Therefore, we also consider another composition asymmetry measure

\[
A_{\alpha,\beta} \equiv \left\langle \frac{2}{\pi} \tan^{-1}\left( \frac{\phi_1^{\alpha}}{\phi_2^{\alpha}} \right) - \tan^{-1}\left( \frac{\phi_1^{\beta}}{\phi_2^{\beta}} \right) \right\rangle
\]  

(26)

that avoids this potentially problematic feature by replacing the ratio \(\phi_1/\phi_2\) with its arctangent value normalized by \(\pi/2\) such that \(0 \leq A_{\alpha,\beta} \leq 1\).

Summarizing our findings using two-sequence RPA theory, figure 4 shows the variation of \(\Delta_{\alpha,\beta}(\phi_1/\phi_2)\) as well as \(A_{\alpha,\beta}\) with the difference in SCD values of the six sequence pairs in figure 3. A reasonable correlation is seen for both composition asymmetry measures, with the \(A_{\alpha,\beta}\) measure exhibiting a better correlation by varying monotonically with SCD difference, indicating that compositional asymmetry or degree of demixing of phase-separated populations as quantified by \(A_{\alpha,\beta}\) is positively correlated with the difference in charge patterns as quantified by difference in SCD values. This plot illustrates graphically how a stochastic, multivalent form of molecular recognition that arises from the interactions among the diverse conformations in a multiple-chain ensemble can lead to demixing of different IDP species into different coexisting phases.

5. Comparison with phase separation in FH models

We next seek a deeper understanding of the RPA results and their ramifications by comparing them with the predictions of a variety of FH models. As emphasized, unlike RPA, FH by itself does not address the physics of sequence dependence [20, 62]. Accordingly, FH \(\chi\) interaction parameters for IDP sequences have to be provided phenomenologically by experiment or theoretically by microscopic physical theory. For example, as will be discussed further below, an intuitive and semi-quantitative connection between RPA and FH is provided by the expansion in equation (24). It should also be noted that FH Neglects interaction terms that are higher than

\[\chi^\text{eff} = |2| I \text{NaCl} = 4.3 \text{ Å} \text{ instead of the correct value of 3.04 Å. To facilitate comparison with [20], however, the effective Debye length in figure 8 of [62] was also set to 4.3 Å.} \]
quadratic order in IDP volume fractions/concentrations ($\phi$’s) such as the $O(G(k)^2)$ terms in equation (24) because $G(k) \propto \phi$ (equations (9a) and (9b)). This approximation can be problematic when IDP concentrations are high. Nonetheless, by treating the three parameters $\chi_{11}$, $\chi_{22}$, and $\chi_{12}$ in the three-component FH interaction term for two IDP species plus solvent

$$f_{\text{FH}} = -(\chi_{11}\phi_1^2 + \chi_{22}\phi_2^2 + 2\chi_{12}\phi_1\phi_2)$$

as free (arbitrary) variables, we can either match FH behavior to that of RPA to gain conceptual insights or explore other interaction scenarios that might be physically plausible when interactions other than the rudimentary electrostatics embodied in RPA are included in the physical picture.

5.1. FH models that imitate RPA theory by having two independent $\chi$’s

We begin this analysis by first constructing FH models with interaction schemes similar to RPA, then tuning the interaction parameters to produce phase behaviors similar to those predicted by RPA in figure 3. If we identify the $\int dk \ k^2/4\pi^2$ integral (equation (3)) of the order $G(k)^2$ terms in the RPA expansion equation (24) with the FH interaction term in equation (27), we may define $\tilde{\chi}_{11}(k)$ and $\tilde{\chi}_{22}(k)\equiv 2\{\langle |\tilde{G}_{11}(k)|^2 \rangle [k(1+k^2)T^*]^2\}$ and $\tilde{\chi}_{12}(k)\phi_2^2 \equiv 2\{\langle |\tilde{G}_{12}(k)|^2 \rangle [k(1+k^2)T^*]^2\}$ such that $\chi_{11} = \int dk \ \tilde{\chi}_{11}(k)$, $\chi_{22} = \int dk \ \tilde{\chi}_{22}(k)$, and $\chi_{12} = \int dk \ \sqrt{\tilde{\chi}_{11}(k)}\sqrt{\tilde{\chi}_{22}(k)}$, from which it is clear that only two set of interaction parameters $\tilde{\chi}_{11}(k)$ and $\tilde{\chi}_{22}(k)$ as functions of $k$ are independent. Here we approximate this dependence by constraining $\chi_{12} = \sqrt{\chi_{11}\chi_{22}}$.

We then construct three FH systems that have $\chi_{11} = \chi_{22}$, $\chi_{11} \gg \chi_{22}$, and $\chi_{12} \gg \chi_{22}$, corresponding respectively to sequence pairs with small, intermediate, and large charge pattern (SCP) differences. The phase diagrams of these models (figure 5) exhibit a trend similar to that seen in the RPA-predicted figure 3. Specifically, figure 5(a) is similar to figure 3(a), figure 5(b) to figure 3(c), and figure 5(c) to figure 3(f). This correspondence offers conceptual clarity because the degree to which the interaction between the two IDP species is favorable is explicit in FH. When $\chi_{11} = \chi_{22} = \chi_{12}$, the two species are miscible and their phase separation propensities are identical, resulting in the coexistence of one both-dilute phase and one both-condensed phase. The similarity between figures 5(a) and 3(a) indicates that this behavior can be achieved physically by two IDP species with similar charge patterns. In contrast, when $\chi_{11} \gg \chi_{12} \gg \chi_{22}$, miscibility of the two species is poor and their phase separation propensities are quite different, resulting in a high degree of population demixing. The similarity of this behavior shown in figure 5(c) with that in figure 3(f) underscores once again that charge-pattern mismatches between IDPs can lead to substantial weakening of attractive interactions.

5.2. FH models with three independent $\chi$’s

We next consider the general case in which the three $\chi$’s in equation (27) are independent. Although this modeling setup does not have a simple correspondence with IDP sequences interacting via physical forces like that described above, the expanded variety of scenarios explored here would be valuable when behaviors more complex than those allowed by our current RPA formulation are considered in more comprehensive and detailed physical theories. As simple examples of the rich possibilities, here we focus on FH models with $\chi_{11} = \chi_{22}$ (≡ $\chi$) and variable $\chi_{12}$ values that are not related to $\chi$. Figure 6 shows the phase diagrams of three representative models with (a) $\chi_{12} \gg \chi$, (b) $\chi_{12} \ll \chi$, and (c) $\chi_{12} < \chi$. Compared to the $\chi \equiv \chi_{11} = \chi_{22} = \chi_{12}$ case in figure 3(a), it is clear that a stronger inter-component attraction (larger $\chi_{12}$) makes the phase-separated

![Figure 5](image-url)
region bulge (figure 6(a)), whereas a weaker inter-component attraction (smaller $\chi_{12}$) shrinks it (figure 6(b)). Nonetheless, the tie-line slopes are positive in both situations, indicating that the two components are largely miscible.

Figure 6(b) indicates that a weakened $\chi_{12}$ shrinks the phase-separated region most around $\phi_1 = \phi_2$. As $\chi_{12}$ decreases further, inter-component attraction all but vanishes, imibility disappears, resulting in the phase-separated region being broken into two parts, one for $\phi_1 \gg \phi_2$ and the other for $\phi_1 \ll \phi_2$ (figure 6(c), shaded area and inset). The tie lines in these two regions are almost parallel to either the $\phi_1$- or the $\phi_2$-axis, implying that one component is dominant while the concentration of the other component barely changes upon phase separation. With such an effective inter-component repulsion (i.e., less favorable inter-component attraction vis-à-vis the strengths of intra-component cohesion), an additional phase-separated regime of poor miscibility similar to that in figures 3(f) and 5(c) is induced, wherein all tie-line slopes are negative, signaling substantial demixing (figure 6(c), region close to $\phi_1 + \phi_2 = 1$ with tie-line slopes = $-1$).

5.3. Ternary coexistence in FH model
If $\chi_{12}$ is made even weaker than that in figure 6(c), the region of poor miscibility below the $\phi_1 + \phi_2 = 1$ boundary would grow. Finally, the three phase-separated regions intersect and a new ternary coexistence region emerges in-between.

According to the Gibbs phase rule, an $n$-component system can separate into at most $n$ coexisting phases, when all other environmental conditions, e.g. temperature and pressure, are kept constant [62, 89]. In our system of $n = 3$ (two sequences plus solvent), whether the system will separate to two or three coexisting phases may be deduced by observing the variation of tie-line slopes in putative regions of binary coexistence: if the tie lines have to become parallel to the $\phi_1$-axis, $\phi_2$-axis, or the $\phi_1 + \phi_2 = 1$ line when they approach these boundaries respectively, the tie-line slopes have to be able to vary smoothly to satisfy these constraints in order for binary coexistence to be stable. In that case, ternary coexistence is unlikely. Conversely, if there are conflicts that prevent a smooth change of tie-line slope, a ternary coexistence region ensues.

An example is provided by using figure 6(c) as starting point. Here the three phase-separated regions are close to the three boundaries, and their tie lines are essentially parallel to the respective boundaries. Under this circumstance, when effective inter-component repulsion is enhanced by weakening $\chi_{12}$ to cause the three regions to evolve toward merging, the conflict among the three different trends of tie-line slopes necessitates reconciliation by a region of ternary phase separation (figure 7).

In order to determine the three phases in ternary coexistence mathematically, we extend the phase-separated free energy expression in equation (11b) for phases ($\alpha$, $\beta$) to including one additional phase $\gamma$, viz,

$$f_{\text{ternary}} = \nu_\alpha f(\phi_1^\alpha, \phi_2^\alpha) + \nu_\beta f(\phi_1^\beta, \phi_2^\beta) + (1 - \nu_\alpha - \nu_\beta)f(\phi_1^\gamma, \phi_2^\gamma),$$

where the fractional volumes $\nu_\alpha$, $\nu_\beta$ are functions of the six $\phi$’s by virtue of volume conservation,

$$\sum_y \nu_y \phi_x^y + \left(1 - \sum_y \nu_y\right) \phi_x^\gamma = \phi_x^\alpha,$$

where $x = 1, 2$ and $\sum_y$ is over $y = \alpha, \beta$. Now the equalities in equation (18) have to include the addition phase to become
and

\[ \sum_{i=1}^{3} \phi_i = 1 \]

by the vertices of the triangle (as in figure 1(b)). Outside of the triangle there are three binary coexistence regions that converge to the \( \phi_x \) axis, \( \phi_y \) axis, and the \( \phi_x + \phi_y = 1 \) boundary, respectively, wherein dashed tie lines connect pairs of coexisting phases as in figures 3, 5, and 6. Bulk states within the bottom-left small white region near the \( \phi_x, \phi_y \) origin does not phase separate. The green dashed lines connect three examples of bulk state \( O_1, O_2, O_3 \) to the same ternary phases \( \alpha, \beta, \gamma \) because the compositions of these phases are independent of bulk-state volume fractions. For any given bulk state \( O_1, O_2, O_3 \) and \( v_x \) are given, respectively, by the areas of the triangles \( O_1 O_2 O_3, O_1 O_3 O_2 \) (all have one solid blue and two green dashed sides) as a fraction of the total triangular area of \( \alpha, \beta, \gamma \) bound by three solid blue lines (see text for details). (b) Free energy landscape of \( \Delta f (\phi_x, \phi_y) \equiv f_{\text{ternary}} (\phi_x, \phi_y) = [a_0 + a_1 \phi_x + a_2 \phi_y] \), where the bracketed function linear in the \( \phi_i \)'s specifies the plane defined by \( \alpha, \beta, \gamma \). In other words, the \( a_i \)'s are determined by solving \( f_{\text{ternary}} (\phi_x, \phi_y, \phi_z) = a_0 + a_1 \phi_x + a_2 \phi_y + a_3 \phi_z \) for \( z = \alpha, \beta, \gamma \). Note that \( \Delta f > 0 \) for all \( (\alpha, \beta, \gamma) \) in the plotted region, indicating that phase separation is preferred if the volume conservation condition for either binary equation (12) or ternary equation (29) can be satisfied. The logarithmic color scale on the right is for both (b) and (c). The turquoise lines in (b) and (c) mark the same ternary phase boundaries as those in (a). (c) Contour plot of \( \Delta f \). The three ternary coexistence phases (\( \Delta f = 0 \)) are seen to be situated in three different basins with small \( \Delta f \).

\[ f_{\text{ternary}} (\alpha, \beta, \gamma) = f_1^\alpha + f_2^\beta + f_3^\gamma, \]

(30a)

\[ f_{\text{ternary}} (\alpha, \beta, \gamma) = f_1^\alpha + f_2^\beta + f_3^\gamma, \]

(30b)

\[ \mu_{w}^\alpha = \mu_{w}^\beta = \mu_{w}^\gamma. \]

(30c)

Because here we have six equalities in equation (30) for six phase-separated \( \phi_i \)'s, the solution for ternary coexistence of \( (\alpha, \beta, \gamma) \) is unique irrespective of the bulk-state volume fractions insofar as they fall within the ternary coexistence region. This situation is different from that of binary coexistence in which the separated phases \( (\alpha, \beta) \) can be different for different bulk-state \( (\phi_1^0, \phi_2^0) \)'s when they are on different tie lines.

Similar to the binary coexistence case in section 4.3, we proceed to demonstrate that minimizing equation (28) is equivalent to solving the equations in equation (30). Using essentially the same approach, we rewrite \( f_{\text{ternary}} \) as a function of six independent variables: \( v_x, v_y, \phi_x^1, \phi_x^2, \phi_y^1, \phi_y^2 \) by first utilizing equation (29) to express \( \phi_x^1 \) and \( \phi_x^2 \) as

\[ \phi_x = \frac{\phi_x - \sum_y v_y \phi_y^x}{1 - \sum_y v_y}, \]

(31)

where \( x, y \) and \( \sum_y \) have the same meanings as above. In this notation, equation (28) becomes

\[ f_{\text{ternary}} (v_x, v_y, \phi_x^1, \phi_x^2, \phi_y^1, \phi_y^2) = \sum_y v_y f^y + \left( 1 - \sum_y v_y \right) f^x. \]

(32)

Similarly to equation (21) for binary phase separation, we calculate the six derivatives of \( f_{\text{ternary}} \) and set them to zero as in equation (22) as necessary conditions for the minimization of \( f_{\text{ternary}} \) resulting in

\[ \frac{\partial f_{\text{ternary}}}{\partial \phi_x^y} = v_y f_x^y - \left( 1 - \sum_y v_y \right) f_x^y, \]

(33a)

\[ \frac{\partial f_{\text{ternary}}}{\partial v_y} = f^y - f^x + \left( 1 - \sum_y v_y \right) \left[ \sum_x f_x^y \cdot \left( 1 - \sum_y v_y \phi_y^x \right) \right] \]

\[ = f^y - f^x + \sum_x f_x^y \phi_y^x - \sum_x f_x^y \phi_x^y, \]

(33b)

where \( \sum_y \) sums over \( y' = \alpha, \beta \). Substituting \( x = 1, 2 \) in equation (33a) yields equations (30a) and (30b), whereas substituting \( y = \alpha, \beta \) in equation (33b) yields equation (30c).

Q.E.D.
Figure 7 provides an FH phase diagram with both binary and ternary coexistence. In this example, $\chi_{12}$ is significantly smaller than $\chi_{11}$, resulting in strong effective repulsion between the two sequences. Consequently, the two islands of binary coexistence around the $\phi_1$- and $\phi_2$-axes intersect with the top-right binary region with $(\phi_1^a, \phi_2^a) = (\phi_1^b, \phi_2^b)$, resulting in a ternary phase separation region corresponding to the $\alpha\beta\gamma$ triangle and its interior (marked by blue lines in figure 7(a) and turquoise lines in figures 7(b) and (c)). The thermodynamic stability of the ternary phase-separated state within this region is illustrated by the $\Delta f$ quantity plotted in figures 7(b) and (c); $\Delta f(\phi_1, \phi_2)$ is the bulk (not-phase-separated) free energy $f_{\text{bulk}}$ minus the free energy value for the same $\phi_1$, $\phi_2$ on a plane defined by the ternary phases (i.e., $\Delta f = 0$ for the three points corresponding to the $\alpha$, $\beta$, $\gamma$ phases). Because $\Delta f > 0$ for any other point within the triangular region, the ternary state is more stable than the bulk state in this region. Moreover, the free energy $f_{\text{exp}}$ of any putative binary coexistence state of a bulk state within the triangular region must lie on a tie line joining two points on the landscape in figures 7(b) and (c). Because $\Delta f > 0$ for any point other than the three ternary phases in the entire plotted region—including points outside the triangular region, $\Delta f > 0$ holds also for any putative binary coexistence state for the bulk state within the triangular region, implying that they are less stable than the ternary phase-separated state in the region.

For any given bulk-state $(\phi_1^0, \phi_2^0)$ in the ternary region, fractional volumes $v_{00}, v_{01}$ and $v_{11}$ in the respective coexisting phases $\alpha$, $\beta$, and $\gamma$ are determined by solving equation (29) and setting $v_0 = 1 - v_{00} - v_{01}$; In terms of the three-dimensional vectors $\Phi_0 \equiv (\phi_1^0, \phi_2^0, 0)$, $\Phi_1 \equiv (\phi_1^1, \phi_2^1, 0)$, $\Phi_{12} \equiv \Phi_1 - \Phi_0$, and $\Phi_{13} \equiv \Phi_2 - \Phi_1$, where $z_1, z_1, z_2 = \alpha, \beta, \gamma$; $v_0 = |\Phi_{00} \times \Phi_{01}|/|\Phi_{10} \times \Phi_{11}|$, $v_{01} = |\Phi_{00} \times \Phi_{01}|/|\Phi_{11} \times \Phi_{12}|$, and $v_{11} = |\Phi_{01} \times \Phi_{12}|/|\Phi_{10} \times \Phi_{11}|$. Because the area of a triangle defined by two vectors is equal to half of the magnitude of their cross product, these fractional volumes correspond to specific ratios of triangular areas as described in the caption for figure 7.

6. Discussion

6.1. Insights into cellular binary and ternary IDP phase coexistence

The present theoretical development bears on the sequence dependence of multicomponent phase separation in the cell. However, because the cellular processes involve many species of biomolecules and are extremely complex [27, 71], development of treatments more elaborated than our simple theories will be needed for quantitative comparison with experiments. Nonetheless, it is instructive to explore whether our RPA and FH results are qualitatively consistent with what has been observed experimentally.

Of interest are fibrillarin FIB1 (323 residues) and nucleophosmin NPM1 (299 residues) from frog (Xenopus laevis) oocytes. These IDPs tend to demix, exhibiting phase behaviors that likely underpin the assembly of nucleolar subcompartments [27]. Treating histidine sidechains at pH $\geq 7$ as neutral, the net charge of FIB1 is 19 and of NPM1 is $-22$. Their charge patterns, as quantified by SCD (equation (23)) = 4.126 and $-0.119$, respectively, are substantially different. Thus, the tendency for FIB1 and NPM1 to demix is qualitatively in line with the RPA-predicted trend in figures 3 and 4.

Aqueous solutions with both FIB1 and NPM1 undergo both binary and ternary liquid–liquid phase separations. In this respect, their experimental phase diagram in figure 4(D) of Feric et al [27] is similar to our FH phase diagram in figure 7. The two regions of binary coexistence of one condensed (around $\alpha$ or around $\beta$) and one dilute (around $\gamma$) phases in figure 7 correspond to their ‘FIB1 rich/NPM1 lean’ and the ‘NPM1 rich/FIB1 lean’ areas, whereas the ternary coexistence region in figure 7 with two condensed ($\alpha$, $\beta$) and one dilute ($\gamma$) phases (as in figure 1(f)) corresponds to their ‘3 Phase’ area [27, 32].

It is noteworthy that our attempts to seek numerical solutions to ternary coexistence in the RPA models studied in figure 3 by minimizing equation (28) either ended in failure or resulted in solutions with two (among three) phases essentially identical and thus reduces the solution to that of binary coexistence. Apparently, ternary coexistence requires an effective intercomponent repulsion that is substantially stronger, as in figure 7, than that posited by RPA. The reason is that RPA constrains the intercomponent interaction strength $\chi_{12}$ to approximately the geometric mean of the two intracomponent interaction strengths $\chi_{11}$ and $\chi_{22}$ (section 5.1) and therefore $\chi_{12} \ll \chi_{11}$, $\chi_{22}$, is highly unlikely if not impossible.

This consideration suggests that difference in charge pattern alone may be insufficient to account for the rather strong effective repulsion between FIB1 and NPM1, although the impact of them being not very close to neutral remains to be investigated. (Unlike the model KE sequences in figure 3 with zero net charge or Ddx4 with a charge ratio = (net charge/chain length) = $-1.7\%$ [61], their charged ratios are, respectively, +5.9% and $-7.4\%$). In addition to the electrostatic interactions among FIB1 and NPM1, other driving forces surely also contribute to their phase behaviors. For example, the presence of ribosomal RNA (rRNA) appears to be important; and the presence of rRNA and inter-phase surface tensions have been modeled in lattice simulations to
6.2. Cooperativity driven by concentration-dependent relative permittivity

Most analytical formulations for charged polymer solutions, including common RPA theories, treat the relative permittivity \( \varepsilon_r \) of the solution as a constant independent of polymer concentration. However, as we noted recently [62], because of the significantly different permittivities of water \( (\varepsilon_w \approx 80) \) and protein \( (\varepsilon_p \approx 2-4) \) [94], the effective \( \varepsilon_r \) of a protein solution can change dramatically with protein concentration. Indeed, protein-dependent variations of dielectric properties of the aqueous medium have been shown to be relevant to globular protein stability in thermophilic species [95–98]. Because of the anticipated importance of dielectric properties to IDP energetics such as enabling a greatly enhanced propensity to phase separate [62], here we expand our theoretical exploration to two-IDP systems and consider in more detail the physical basis of various effective medium approximations that may be applied to estimate \( \varepsilon_r \) of IDP solutions. Since the scope of this exploration is limited to establishing certain general principles, for simplicity of the discussion we let \( \varepsilon_p \) be the relative permittivity of both IDP species in the system such that \( \varepsilon_r(\phi_1, \phi_2) = \varepsilon_r(\phi) \) where \( \phi = \phi_1 + \phi_2 \).

Previously we considered two models for \( \varepsilon_r \) [62], namely the ‘slab model’ derived by considering a planar electric capacitor, wherein

\[
\varepsilon_{r}^{\text{Slab}}(\phi) = \frac{\varepsilon_p \varepsilon_w}{\phi \varepsilon_w + (1 - \phi) \varepsilon_p},
\]

which corresponds to equation (3) of Bragg and Pippard when the depolarizing coefficient \( L = 0 \) [99], and the Clausius–Mossotti (CM) model predicting

\[
\varepsilon_{r}^{\text{CM}}(\phi) = \frac{1 + 2[(1 - \phi) \gamma_w + \phi \gamma_p]}{1 - [(1 - \phi) \gamma_w + \phi \gamma_p]},
\]

where

\[
\gamma_w \equiv \frac{\varepsilon_w - 1}{\varepsilon_w + 2}, \quad \gamma_p \equiv \frac{\varepsilon_p - 1}{\varepsilon_p + 2}
\]

are proportional to the CM expression for molecular polarizability [100]. The \( \phi \)-dependences of \( \varepsilon_{r}^{\text{Slab}}(\phi) \) and \( \varepsilon_{r}^{\text{CM}}(\phi) \) are very similar (figure 11 of [62]). Both models recognize amino acid residues and water as materials with excluded volume such that the contributions of their molecular polarizabilities to the overall dielectric property of the medium are against a vacuum background.

An alternate approach to effective medium is the Maxwell Garnett (MG) model [84] that pictures the effective dielectric property of a composite material as arising from embedding a component (IDP in our case) into an all-permeating background medium (water in our case). IDP excluded volume is not taken into account in this approach. Applying this method to our IDP solution yields

\[
\varepsilon_{r}^{\text{MG}}(\phi) = \frac{\varepsilon_w + 2 \gamma_{\text{MG}}}{1 - \phi \gamma_{\text{MG}}},
\]

where

\[
\gamma_{\text{MG}} \equiv \frac{\varepsilon_p - \varepsilon_w}{\varepsilon_p + 2 \varepsilon_w}.
\]

Comparing the \( \gamma_p \) expression in equation (36) with equation (38) indicates that \( \gamma_{\text{MG}} \) corresponds—up to a constant—to an effective molecular polarizability of IDP material, not against vacuum but rather in a water background \( \gamma_p \) is mathematically equal to \( \gamma_{\text{MG}} \) when \( \varepsilon_w \to 1 \), thus MG is related to CM in this respect [101]. Another approach known as the Bruggeman (BG) model [84] is an \( \varepsilon_p \to \varepsilon_w \) symmetrized form of MG. The BG \( \phi \)-dependent permittivity is given by

\[
\varepsilon_{r}^{\text{BG}}(\phi) = \frac{b_{\text{BG}}(\phi) + \sqrt{8 \varepsilon_w \varepsilon_p + b_{\text{BG}}(\phi)^2}}{4},
\]

where

\[
b_{\text{BG}}(\phi) = [2\phi - (1 - \phi)] \varepsilon_p + [2(1 - \phi) - \phi] \varepsilon_w.
\]

A graphical illustration of the different physical pictures assumed by the slab/CM versus the MG/BG approaches is provided by figure 8. Leaving aside questions about the appropriateness of their respective physical pictures for the moment, we first examine the properties of these \( \varepsilon_r(\phi) \)’s and their implications for IDP phase separation.
Variations of several properties of the $\epsilon_i(\phi)$'s are shown in figure 9. Although all four models give $1/\epsilon_i(\phi) = 1/\epsilon_w$ and $1/\epsilon_i(\phi = 1) = 1/\epsilon_p$ as they should, the $\phi$-dependences of the slab/CM models are very different from that of the MG/BG models. Here we are more interested in $1/\epsilon_i$ than $\epsilon_i$ itself because $1/\epsilon_i$ is directly proportional to Coulomb energy. Figure 9(a) shows that $1/\epsilon_i$ of the slab model is linear in $\phi$, that of CM is approximately linear; but the $1/\epsilon_i$'s of MG and BG increase very little when $\phi$ is small and exhibit rapid increases toward the $1/\epsilon_i$ value for $\phi = 1$ only for $\phi \gtrsim 0.8$ and 0.6, respectively. The linear and near-linear $\phi$-dependences of the $1/\epsilon_i$'s for the slab and CM models are underscored by their small first and second derivatives in $\phi$, whereas the sharp rises of $1/\epsilon_i$ near $\phi \approx 1$ for the MG and BG models are illustrated by their large $\phi$-derivatives for $\phi \gtrsim 0.7$ (figures 9(b) and (c)).

As described in our previous work [62], when permittivity becomes a function of IDP concentration, the last subtraction of $\mathcal{G}(k)$ in equation (3) has to be modified because $\mathcal{G}(k)$ is no longer linear in $\phi$’s [62]. A straightforward generalization of equations (68) and (69) of [62] to the present case with two IDP sequences (but now with neither salt nor counterions) leads to the following replacement for the RPA expression in equation (3) to accommodate a $\phi$-dependent $\epsilon_i$:

$$f_\Omega(\phi_1, \phi_2) = \int_0^\infty \frac{d \mathbf{k}^2}{4 \pi^2} \left[ \ln \left[ 1 + \mathcal{G}_1(k) \right] - \mathcal{G}_2(k) \right],$$

(41)
new J. Phys., 19 (2017) 115003

Y-H Lin et al

Figure 10. RPA-predicted spinodal instability of a two-sequence aqueous solution with an IDP-concentration-dependent or an IDP-concentration-dependent relative permittivity. Results in (a)–(d) are for the (seq1, seq2) system at $T_0^* = 0.05$ computed using the four different models of $\epsilon_\ell(\phi)$ described in the text, with $\epsilon_w = 80$, $\epsilon_0 = 4$, and $\phi = \phi_1 + \phi_2$. Spinodal regions are shaded and bounded by solid blue lines. The inclined solid black line $\phi_1 + \phi_2 = 1$ is the volume-conservation boundary. (a) Slab model, exhibiting only a single spinodal region; (b) Clausius–Mossotti model, also showing a single spinodal region; (c) Maxwell–Garnett model, two spinodal regions are observed; note that the upper spinodal boundary is very close to but not identical with $\phi_1 + \phi_2 = 1$. (d) Bruggeman model, showing also two spinodal regions. (e) Comparing the dilute spinodal boundary in the slab model (blue solid line) against the spinodal boundaries for RPA models with $\phi$-independent permittivity $\epsilon_\ell = 40$, 60, and 80 (dashed lines) at the same temperature $T_0^* = 0.05$. Similar to (e) but here we consider the MG model instead of the slab model. In (e) and (f), all dash lines higher than the blue solid line are condensed-phase boundaries for the constant-$\epsilon_\ell$ models. Insets in (a)–(f) are zoom-in’s that provide a closer look at the dilute spinodal phase boundaries.

where

$$G_1(k) = \frac{4\pi}{k^2(1 + k^2)T_0^* \epsilon_\ell(\phi)} \left[ \langle \sigma_1 \hat{G}_{11}(k) | \sigma_1 \rangle + \langle \sigma_2 | \hat{G}_{22}(k) | \sigma_2 \rangle \right],$$

$$G_2(k) = \frac{4\pi}{k^2(1 + k^2)T_0^* \epsilon_\ell(\phi)} \left[ \frac{1}{N_1} \sum_i \sigma_1^{(i)} | + \frac{\phi_e}{N_2} \sum_i \sigma_2^{(i)} | \right],$$

and, following equation (67) of [62],

$$T_0^* \equiv 4\pi \epsilon_0 k_0 T/\epsilon^2 = T^*/\epsilon_\ell.$$  

The resulting spinodal phase diagrams predicted by the four $\epsilon_\ell(\phi)$ models are shown in figure 10 for the (seq1, seq2) pair at $T_0^* = 0.05$. This temperature is chosen to facilitate comparison with constant-$\epsilon_\ell$ results because $T_0^* = 0.05$ corresponds to $T^* = 4$ when $\epsilon_w = \epsilon_\ell = 80$.

Figures 10(a)–(d) shows that all four $\epsilon_\ell(\phi)$ models have large spinodal regions extending to $\phi_1 + \phi_2 \approx 0.8$. These spinodal regions are much larger than those predicted by constant-$\epsilon_\ell$ theories (figures 10(e), (f)), pointing once again to a significant cooperative effect arising from a decrease in permittivity upon IDP condensation, which, in turn, increases electrostatic attraction and hence leads to more IDP condensation [62]. For instance, the condensed-phase volume fractions of the slab and CM models $\approx 0.8$, which represents a $\sim 20$-fold increase from the condensed-phase volume fraction of $\approx 0.03$ for a constant $\epsilon_\ell = \epsilon_w = 80$ (figure 10(e)). The corresponding enhancement in the MG and BG models is even more prominent—their condensed-phase volume fraction almost reaching the $\phi_1 + \phi_2 = 1$ limit (figures 10(c), (d))—because of the sharp rises of their $\epsilon_\ell(\phi)$’s when $\phi \rightarrow 1$ (figure 9). Although the precise quantitative impact of $\epsilon_\ell(\phi)$ remains to be ascertainable experimentally, our theoretical results suggest strongly that $\phi$-dependent relative permittivity should play a significant role in IDP phase separation, and that such a physical cooperative effect may help rectify some of the RPA-predicted condensed-phase volume fractions (e.g. those in figures 3(a)–(d)) that seem unrealistically low.
Interestingly, while the slab and CM models enlarge a single spinodal region vis-à-vis that for constant-$\epsilon$, the MG and BG models produce an additional spinodal region close to the $\phi_1$-$\phi_2$ origin. This region is similar in scope to the constant-$\epsilon$ spinodal region, and is well separated from the MG and BG models’ lower boundaries at $\phi_1 + \phi_2 \approx 0.6$ (MG) or 0.4 (BG) for their main (much larger) spinodal regions (figures 10(c)–(l)). This feature likely arises from the fact that $1/\epsilon_s$’s for MG and BG barely change for $\phi \lesssim 0.2$ (figure 9(a)), thus the behaviors of these models at small $\phi$’s should be similar to those of constant-$\epsilon$, models. For larger $\phi$’s, however, because of the rapid increase of their $1/\epsilon_s$’s with $\phi$, the MG and BG models become similar to the slab and CM models.

Although the slab/Cs and MG/BG models produce similarly expanded spinodal regions for the example in figure 10, the difference in their predicted $\epsilon_s(\phi)$’s do have important implications on the energetics of IDP phase separation. For example, the condensed-phase volume fraction of Ddx4 was recently determined to be $\sim 15$–0.2 [86]. At $\phi \approx 0.2$, the slab/Cs models posit a significant cooperative effect in favor of phase separation, but the MG/BG models suggest that such cooperativity is negligible unless $\phi \gtrsim 0.6$ (figure 9). Issues related to effective medium approximations can be intricate, as witnessed by the extensive materials-science literature on the topic [84, 101]. Nevertheless, for IDP solutions, our intuition is that the slab/Cs scenario is more physically plausible than the MG/BG scenario. Consistent with the slab/Cs picture in figure 8(a), just like dissolved folded proteins, dissolved IDPs occupy volumes excluded to water. Dissolved IDPs and folded proteins have similar densities (figure 11) of 1.32–1.52 g cm$^{-3}$ [102] and hence similar partial molar volumes on a per-gram basis. In contrast, the MG/BG picture in figure 8(b) invokes a negative effective molecular polarizability for IDP that counteracts an all-permeating water medium (\(\gamma_{\text{MG}} < 0\) because $\epsilon_p < \epsilon_w$). This scenario is apparently at odds with the reality of IDP excluded volume, suggesting that while the MG/BG models may be applicable in certain solid-state situations [101], they may be problematic for IDP solutions. A definitive resolution of this question awaits further theoretical and experimental investigation.

7. Conclusions

To recapitulate, we have taken a step to address the sequence-phase relationship in regard to how mixing/demixing of IDP components in membraneless organelles are governed by the IDPs’ amino acid sequences. Going beyond mean-field FH and OV approaches, RPA provides an approximate physical account of sequence effects [61–63]. Our findings point to a multivalent, stochastic, ‘fuzzy’ mode of molecular recognition in that mixing the populations of a pair of IDP sequences is favorable if their charge patterns are similar whereas population demixing is promoted when their charge patterns are very different. For the examples studied, a quantitative correlation is observed between the RPA-predicted tendency for a pair of sequences to demix in two (binary) coexisting phases and the difference in their SCD parameters. This predicted trend is qualitatively in line with the observed demixing of the nucleolar IDPs FIB1 and NPM1 because they have very different SCD’s, although a comparison of the experimental FIB1/NPM1 phase diagram [27] with our RPA and FH results indicates that inclusion of non-electrostatic interactions as well as more biomolecular species in the analysis will be necessary for a quantitative theoretical account of sequence-specific ternary coexistence. It should also be
noted that our current RPA formulation does not consider counterion condensation, which can be important for IDP sequences with high net charges. A recent transfer matrix theory [103] should be helpful in tackling such situations, although as it stands this theory does not address sequence dependence. Despite our theory’s limitations, the simple principles of sequence dependence suggested by the present effort should already be testable by in vitro experiments on IDP polymers [70]. As illustrated by our consideration of IDP-concentration-dependent permittivity, theoretical study of IDP phase separation is only in its infancy. The logical next steps in the development of RPA theory include extending to systems with more than two sequences and sequences with non-zero net charges. Much biophysics of IDP phase separation remains to be discovered.

Acknowledgments

We thank Alaji Bah, Robert Vernon (Hospital for Sick Children), Lewis Kay (University of Toronto), Che-Ting Chan (Hong Kong University of Science and Technology), Kingshuk Ghosh (University of Denver), Pak-Ming Hui (Chinese University of Hong Kong), and Huan-Xiang Zhou (University of Illinois at Chicago) for helpful discussions. We are also grateful to an anonymous referee for suggestions that led to the pedagogical graphics in figure 1. This work was supported by Canadian Cancer Society Research Institute grant no. 703477, Canadian Institutes of Health Research grant MOP-84281, and computational resources provided by SciNet of Compute/Calcu Canada.

References

[1] Uversky V N, Gillespie J R and Fink A L 2000 Why are ‘natively unfolded’ proteins unstructured under physiologic conditions? Proteins 41 115–27
[2] Dunker A K et al 2001 Intrinsically disordered proteins J. Mol. Graph. Model. 19 26–59
[3] Tompa P 2001 Intrinsically unstructured proteins Trends Biochem. Sci. 27 527–33
[4] Gunasekaran K, Tsiak C J, Kumar S, Zanuy D and Nussinov R 2003 Extended disordered proteins: targeting function with less scaffold Trends Biochem. Sci. 28 81–5
[5] Tompa P 2012 Intrinsically unstructured proteins: a 10-year recap Trends Biochem. Sci. 37 509–16
[6] van der Lee R et al 2014 Classification of intrinsically disordered regions and proteins Chem. Rev. 114 6589–631
[7] Liu Z and Huang F 2014 Advantages of proteins being disordered Protein Sci. 23 539–50
[8] Chen T, Song J and Chan H S 2015 Theoretical perspectives on nonnative interactions and intrinsic disorder in protein folding and binding Curr. Opin. Struct. Biol. 30 32–42
[9] Das R K, Ruff K M and Pappu R V 2015 Relating sequence encoded information to form and function of intrinsically disordered proteins Curr. Opin. Struct. Biol. 32 102–12
[10] Wright P E and Dyson H J 2015 Intrinsically disordered proteins in cellular signalling and regulation Nat. Rev. Mol. Cell Biol. 16 18–29
[11] Czajkowski V, Follis A V, Kriwacki R W and Forman-Kay J D 2016 Dynamic protein interaction networks and new structural paradigms in signalling Chem. Rev. 116 6424–62
[12] Bah A and Forman-Kay J D 2016 Modulation of intrinsically disordered protein function by post-translational modifications J. Biol. Chem. 291 6696–705
[13] Brangwynne C P, Eckmann C R, Courseon D S, Rybarska A, Hoege C, Gharaikhani J, Julicher F and Hyman A A 2009 Germline P granules are liquid droplets that localize by controlled dissolution/condensation Science 324 1729–32
[14] Li P et al 2012 Phase transitions in the assembly of multivalent signalling proteins Nature 483 336–40
[15] Kato M et al 2012 Cell-free formation of RNA granules: low complexity sequence domains form dynamic fibers within hydrogels Cell 149 753–67
[16] Lee C F, Brangwynne C P, Gharaikhani J, Hyman A A and Julicher F 2013 Spatial organization of the cell cytoplasm by position-dependent phase separation Phys. Rev. Lett. 111 088101
[17] Hyman A A, Weber C A and Julicher F 2014 Liquid–liquid phase separation in biology Annu. Rev. Cell. Dev. Biol. 30 39–58
[18] Toretsky J A and Wright P E 2014 Assemblages: functional units formed by cellular phase separation J. Cell. Biol. 206 579–88
[19] Wang T J, Smith J, Chen B C, Schmidt H, Rasoloson D, Paix A, Lambrus B G, Calidas D, Betzig E and Seydoux G 2014 Regulation of RNA granule dynamics by phosphorylation of serine-rich, intrinsically disordered proteins in C. elegans eLife 3 e04591
[20] Nott T J et al 2015 Phase transition of a disordered nuage protein generates environmentally responsive membraneless organelles Mol. Cell 57 936–47
[21] Elbaum-Garfinkle S, Kim Y, Szczepaniak K, Chen C C-H, Eckmann C R, Myong S and Brangwynne C P 2015 The disordered P granule protein LAF-1 drives phase separation into droplets with tunable viscosity and dynamics Proc. Natl. Acad. Sci. USA 112 7189–94
[22] Mollick D, Temirov I, Lee J, Coughlin M, Kanagaraj A P, Kim H J, Mittag T and Taylor J P 2015 Phase separation by low complexity domains promotes stress granule assembly and drives pathological fibrillization Cell 163 123–33
[23] Kroschwald S, Maharan S, Mateju D, Malinovská L, Nuske E, Poser I, Richter D and Alberts S 2015 Promiscuous interactions and protein disaggregates determine the material state of stress-inducible RNP granules eLife 4 e06807
[24] Nott T J, Crager T D and Baldwin A J 2016 Membraneless organelles can melt nuclear acid duplexes and act as biomolecular filters Nat. Chem. 8 569–75
[25] Banani S F, Rice A M, Peeples W B, Lin Y, Jain S, Parker R and Rosen M K 2016 Compositional control of phase-separated cellular bodies Cell 166 651–63
[26] Pak C W, Kono M, Holehouse A S, Padrick S B, Mittal A, Ali R, Yunus A A, Liu D R, Pappu R V and Rosen M K 2016 Sequence determinants of intracellular phase separation by complex coacervation of a disordered protein Mol. Cell 63 72–85
[27] Feric M, Vaida N, Harmon T S, Mitrea D M, Zhu L, Richardson T M, Kriwacki R W, Pappu R V and Brangwynne C P 2016 Coexisting liquid phases underlie nucleolar subcompartments Cell 165 1686–97
[28] Mitrea D M, Cika J A, Guy C S, Ban D, Banerjee P R, Stanley C B, Nourse A, Deniz A A and Kriwacki R W 2016 Nucleophosmin integrates within the nucleolus via multi-modal interactions with proteins displaying R-rich linear motifs and RNA life, J Cell 164 e13571
[29] Wu H and Fuxreiter M 2016 The structure and dynamics of higher-order assemblies: amyloids, signalosomes, and granules Cell 165 1055–106
[30] Bergeron-Sandoval L-P, Safaee N and Michnick S W 2016 Mechanisms and consequences of macromolecular phase separation Cell 165 1067–1079
[31] Riback J A, Katanski C D, Kear-Scott J L, Pilipenko E V, Bojek A E, Osnich T R and Drummond D A 2017 Stress-triggered phase separation is an adaptively tuned response Cell 168 1026–1040
[32] Shin Y and Brangwynne C P 2017 Liquid phase condensation in cell physiology and disease Science 357 aaf4382
[33] Meikar O, Da Ros M, Korhonen H and Kotttaz J N 2011 Chromatid body and small RNAs in male germ cells Reproduction 142 195–209
[34] Wang J T and Seydoux G 2014 P granules Curr. Biol. 24 R367–8
[35] Kederesna N, Ivanov P and Anderson P 2013 Stress granules and cell signaling: more than just a passing phase? Trends Biochem. Sci. 38 494–506
[36] Anderson P, Kederesna N and Ivanov P 2015 Stress granules, P–bodies and cancer Biochim. Biophys. Acta 1849 861–70
[37] Li Y R, King O D, Smitter J L and Gitter A D 2013 Stress granules as crucibles of ALS pathology J. Cell Biol. 201 361–72
[38] Ramaswami M, Taylor J P and Parker R 2013 Altered ribostastic RNA-protein granules in degenerative disorders Cell 154 727–36
[39] Zeng M, Shang Y, Araki Y, Guo T, Huganir R L and Zhang M 2016 Phase transition in postsynaptic densities underlies formation of synaptic complexes and synaptic plasticity Cell 166 1163–75
[40] Mittag T and Forman-Kay J D 2013 From sequence and forces to structure, function, and evolution of intrinsically disordered proteins Structure 21 1492–9
[41] Krezminski M, Marsh J A, Neale C, Choy W Y and Forman-Kay J D 2013 Characterization of disordered proteins with ENSEMBLE Bioinformatics 29 598–9
[42] Holehouse A S, Das R K, Ahmad N, Richardson M O G and Pappu R V 2017 CIDER: resources to analyze sequence-ensemble relationships of intrinsically disordered proteins Biophys. J. 112 16–21
[43] Holehouse A S and Pappu R V 2015 Protein polymers: encoding phase transitions Nat. Mater. 14 1083–4
[44] Chen J 2012 Towards the physical basis of how intrinsically disordered mediates protein function Arch. Biochem. Biophys. 524 123–31
[45] Song J, Ng C S, Tompa P, Lee K A W and Chan H S 2013 Polycation–π interactions are a driving force for molecular recognition by an intrinsically disordered oncoprotein family PLoS Comput. Biol. 9 e1003239
[46] Baker C M and Best R B 2014 Insights into the binding of intrinsically disordered proteins from molecular dynamics simulation WIREs Comput. Mol. Sci. 4 182–98
[47] Song J, Gomes G N, Gradinaru C C and Chan H S 2015 An adequate account of excluded volume is necessary to infer compactness and asphericity of disordered proteins by Förster resonance energy transfer J. Phys. Chem. B 119 15191–202
[48] Best R B 2017 Computational and theoretical advances in studies of intrinsically disordered proteins Curr. Opin. Struct. Biol. 42 147–54
[49] Levine Z A and Shea J E 2017 Simulations of disordered proteins and systems with conformational heterogeneity Curr. Opin. Struct. Biol. 43 95–103
[50] Rauscher S, Baud S, Miao K, Lee K F W and Pomès P 2006 Proline and glycine control protein self-organization into elastomeric or amyloid fibrils Structure 14 1667–76
[51] Ruff K M, Harmon T S and Pappu R V 2015 CAMELOT: a machine learning approach for coarse-grained simulations of aggregation of block-copolymeric protein sequences J. Chem. Phys. 143 243123
[52] Qin S and Zhou H-X 2016 Fast method for computing chemical potentials and liquid–liquid phase equilibria of macroscopic solutions J. Phys. Chem. B 120 8164–74
[53] Qin S and Zhou H-X 2017 Protein folding, binding, and droplet formation in cell-like conditions Curr. Opin. Struct. Biol. 43 28–37
[54] Kastelic M, Kaluzhnyi Y V, Hribar-Lee B, Dill K A and Vlachy V 2015 Protein aggregation in salt solutions Proc. Natl Acad. Sci. USA 112 6766–70
[55] Flory P J 1953 Principles of Polymer Chemistry (Ithaca, NY: Cornell University Press)
[56] Chen H S and Dill K A 1994 Solvation: effects of molecular size and shape J. Chem. Phys. 101 7007–26
[57] Pappu R V, Wang X, Vitals A and Crick S L 2008 A polymer physics perspective on driving forces and mechanisms for protein aggregation Arch. Biochem. Biophys. 469 132–41
[58] Brangwynne C P, Tompa P and Pappu R V 2015 Polymer physics of intracellular phase transitions Nat. Phys. 11 899–904
[59] Voorn M J 1956 Complex coacervation: I. General theoretical considerations Red. Trav. Chim. Pays-Bas 75 317–30
[60] Overbeek J T G and Voorn M J 1957 Phase separation in polyelectrolyte solutions. Theory of complex coacervation J. Cell. Physiol. 49 7–26
[61] Lin Y-H, Forman-Kay J D and Chan H S 2016 Sequence–specific polyanionpholyte phase separation in membraneless organelles Phys. Rev. Lett. 117 178701
[62] Lin Y-H, Song J, Forman-Kay J D and Chan H S 2017 Random-phase–approximation theory for sequence–dependent, biologically functional liquid–liquid phase separation of intrinsically disordered proteins J. Mol. Liq. 228 176–93
[63] Lin Y-H and Chan H S 2017 Phase separation and single–chain compactness of charged disordered proteins are strongly correlated Biophys. J. 112 2043–6
[64] Bohm D and Pines D 1951 A collective description of electron interactions: I. Magnetic interactions Phys. Rev. 82 625–34
[65] de Gennes P G 1979 Scaling Concepts in Polymer Physics (Ithaca, NY: Cornell University Press) pp 245–64
[66] Malhi K A and Olvera de la Cruz M 2000 Phase diagrams of salt-free polyelectrolyte semidilute solutions Macromolecules 33 7649–54
[67] Ermoshkin A V and Olvera de la Cruz M 2003 Polyelectrolytes in the presence of multivalent ions: gelation versus segregation Phys. Rev. Lett. 90 125504
[68] Das R K and Pappu R V 2013 Conformations of intrinsically disordered proteins are influenced by linear sequence distributions of oppositely charged residues Proc. Natl Acad. Sci. USA 110 13392–7
[69] Sawle I and Ghosh K 2015 A theoretical method to compute sequence dependent configurational properties in charged polymers and proteins J. Chem. Phys. 143 085101
[70] Quiriz F G and Chilkoti A 2015 Sequence heuristics to encode phase behaviour in intrinsically disordered protein polymers Nat. Mater. 14 1164–71
[71] Jain S, Wheeler J R, Walters R W, Agrawal A, Barsic A and Parker R 2016 ATPase–modulated stress granules contain a diverse proteome and substructure Cell 164 487–98

22
[72] Jacobs W M and Frenkel D 2017 Phase separations in biological systems with many components Biophys. J. 112 683–91
[73] Harmon T S, Holehouse A and Pappu R V 2017 To mix, or to demix, that is the question Biophys. J. 112 565–7
[74] Chan HS 1999 Folding alphabets Nat. Struct. Biol. 6 994–6
[75] Chan HS and Dill KA 1996 Comparing folding codes for proteins and polymers Proteins 24 335–44
[76] Wroe R, Bornberg-Bauer E and Chan HS 2005 Comparing folding codes in simple heteropolymer models of protein evolutionary landscape: robustness of the superfunnel paradigm Biophys. J. 88 118–31
[77] Tompa P and Fuxreiter M 2008 Fuzzy complexes: polymorphism and structural disorder in protein–protein interactions Trends Biochem. Sci. 33 2–8
[78] Sharma R, Raduly A, Miskei M and Fuxreiter M 2015 Fuzzy complexes: specific binding without complete folding FERS Lett. 589 2533–42
[79] Miskei M, Antal C and Fuxreiter M 2017 FuzDB: database of fuzzy complexes, a tool to develop stochastic structure–function relationships for protein complexes and higher-order assemblies Nucl. Acids Res. 45 228–35
[80] Borg M, Mittag T, Pawson T, Tyers M, Forman–Kay JD and Chan HS 2007 Polyelectrostatic interactions of disordered ligands suggest a physical basis for ultrasensitivity Proc. Natl Acad. Sci. USA 104 9650–5
[81] Mittag T, Marsh J, Grishaw A, Orlicky S, Lin H, Sicheri F, Tyers M and Forman–Kay JD 2010 Structure/function implications in a dynamic complex of the intrinsically disordered S1 with the Cdc4 subunit of an SCF ubiquitin ligase Structure 18 494–506
[82] Dejardins G, Meeker CA, Bhachech N, Currie SL, Okon M, Graves B J and McIntosh LP 2014 Synergy of aromatic residues and phosphoserines within the intrinsically disordered DNA-binding inhibitory elements of the Ets-1 transcription factor Proc. Natl Acad. Sci. USA 111 11019–24
[83] Csizmok V et al 2017 An allosteric conduit facilitates dynamic multisite substrate recognition by the SCF(Cdc4) ubiquitin ligase Nat. Commun. 8 15943
[84] Markel V A 2016 Introduction to the Maxwell Garnett approximation: tutorial J. Opt. Soc. Am. A 33 1244–56
[85] Delaglio F, Grzesiek S, Vuister G W, Zhu G, Pfeifer J and Bax A 1995 NMRPipe: a multidimensional spectral processing system based on UNIX pipes J. Biomol. NMR 6 277–93
[86] Brady JP et al 2017 Structural and hydrodynamic properties of an intrinsically disordered region of a germ-cell specific protein upon phase separation Proc. Natl Acad. Sci. USA 114 E194–203
[87] Gill SC and von Hippel PH 1989 Calculation of protein extinction coefficients from amino acid sequence data Anal. Biochem. 182 319–26
[88] Voorn MJ 1956 Complex coacervation: III. Thermodynamic calculations on three-component systems Reel. Trav. Chim. Pays-Bas 75 427–46
[89] Atkins P and de Paula J 2009 Physical Chemistry 9th edn (New York: Freeman)
[90] Kastelic M, Kalyuzhnyi Y V and Vlachy V 2016 Modeling phase transitions in mixtures of β–α lens crystallins Soft Matter 12 7289–98
[91] Grosberg A Y and Khokhlov A R 1994 Statistical Physics of Macromolecules (Woodbury, NY: AIP)
[92] Kraft D 1988 A Software Package for Sequential Quadratic Programming, Deutsche Forschungs- und Versuchsanstalt für Luft- und Raumfahrt Köln: Forschungsbericht (Köln: DFVLR)
[93] Jones E 2001 SciPy: open source scientific tools for Python (Accessed: 20, October 2017, http://scipy.org/)
[94] Leach AR 1996 Molecular Modeling. Principles and Applications (Singapore: Addison Wesley Longman Limited) p 170
[95] Elcock AH 1998 The stability of salt bridges at high temperatures: implications for hyperthermophilic proteins J. Mol. Biol. 284 489–502
[96] Zhou H-X 2002 Toward the physical basis of thermophilic proteins: linking of enriched polar interactions and reduced heat capacity of unfolding Biophys. J. 83 3126–33
[97] Dominy BN, Minoux H and Brooks CL 2004 An electrostatic basis for the stability of thermophilic proteins Proteins 57 128–41
[98] Sawle L, Huihui J and Ghosh K 2017 All-atom simulations reveal protein charge decoration in the folded and unfolded ensemble is key in thermophilic adaptation J. Chem. Theor. Comput. 13 5065–75
[99] Bragg W L and Pippard AB 1953 The form birefringence of macromolecules Acta Cryst. 6 865–7
[100] Jackson JD 1975 Classical Electrodynamics 2nd edn (New York: Wiley) pp 154–5
[101] Tsukerman I 2017 Classical and non-classical effective medium theories: New perspectives Phys. Lett. A 381 1244–56
[102] Fischer H, Polikarpov I and Craievich AF 2004 Average protein density is a molecular-weight-dependent function Protein Sci. 13 2625–8
[103] Lytle T K and Sing CE 2017 Transfer matrix theory of polymer complex coacervation Soft Matter 13 7001–12