Assembly of model postsynaptic densities involves interactions auxiliary to stoichiometric binding

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ABSTRACT The assembly of functional biomolecular condensates often involves liquid-liquid phase separation (LLPS) of proteins with multiple modular domains, which can be folded or conformationally disordered to various degrees. To understand the LLPS-driving domain-domain interactions, a fundamental question is how readily the interactions in the condensed phase can be inferred from interdomain interactions in dilute solutions. In particular, are the interactions leading to LLPS exclusively those underlying the formation of discrete interdomain complexes in homogeneous solutions? We address this question by developing a mean-field LLPS theory of two stoichiometrically constrained solute species. The theory is applied to the neuronal proteins SynGAP and PSD-95, whose complex coacervate serves as a rudimentary model for neuronal postsynaptic densities (PSDs). The predicted phase behaviors are compared with experiments. Previously, a three SynGAP/two PSD-95 ratio was determined for SynGAP/PSD-95 complexes in dilute solutions. However, when this 3:2 stoichiometry is uniformly imposed in our theory encompassing both dilute and condensed phases, the tie-line pattern of the predicted SynGAP/PSD-95 phase diagram differs drastically from that obtained experimentally. In contrast, theories embodying alternate scenarios postulating auxiliary SynGAP-PSD-95 as well as SynGAP-SynGAP and PSD-95-PSD-95 interactions, in addition to those responsible for stoichiometric SynGAP/PSD-95 complexes, produce tie-line patterns consistent with experiment. Hence, our combined theoretical-experimental analysis indicates that weaker interactions or higher-order complexes beyond the 3:2 stoichiometry, but not yet documented, are involved in the formation of SynGAP/PSD-95 condensates, implying future efforts to ascertain the nature of these auxiliary interactions in PSD-like LLPS and underscoring a likely general synergy between stoichiometric, structurally specific binding and stochastic, multivalent “fuzzy” interactions in the assembly of functional biomolecular condensates.

INTRODUCTION

Biomolecular condensates composing of various proteins, nucleic acids, and small molecules are important for biological functions (1). Examples include intracellular compartments, often referred to as “membraneless organelles,” e.g., P-granules, nuclear speckles, and Cajal bodies (2,3), and extracellular materials such as the precursory tropoelastin coacervates of elastic connective tissues (4). Some membraneless organelles, such as nucleoli and stress granules, are organized further into subcompartments (5,6). Biomolecular condensates serve diverse functions, including but not limited to gene regulation, cell growth, and synaptic activities (1,7–9). Membraneless organelles reduce biomolecular noise in the cell (10) and provide regulated local

**SIGNIFICANCE** It has become increasingly clear that functional biomolecular condensates underpinned by liquid-liquid phase separation (LLPS) are stabilized by dynamic multivalent interactions as well as by structurally specific interactions. To gain insights into the role of stoichiometric binding in biomolecular LLPS, we develop theories for LLPS driven solely by interactions that stabilize stoichiometric protein complexes in dilute solution and for alternate scenarios in which auxiliary interactions also contribute. Application of our formulations to experimental measurements of dilute- and condensed-phase protein concentrations of the SynGAP/PSD-95 condensate model of postsynaptic densities reveals that its assembly involves interactions auxiliary to those stabilizing the 3:2 SynGAP/PSD-95 complex in dilute solution, exemplifying a synergy between specific and stochastic interactions in the assembly of biomolecular condensates.
environments that facilitate specific biochemical processes, yet they can also assemble or disassemble rapidly in response to environmental changes (11). Biophysically, liquid-liquid phase separation (LLPS) is recognized as a major—albeit not the only (12)—mechanism in the assembly, including subcompartmentalization (5), of many of these condensates (13).

Main ingredients of biomolecular condensates can be intrinsically disordered proteins (IDPs), folded globular proteins, RNA, and/or proteins with folded domains and intrinsically disordered regions (IDRs) (5,14–16). When only parts of the proteins (e.g., folded domains (17)) engage in significant favorable interchain interactions while other parts (e.g., parts of or entire IDRs) do not, LLPS energetics may be conceptualized using a “stickers and linkers” picture (18–20). The sticker or linker distinction is not always clear-cut, however, because every part of a protein chain may contribute to LLPS-driving interactions. For folded domains, the interactions can be, though not always, structurally specific or entail “folding upon binding” when IDR-interacting partners are involved (21,22). In contrast, for interactions among IDPs and IDRs, the molecular recognition mechanisms underlying their sequence-dependent LLPS (14,23–26) are stochastic or “fuzzy” (27) in that they involve diverse, dynamic conformations and transient interactions (28–31). However, some of the transient interactions may entail fluctuations between disordered chain configurations with labile, but nonetheless specific, fibril-like structures (32,33).

It has been known for some time that proteins participating in signaling pathways are often organized structurally and sequentially in a modular fashion, with folded domains such as SH2 and SH3 acting as modules playing key regulatory roles (34). Recently, multivalent interactions involving some of these domains are found to be important for the assembly of biomolecular condensates. Thus, LLPS can be a key physicochemical mechanism exploited by nature for physiological regulation, as LLPS’s essentially all-or-none features entail sharp transitions among conformationally and hence functionally distinct states of these proteins (17).

In view of the central role of multivalency in biomolecular condensates (17,18,35), network concepts such as percolation transition (18–20,36) and graph theory (37,38) have been applied to provide rationalizations and theoretical formulations of LLPS and gelation (39). In this perspective, the nodes or “stickers” are folded or labile IDR-interacting domains, and the LLPS or gelation transition is determined in large measure by the number of connections (valence of attractive interactions) that can be effectuated between the nodes. Recent applications of this approach include elucidating the role of RNA-binding valence in creating sufficient connectivity of a ribonucleoprotein network (37) in the assembly of stress granules (40,41), and in the competition between gelation and LLPS of the cancer-related protein SPOP with its substrate (38).

### Postsynaptic densities

With these general considerations in mind, this work aims to gain insight into the LLPS-driving interactions of two modular proteins, synaptic Ras GTPase-activating protein (SynGAP) and postsynaptic density protein 95 (PSD-95), which are major constituents of postsynaptic densities (PSDs) (42). Recent advances indicate that biomolecular condensates play critical roles in neural function, probably because of their ability to respond rapidly to stimuli; thus, misregulated LLPS can lead to neurological diseases (43–46). In particular, phase separation is important for proper communication between neurons because LLPS is involved in the assembly of synaptic vesicles attached on presynaptic plasma membranes (47,48) as well as the PSDs beneath postsynaptic membranes (8,42,49,50).

Each PSD is a disk-shaped membraneless protein-rich compartment, an assembly made up of nucleic acids and thousands of different types of proteins (43,51). These include RNA-binding proteins (52); transmembrane layer proteins such as N-methyl-D-aspartate receptor (NMDAR) (53); actins in the cytoskeleton layer; scaffold proteins such as PDZ-domain-containing PSD-95s, which are membrane-associated guanylyl kinase (GK)-associated protein (GKAP) (55); guanylate kinase (GK)-associated protein (GKAP) (55); SH3 and multiple ankyrin repeat domain proteins (Shanks) (56); and the Homer family of adaptor proteins (57), as well as the SynGAP protein known to be highly enriched in PSDs (8,58). Situated adjacent to postsynaptic membranes, PSDs can exchange materials with the cytoplasm in synaptic spines and accordingly serve to provide spatial and temporal organization of the neurotransmitter receptors at the synapse (43). The functional importance of PSDs is highlighted by intriguing recent observations that PSDs are downsized during sleep (59,60), which suggest that synaptic strength is renormalized during sleep and that a SynGAP/PSD-95 condensate model of PSD (42) can be disassembled by moderate hydrostatic pressure (16,61), which points to a possible biophysical origin of the high-pressure neurological syndrome experienced by divers (62).

A minimal experimental model of PSD constructed using SynGAP and PSD-95, both of which are very abundant in PSDs, has been observed to self-organize into highly condensed, PSD-like droplets in vitro, offering the first indication that LLPS is a significant biophysical underpinning of PSD assembly (42). Subsequently, more realistic in vitro models of PSD—encompassing SynGAP and three scaffold proteins GKAP, Shank3, and Homer3 (in addition to PSD-95), as well as NR2B as another glutamate receptor—have also been constructed as a versatile research platform (8). Here, as a first step toward elucidating the statistical mechanics of PSD assembly, we focus on the simpler two-component SynGAP/PSD-95 construct, the same system we have utilized recently for studying the effect of hydrostatic pressure on PSD stability (61). SynGAP...
and PSD-95 undergo LLPS together when mixed, but each of the individual components, SynGAP or PSD-95 (up to 100 μM concentration each), does not phase separate by itself (42). It follows that the SynGAP/PSD-95 condensate is a complex coacervate, the formation of which must involve favorable interactions between SynGAP and PSD-95 molecules.

**LLPS of stoichiometric SynGAP/PSD-95 complexes as a possible mechanism of PSD assembly**

Wild-type SynGAP has a long coiled-coil (CC) domain that can dock onto two other such domains in an intertwining manner, resulting in a highly stable SynGAP trimer, which we term $S_3$ hereafter. The SynGAP trimer can, in turn, form a complex with two PSD-95 molecules (42). We refer to this SynGAP/PSD-95 complex with a 3:2 stoichiometry, which is observed in dilute solution of SynGAP and PSD-95, as $S_3P_2$. Experimental studies indicate that formation of the trimer $S_3$ is necessary for PSD-like complex coacervation because a SynGAP mutant that abolishes SynGAP’s ability to trimerize does not undergo LLPS in the presence of PSD-95 (42). Taken together, these observations raise the tantalizing possibility that units of the $S_3P_2$ complex act as the only nodes of favorable interactions in a SynGAP/PSD-95 condensate. In other words, the condensate is stabilized solely by interactions among the $S_3P_2$ complexes. Examples of how inter-$S_3P_2$ favorable interactions may arise from known interactions among SynGAP and PSD-95 domains (42,50) are provided in Fig. 1. Recent experiments indicate that the GK domain of a $P$ molecule interacts favorably with one of the PDZ domains of another $P$ molecule when that PDZ domain is bound to one of the PDZ binding domains (PBMs) of an $S_3$ (50). These GK-PDZ-PBM interactions can lead to a network of favorably interacting $S_3P_2$ complexes (Fig. 1, b and c). Although such interactions may lead merely to a linear array of favorably interacting $S_3P_2$ complexes (Fig. 1 b) as previously envisioned (50), they may also result in a three-dimensional network of favorably interacting $S_3P_2$ complexes (Fig. 1 c) and may therefore stabilize a condensed phase of $S_3P_2$ complexes. The hypothetical scenario that SynGAP/PSD-95 condensate is stabilized entirely by favorable interactions among units of $S_3P_2$ complexes represents a strong coupling between the $S_3 + 2P \rightleftharpoons S_3P_2$ binding equilibrium and LLPS-driving interactions. As such, it is instructive to ascertain, theoretically, the implications of this assumed scenario on the predicted LLPS properties. This knowledge would be useful, in general, for studying not only the SynGAP/PSD-95 systems but also other coacervate systems that involve a similar binding equilibrium or a chemical equilibrium of reversible reactions that takes an equivalent mathematical form as a binding equilibrium. The theoretically predicted mesoscopic phase properties, such as coexistence curves and phase diagrams, may then be compared against corresponding experimental observations to assess whether the assumed microscopic
stoichiometric pattern of network interactions is indeed the case or, alternatively, whether auxiliary nodes of favorable interactions in addition to S3P2 complexes are involved in stabilizing the condensed coacervate phase.

**Probing interaction networks in SynGAP/PSD-95 condensates by theory and LLPS experiment**

To this end, we endeavor to develop a theoretical formulation for protein complex coacervation with an emphasis on the interactions among the folded protein domains under constraints similar to those imposed by the modular organization, multivalency, and stoichiometry of the SynGAP/PSD-95 system. In contrast, most of the recent attention of theoretical and computational developments on biomolecular condensates has been on IDPs and IDRs. Although insights have been gained into multiple-component LLPS by several recent theoretical and modeling studies (e.g., (5,19,29,31,39,63)) and some folded domains such as the helicase domain in LAF-1 have been considered in explicit-chain simulations (64), the preponderance of the efforts thus far are on systems with a single IDP or IDR species. For instance, sequence-dependent IDP LLPS has been studied using analytical theory (23,65), coarse-grained explicit-chain lattice (20,66,67), and continuum (24,26,64,68) as well as field-theoretic (31,69) simulations. These efforts have focused on how the LLPS propensity of heteropolymeric chain molecules depends on the sequence of charged monomers (23,70,71) in general (as characterized, e.g., by quantitative charge pattern parameters (72,73)); the effects of sequence patterns of hydrophobic (74), aromatic (25), and other amino acid residues (26,64) on IDP phase behaviors; and the impact of sequence patterns on the phase properties of synthetic polymers (75,76).

The microscopic complexity of the SynGAP/PSD-95 system makes modeling its LLPS in atomic details with quantitative precision infeasible. To gain insight into the essential biophysics of the system, we construct a highly simplified model that treats each S3 or P molecule as a single particle that can engage in certain interactions suggested by experimental observations. The model is for LLPS driven largely by folded domain association. As such, it is substantially different from the aforementioned theories for IDP LLPS that take into account the conformational diversity of IDP chains. In this respect, the conceptual underpinnings of this model are akin to those of patchy-particle models of colloids (77–79), which have been applied recently to study phase behaviors of folded proteins such as lens crystallins (80), ribonucleoprotein droplets in which both the proteins and RNA are modeled as patchy articles (81), and the relationship between patchy-particle and coarse-grained explicit-chain models of biomolecular LLPSs including assembly of multicomponent condensates (82). The model developed herein is structurally and energetically more simplified than patchy-particle formulations in that the valencies of S3 and P are now treated in a mean-field manner without considering the interactions’ anisotropic directionality. This simplicity notwithstanding, our tractable model provides a critical assessment of SynGAP/PSD-95 coacervation scenarios because the model features a possible coupling between the formation of the 3:2 S3P2 complex and phase separation. We find that the constraints imposed by an exclusively complex-driven phase separation entail drastically different S3 and P compositions of coexisting phases from those predicted by classical Flory-Huggins theories (83,84) in the absence of such coupling. Therefore, whether SynGAP/PSD-95 coacervation is driven solely by interactions among S3P2 complexes can be assessed by comparing our theoretical predictions against experiment. With this recognition, we have now conducted extensive experimental measurements of S3P compositions of coexisting phases. The results are consistent with classical Flory-Huggins theory, but not the exclusively complex-driven LLPS theory, indicating a hitherto unknown prevalence of auxiliary interactions beside those among the S3P2 complexes in the SynGAP/PSD-95 condensed phase. Details of these findings and their ramifications are provided below.

**MATERIALS AND METHODS**

**A general mean-field theoretical framework for stoichiometric complex-driven LLPS**

Before delving into the particulars of the SynGAP/PSD-95 system, it is useful and instructive to first construct a general LLPS theory for a solution containing two species of solutes A and B that are capable of forming a stoichiometric complex, AB, (where x, y are two fixed positive integers) and whose complex coacervation is contingent upon, i.e., coupled to, the formation of such complexes. The formation and dissociation of the complex may be expressed as a reversible binding or chemical reaction:

\[
xA + yB \leftrightarrow A_x B_y.
\]

Now let the total solution volume be \(V\), and the number of solutes A and B in the solution be, respectively, \(n_A\) and \(n_B\). Proceeding with a Flory-Huggins (FH)-type lattice argument (83,85), we assume for simplicity that individual A and B solute molecules occupy equal volume, denoted by \(v_s\), and discretize the solution volume conceptually to a total of \(M = V/v_s\) lattice sites. Accordingly, a lattice site in the system can be occupied by an A or B solute molecule in its entirety, part of an A,B complex, or an entire solvent molecule. In other words, each A, B, or solvent molecule takes up one lattice site, whereas each A,B complex occupies a contiguous chunk of \(x + y\) sites.

Consider a situation in which \(m\) complexes are formed in the solution. The binding reaction and the configurational freedom of such a state is described by the partition function

\[
Z_{\text{bind}}(m) = \frac{M!e^{\theta_s/m}}{m!(n_A - mx)!(n_B - ym)(M - n_A - n_B)!},
\]

where \(-\theta_s < 0\) is the free energy associated with the favorable complex-forming binding (forward) reaction in Eq. 1. \(\theta_s = k_BT\) is Boltzmann constant, and \(T\) is absolute temperature. The free energy \(\theta_s\) does not account for the part of translational entropy described by the combinatoric factors in Eq. 2. Nonetheless, in general \(\theta_s\) can contain enthalpic as well as entropic components with entropic contributions from, e.g., change in the solvent orientational...
entropy associated with the binding reaction. It should also be noted that for
notational simplicity, a standard FH factor, \((z - 1)M^{n + y - 1}\), to account
for the spatial contiguity of each of the \(A_b B_b\) complexes (where \(z\) is the
coordination number of the lattice for a flexible, polymer-like complex
and \(z - 1 \rightarrow 1\) for a rigid complex) is omitted in Eq. 2 because the
factor may be formally absorbed into the entropic component of \(\epsilon_m\) by rede
fining \(\epsilon_m \rightarrow \epsilon_m + k_BT(x + y - 1)\ln[z - 1]/M\).

We further assume that LLPS of our model system is driven by short
spatial range contact-like interactions between individual bound solute A
or B molecules from different \(A_b B_b\) complexes. For each \(m\)-complex state,
the energetic contribution for these interactions is given by the following free
energy

\[
-k_BT \ln Z_{\text{LLPS}}(m) = -\frac{Z_{\text{LLPS}}}{M}\left(\frac{m}{2}\right)(x + y)^2 + O(m^3),
\]  

(3)

where \(Z_{\text{LLPS}}(m)\) is the pertinent partition function and \(\epsilon_m\) is the contact
free energy of an \(A-A, B-B,\) or \(A-B\) interaction (which are assumed to
carry the same \(\epsilon_m\) for simplicity). In situations in which these interactions
are relatively weak, as is apparently the case for the SynGAP/PSD-95
system according to a recent analysis of experimental data on the sys
tem’s sensitivity to hydrostatic pressure (61), one may consider only
the pairwise \(O(m^2)\) contribution in Eq. 3, which corresponds to the
lowest-order interaction term in coordinate-space polymer lattice cluster
theory (86).

It should be emphasized that the \(\epsilon_m\) for \(A-A, B-B,\) and \(A-B\) contacts in this
formulation is only for intercomplex interactions between different \(A_b B_b\)
units. The parameter \(\epsilon_m\) is not involved in the assembly of an individual
\(A_b B_b\) complex, which is stipulated separately by a chemical equilibrium
(Eq. 1) as will be described further below. In our mean-field formulation,
each unit of \(A_b B_b\) interacts as a whole in an isotropic manner. In this context,
assigning different energies to intercomplex \(A-A, B-B,\) and \(A-B\) contacts
would only amount, mathematically, to a redefinition of \(\epsilon_m\), which is effec
tively a weighted average intercomplex energy over different types of con
.tacts between the complexes’ \(A\) and \(B\) constituents. Going forward, it would be
interesting in future investigations to extend our approach to structurally
and energetically more realistic models that allow for anisotropic interac
tions, such as patchy-particle models (82). Under those extended frame
works, it would then be useful to study the effects of different intercomplex
energies for \(A-A, B-B,\) and \(A-B\) contacts.

Here, with \(Z_{\text{LLPS}}\) and \(Z_{\text{LLPS}}\) in place, the total partition function \(Z\) for our
model solution system that takes into account all possible \(m\)-complex states is
given by

\[
Z = \sum_{m=0}^{m_{\text{max}}} Z_{\text{LLPS}}(m) = \sum_{m=0}^{m_{\text{max}}} e^{-M(m)},
\]  

(4)

where \(m_{\text{max}}\), the integer part of \(\min[n_A x_A, n_B y_B]\), is the maximal number of
\(A_b B_b\) complexes possible for a given solution system and \(f(m)\) is the \(m\)-state
free energy per lattice site in units of \(k_BT\). Application of Stirling’s approxima
tion for the factorials to Eqs. 2 and 3 leads to the following approximate expression for \(Mf(m)\):

\[
Mf(m) \approx (n_A - x_m)\ln(n_A - x_m) + (n_B - y_m)\ln(n_B - y_m)
+ (M - n_A - n_B)\ln(M - n_A - n_B) + m \ln m
+ m\left(x + y - 1 - \frac{\epsilon_m}{k_BT}\right) - \chi \frac{(m(x + y))^2}{M} - M \ln M,
\]  

(5)

where \(\chi = \epsilon_m/(2k_BT)\) is an effective FH parameter (86,87). In arriving at
Eq. 5, we have neglected \(O(1/M)\) and \(O(\ln M/M)\) contributions to \(f(m)\) because these terms vanish in the \(M \rightarrow \infty\) thermodynamic limit. Accord
ingly, the \(ym(x + y)^2\) term from Eq. 3 and terms from the \((2\pi n)^{1/2}\) part of
the Stirling approximation \(n! = (2\pi n)^{1/2} e^{-\pi n}\) are not included in Eq. 5.

Following common notation in FH phase-separation theory, we define \(\phi_A \equiv n_A/M\) and \(\phi_B \equiv n_B/M\) as overall volume fractions for \(A\) and \(B\), respec
tively; \(\psi \equiv m(x + y)/M\) as the volume fraction for \(A_b B_b\); and \(\phi_A -
x\psi/(x + y)\) and \(\psi_B \equiv \phi_B - y\psi/(x + y)\) as volume fractions, respectively,
of unbound \(A\) and \(B\) (i.e., that do not form \(A_b B_b\)). Using this notation, Eq. 5 may be rewritten as

\[
f(\psi) = \psi_A \ln \psi_A + \psi_B \ln \psi_B + \frac{\psi}{x + y} \ln \frac{\psi}{x + y}
+ (1 - \phi_A - \phi_B) \ln(1 - \phi_A - \phi_B)
\]

(6)

where

\[
\K^0 = e^{-\epsilon_m/k_BT} = \frac{M^{x + y - 1}}{M^x + y - 1}
\]  

(7)

is a reduced (dimensionless) dissociation constant of \(A_b B_b\) in dilute solution
(the “0” superscript symbolizes dilute solution) of which we will be clarified below.

We are now ready to derive an approximate expression for the system
described by \(Z\) by replacing the summation in Eq. 4 with a single term at the saddle point \(\overline{\psi}\) of \(f(\psi)\). In other words, the free energy of the system

\[
\mathcal{F} = -k_BT \ln Z \approx Mf(\overline{\psi}),
\]  

(8)

where \(\overline{\psi}\) is obtained by solving

\[
0 = \frac{\partial f(\psi)}{\partial \psi} \bigg|_{\psi = \overline{\psi}} = -\frac{x}{x + y} \ln \overline{\psi}_A - \frac{y}{x + y} \ln \overline{\psi}_B
+ \frac{1}{x + y} \left[ \ln \overline{\psi}_A + \ln \K^0 \right] - 2\chi \overline{\psi}_A.
\]  

(9)

Multiplying every term on the right-hand side of Eq. 9 by \(x + y\) and taking the exponential of all terms result in the equivalent condition

\[
\frac{\overline{\psi}_A \overline{\psi}_B}{\overline{\psi}(x + y)} = \K^0 e^{-2\chi(x + y)\overline{\psi}_A}
\]  

(10)

which can be further rewritten as

\[
\frac{[A]^{y}[B]^y}{[A_b B_b]} = \K^0 e^{-2\chi(x + y)\overline{\psi}_A}\overline{\psi}_B,
\]  

(11)

where \([A], [B],\) and \([A_b B_b]\) are the equilibrium concentrations, respec
tively, of \(A\), \(B\), and \(A_b B_b\), and \(\K^0\) is the dissociation constant of \(A_b B_b\) in dilute solution. These quantities are given by

\[
[A] = \frac{\overline{\psi}_A}{V_x(x + y)} = \frac{\overline{\nu}_A}{V},
\]  

(12a)

\[
[B] = \frac{\overline{\psi}_B}{V_y(x + y)} = \frac{\overline{\nu}_B}{V},
\]  

(12b)
Experimental materials and methods

Protein expression and purification

The PSD-95 construct containing PDZ, SH3, and GK tandem (amino acid residues 306–1176 of the protein, referred to as PSD-95 in this work) was polymerase chain reaction (PCR) amplified from human complementary DNA (cDNA) library (NCBI: NP_001122359). The SynGAP construct containing a trimeric CC domain and a PSD-95 PDZ binding domain (PBM) (encoding amino acids 1147–1308, lacking 1192V–1197K and 1293E–1295G of the protein; referred to as SynGAP in this work) was PCR amplified from mouse cDNA library (UniProt: J3QQ18) (42). The genes encoding PSD-95 and SynGAP were individually cloned into vectors containing an N-terminal His6-affinity tag. Recombinant proteins were expressed in Escherichia coli BL21 (DE3) cells in Luria Broth (LB) medium at 16°C overnight and purified using a nickel-nitritotriacetic acid (Ni-NTA) agarose affinity column followed by size-exclusion chromatography (Superdex 200, GE Healthcare Life Sciences, Piscataway, NJ) with a column buffer containing 50 mM Tris (pH 8.2), 100 mM NaCl, 1 mM EDTA, 1 mM dithiothreitol (DTT). The fusion tag of each protein was cleaved by human rhinovirus (HRV) 3C protease and separated by another step of size-exclusion chromatography.

Sedimentation-based assay of phase separation and phase diagram

All proteins were precleared through high-speed centrifugation (16,873 × g 10 min) at 25°C before sedimentation assay. Proteins were mixed at each designed concentration in 50 μL total volume. After 10 min equilibration at room temperature, the mixture was centrifuged at 16,873 × g at 25°C for 10 min. The supernatant was collected, and pellet was thoroughly resuspended in 50 μL buffer. Proteins from supernatant and pellet fractions were analyzed through sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) with Coomassie blue R250 staining.

The phase diagram was constructed from the sedimentation-based phase-separation assay. Phase separation of a SynGAP/PSD-95 mixture was considered to occur if the concentrations of SynGAP and PSD-95 recovered from the pellet of the mixture were higher than the pellet fraction of the individual protein alone in each sedimentation assay.

Quantification of protein concentration in the condensed phase

Concentrations of PSD-95 in the condensed phase were measured according to the previous reported protocol (8). Briefly, Cy3-labeled PSD-95 was diluted into the final concentration of 1% by mixing with unlabeled protein. 1% Cy3-labeled PSD-95 was further mixed with SynGAP to form condensates. The mixture was injected into a 96-well glass-bottomed plate (Thermo Fisher Scientific, Waltham, MA). Confocal fluorescence images were captured using a Zeiss LSM880 confocal microscope with a 63× objective lens (Carl Zeiss, Oberkochen, Germany). The confocal slice images spanning the middle of each droplet were used to quantify the concentration of PSD-95.

To calculate concentration of PSD-95 in each droplet, a standard calibration curve was generated. Cy3-labeled PSD-95 alone in dilute solutions at different concentrations were added into a glass-bottomed plate. The fluorescence intensity of each confocal slice image was measured using the same settings as that for the droplet quantifications. According to the standard curve, the fluorescence intensity of PSD-95 in each condensate can be converted into absolute molar concentration.

RESULTS AND DISCUSSION

A mean-field theory of stoichiometric complex-driven SynGAP/PSD-95 LLPS

We are now in a position to apply the general theory developed above to the SynGAP/PSD-95 system, with the amino acid sequence information of the SynGAP and PSD-95 constructs (157 and 416 residues, respectively) specified above in Experimental materials and methods. For wild-type SynGAP, because of the extremely strong propensity to trimerize (42), we consider, as a very good approximation, that all SynGAP molecules exist as part of S3 trimers in solution. Accordingly, we set A = S3, B = P, x = 1, and y = 2 (Fig. 2 a) in the formalism of the last section to obtain from Eq. 6 the free energy for the SynGAP/PSD-95 system:

\[ f(S_3, P) = \frac{1}{3} \ln \frac{S_3}{P} + \frac{1}{3} \ln \frac{P}{S_3} + \left(1 - S_3 - P\right) \ln \left(1 - S_3 - P\right) + \frac{3}{2} \ln \left(1 - S_3 - P - \frac{S_3}{P}\right) - \psi \sqrt{3}, \]

where

\[ \psi_3 = S_3 - \psi / 3, \]
\[ \psi_p = P - 2\psi / 3, \]

and

\[ \frac{3}{2} \psi_3^2 = \psi_3 \psi_p \psi_0 \psi_4 \]

follow from Eq. 11. The LLPS-driving interaction network envisioned by this formulation is schematically illustrated in Fig. 2 b.

The corresponding phase-coexistence behaviors are solved by the numerical method described in (29). In general, for a solution with overall average protein volume fractions \((\phi_3^0, \phi_p^0)\), the free energy without phase separation is given by
The system will phase separate if there exists at least a set of \((v, \phi_{S_3}^0, \phi_P^0)\) values for which \(f_{\text{sep}} < f_{\text{bulk}}\). In that case, the final protein volume fractions—i.e., the binary coexistence phase boundary—are the volume fractions that minimize \(f_{\text{sep}}\). This minimization is achieved numerically by implementing a three-variable sequential least-squares programming algorithm (88) using the scipy.optimize.minimize function in SciPy, a Python-based numerical package for scientific computation (89). To ensure that there are no numerical errors in our calculation, we check the optimized sequential least-squares programming solutions against the following equations for chemical potential balance of coexisting phases to confirm that these conditions are indeed satisfied:

\[
\begin{align}
 f_{\text{sep}}(v, \phi_{S_3}^0, \phi_P^0) &= v f(\phi_{S_3}^0, \phi_P^0) + (1-v) f(\phi_{S_3}^0, \phi_P^0) \\
 +&(1-v)f\left(\frac{\phi_{S_3}^0 - v \phi_{S_3}^0}{1-v}, \frac{\phi_P^0 - v \phi_P^0}{1-v}\right). 
\end{align}
\] (18)

Whether the system phase separates is determined by the difference between \(f_{\text{bulk}}\) and the free energy \(f_{\text{sep}}\) of a phase-separated system with two coexisting phases labeled by \(\alpha\) and \(\beta\), viz.,

\[
f_{\text{sep}} = v f(\phi_{S_3}^0, \phi_P^0) + (1-v) f(\phi_{S_3}^0, \phi_P^0),
\] (16)

where \(v\) is the fraction of the total system volume in phase \(\alpha\). By definition, \(v\) satisfies \(0 < v < 1\) and the volume conservation conditions

\[
v \phi_{S_3}^0 + (1-v) \phi_{S_3}^0 = \phi_{S_3}^0
\] (17a)

and

\[
v \phi_P^0 + (1-v) \phi_P^0 = \phi_P^0,
\] (17b)

which, in turn, can be used to relate volume fractions in the \(\beta\) phase to those in the \(\alpha\) phase, allowing \(f_{\text{sep}}\) for overall volume fractions \(\phi_{S_3}^0, \phi_P^0\) to be rewritten as a function of \(v, \phi_{S_3}^0\), and \(\phi_P^0\).

\[
\begin{align}
 f_{\text{sep}}(v, \phi_{S_3}^0, \phi_P^0) &= v f(\phi_{S_3}^0, \phi_P^0) \\
 +&(1-v)f\left(\frac{\phi_{S_3}^0 - v \phi_{S_3}^0}{1-v}, \frac{\phi_P^0 - v \phi_P^0}{1-v}\right). 
\end{align}
\] (18)

The trend of phase behaviors predicted by the free energy given by Eq. 13 for this model of stoichiometric complex-driven LLPS is illustrated by the phase diagram in Fig. 2 c, which features an L-shape phase-separated regime (bound by the light blue solid lines and the diagonal black solid line with slope = -1). Apparently, a hallmark of stoichiometric complex-driven LLPS is a trend of converging tie lines (dark blue dashed lines in Fig. 2 c) from the dilute, protein-depleted phase boundary (large light blue L in Fig. 2 c) to the condensed, protein-rich phase boundary (small light blue L). When the overall concentrations (volume

\[
\begin{align}
 f_{\text{sep}}(v, \phi_{S_3}^0, \phi_P^0) &= v f(\phi_{S_3}^0, \phi_P^0) \\
 +&(1-v)f\left(\frac{\phi_{S_3}^0 - v \phi_{S_3}^0}{1-v}, \frac{\phi_P^0 - v \phi_P^0}{1-v}\right). 
\end{align}
\] (18)

The system will phase separate if there exists at least a set of \((v, \phi_{S_3}^0, \phi_P^0)\) values for which \(f_{\text{sep}} < f_{\text{bulk}}\). In that case, the final protein volume fractions—i.e., the binary coexistence phase boundary—are the volume fractions that minimize \(f_{\text{sep}}\). This minimization is achieved numerically by implementing a three-variable sequential least-squares programming algorithm (88) using the scipy.optimize.minimize function in SciPy, a Python-based numerical package for scientific computation (89). To ensure that there are no numerical errors in our calculation, we check the optimized sequential least-squares programming solutions against the following equations for chemical potential balance of coexisting phases to confirm that these conditions are indeed satisfied:

\[
\begin{align}
 f_{\text{sep}}(v, \phi_{S_3}^0, \phi_P^0) &= v f(\phi_{S_3}^0, \phi_P^0) + (1-v) f(\phi_{S_3}^0, \phi_P^0) \\
 +&(1-v)f\left(\frac{\phi_{S_3}^0 - v \phi_{S_3}^0}{1-v}, \frac{\phi_P^0 - v \phi_P^0}{1-v}\right). 
\end{align}
\] (18)

The trend of phase behaviors predicted by the free energy given by Eq. 13 for this model of stoichiometric complex-driven LLPS is illustrated by the phase diagram in Fig. 2 c, which features an L-shape phase-separated regime (bound by the light blue solid lines and the diagonal black solid line with slope = -1). Apparently, a hallmark of stoichiometric complex-driven LLPS is a trend of converging tie lines (dark blue dashed lines in Fig. 2 c) from the dilute, protein-depleted phase boundary (large light blue L in Fig. 2 c) to the condensed, protein-rich phase boundary (small light blue L). When the overall concentrations (volume
fractions) of S₃ and P are in the dilute-solution stoichiometric ratio of 1:2 ($φ₀^S/φ₀^P = 1/2$), the tie line has slope = 1/2, coinciding exactly with the black solid line in Fig. 2c with the same slope, indicating that this 1:2 stoichiometric ratio is maintained in both the dilute and condensed phases. When $φ₃^S/φ₃^P ≠ 1/2$, the molecular species (S₃ or P) that is in excess of the 1/2 ratio has no effect on phase separation because they do not generate more S₃P₂ complexes, and, by model construction, only interactions among S₃P₂ complexes are capable of driving LLPS. Consequently, the tie lines connecting coexisting dilute and condensed phases are converging toward the $φ₃/φ₃ = 1/2$ line, meaning that the condensed phase is always closer than the dilute phase to the $φ₃/φ₃ = 1/2$ stoichiometric ratio if $φ₃^S/φ₃^P ≠ 1/2$ because of the larger number of favorable interactions among S₃P₂ complexes in the condensed phase. With this ramification of a hypothetical stoichiometric complex-driven LLPS elucidated, we now proceed to test experimentally whether the real SynGAP/PSD-95 system indeed undergoes such a complex coacervation process.

**Experimental measurements of the PSD-95 concentrations in the condensed phase of SynGAP/PSD-95 mixtures**

Our previous study showed that SynGAP and PSD-95 could undergo phase separation together. In the dilute phase, the two proteins form a homogenous 3:2 complex (42). However, the binding mode of SynGAP and PSD-95 in the condensed phase is unknown. To explore whether the two proteins also form a stoichiometric 3:2 complex in the condensed phase, we measured concentrations of the two proteins in the dilute phase and in the condensed phase (labeled in Fig. 3 as “S” for supernatant and “P” for pellet, respectively, as in Fig. 4 of (42), not to be confused with the notation S for SynGAP and P for PSD-95). SynGAP and PSD-95 were mixed at different molar ratios. Sedimentation-based assay was used to assess their distributions in the dilute and condensed phases by SDS-PAGE with Coomassie blue staining (Fig. 3a). The percentages of SynGAP and PSD-95 recovered in the pellet were analyzed. A phase diagram of the SynGAP/PSD-95 system was then constructed according to the data obtained from the sedimentation-based assay. Here, the phase diagram in Fig. 3b shows that complex coacervation of SynGAP and PSD-95 is highly sensitive to the concentrations of the two proteins. When concentrations of the two components were higher than certain threshold values, phase separation of the mixtures occurred persistently. Interestingly, when fixing the concentration of one component at a value below a certain threshold, increasing the concentration of the other component could also induce phase separation. For instance, there was no phase separation when SynGAP was at 30 μM and PSD-95 was at 20 μM. However, when the concentration of SynGAP was increased to 120 μM with [PSD-95] unchanged at 20 μM, phase separation of the mixture was observed (Fig. 3b). This observation suggests that SynGAP and PSD-95 not only interacted with each other through the PDZ domain of PSD-95 and the PBM of SynGAP, but that the interactions might also involve other sites of the two proteins. To quantitatively measure the absolute concentration of PSD-95 in the condensed phase so as to enable the theoretical analysis below that provides a clearer answer to this basic question, we used a confocal fluorescence image-based method developed in our earlier study (8) (Fig. 3c and d). The fluorescence intensity of PSD-95 in the condensed droplets under each condition was converted into the absolute concentration of the protein based on the standard curve of Cy3-labeled PSD-95 obtained in dilute solutions (Fig. 3c). The derived absolute concentrations of PSD-95 under the conditions shown in the phase diagram of the PSD-95 and SynGAP mixtures (Fig. 3b) are shown by the heat map in Fig. 3e.

**Experimental concentrations of SynGAP and PSD-95 in coexisting condensed and dilute phases**

The next step in our investigation is to analyze the experimental data to determine whether they are consistent with the predicted behaviors in Fig. 2c for the hypothetical scenario in which LLPS is driven solely by favorable interactions among stoichiometric S₃P₂ complexes. Consider an aqueous solution of S and P with initial concentrations [S₀] and [P₀]. Using a notation similar to that in Eqs. 16, 17, and 18 for the hypothetical system when it has undergone phase separation, we define $v$ as the fraction of total volume in the dilute phase, and thus $1 - v$ is the fraction of total volume in the condensed phase. It follows from the conservation of S and P that

$$[S^0] = v[S^{dil}] + (1 - v)[S^{cond}],$$

(21a)

and

$$[P^0] = v[P^{dil}] + (1 - v)[P^{cond}],$$

(21b)

where quantities in the dilute and condensed phases are labeled, respectively, by the superscripts “dil” and “cond.” Note that we have defined the concentrations in Eqs. 21a and 21b purposely to correspond to those measured in the experiments (Fig. 3) in that they refer to the total concentrations of S and P molecules or their concentrations in the dilute and condensed phases irrespective of their binding status. Nonetheless, based on prior experiments (42) as noted in the introductory discussion above, we recognize that virtually all S molecules are in the form of S₃ under these experimental conditions such that $[S^0] ≈ 3[S^0]$, $[S^{dil}] ≈ 3[S^{dil}]$, and $[S^{cond}] ≈ 3[S^{cond}]$ for the concentrations, whereas the volume fraction $φ₃ = φ₃$. To construct a binary phase diagram, our goal here is to determine $[S^{dil}]$, $[S^{cond}]$, $[P^{dil}]$, and $[P^{cond}]$ from two sets
of experiments: 1) the 121 centrifugation-based assays providing the fractional numbers of protein molecules in the condensed phase, denoted here as

$$g_S = \frac{(1 - v) [S]^{\text{cond}}}{[S]^0},$$

(22a)

and

$$g_P = \frac{(1 - v) [P]^{\text{cond}}}{[P]^0},$$

(22b)

and 2) the 25 confocal microscopy measurements affording the condensed-phase concentration, $[P]^{\text{cond}}$, of PSD-95 (P), which may be identified, aside from a proportionality constant, to the volume fraction of P:

$$\phi_P^{\text{cond}} \propto [P]^{\text{cond}}.$$  

(23)

Because the $\gamma_S$ and $\gamma_P$ in Eqs. 22a and 22b determined from centrifugation measurements (two experimental data points per condition) are by themselves insufficient to determine both $[S]^{\text{dis}}$ and $[P]^{\text{dis}}$, as well as $v$ (three variables) in Eqs. 21a and 21b, we can only make the full determination of dilute- and condensed-phase protein concentrations for the 25 conditions for which confocal microscopy data are also available, with

$$[S]^0 = 30, 60, 90, 120, 180 \ \mu\text{M},$$

$$[P]^0 = 20, 40, 60, 80, 120 \ \mu\text{M}.$$ 

(22b)

For each of these given conditions, we may first use Eq. 22b to obtain

$$v = 1 - \frac{[P]^0}{[P]^{\text{cond}}}.$$  

(24)
by using $\gamma_P$ from centrifugation measurements and $[P^{\text{cond}}]$ from confocal microscopy. Once $v$ is determined, $[P^{\text{dil}}]$ follows from Eq. 21b as

$$[P^{\text{dil}}] = \frac{[P^0](1 - \gamma_P)}{v},$$

and the SynGAP concentrations in the dilute and condensed phases follow, respectively, from Eqs. 21a and 22a as

$$[S^{\text{dil}}] = \frac{[S^0](1 - \gamma_S)}{v}$$

and

$$[S^{\text{cond}}] = \gamma_S \frac{[S^0]}{1 - v}$$

by using the solved value of $v$ and $\gamma_S$ from centrifugation measurements.

The general trend of binary SynGAP/PSD-95 phase behaviors inferred from experimental data is not consistent with the stoichiometric complex-driven LLPS scenario

The above-described analysis of experimental data yields the $([P^{\text{dil}}], [S^{\text{dil}}])$ and $([P^{\text{cond}}], [S^{\text{cond}}])$ data points in Fig. 4 (orange and blue dots) as well as the tie lines connecting their coexisting values (dashed lines in Fig. 4). Because the protein concentrations of the initial states in the experiments are very close to those of the dilute phases and the dilute-phase concentrations are not measured directly, there is appreciable scatter in the $([P^{\text{dil}}], [S^{\text{dil}}])$ and $([P^{\text{cond}}], [S^{\text{cond}}])$ data points. This is because when initial concentrations of S and P are low, the droplets formed through phase separation are very small. The thickness of individual droplet is often smaller than the confocal optical section thickness (~0.9 μm). In such situations, the fluorescence intensity measured by confocal microscope is not only affected by fluorescence signal in the droplet but also interfered with by the glass of the coverslip and labeled proteins in solution. This basic limitation precludes an accurate quantification of the uncertainties in measured condensed-phase concentrations when overall protein concentrations are low, making it impractical to construct a smooth phase boundary from the plotted data. In future investigations, this difficulty can possibly be overcome by using solutions that lead to larger droplets, in which case the quality of the data may be evaluated by comparing signal intensities for different Z stacks obtained by confocal microscopy. For instance, if the thickness of droplet is larger than the confocal optical section thickness, the fluorescence intensities of several layers spanning the middle of each droplet should be comparable (see, e.g., Fig. 3 of (8)).

Experimental limitations notwithstanding, one unmistakably clear message from Fig. 4a is that the tie lines diverge from the protein-depleted phases (orange data points in Fig. 4a) to the protein-rich phases (blue data points in Fig. 4a), a trend that is opposite to the converging tie lines from the dilute, protein-depleted phase boundary to the condensed, protein-rich phase boundary in Fig. 2c. This information is of critical importance, despite noted experimental uncertainties in measuring concentrations, because if the stabilizing interactions in the SynGAP/PSD-95 coacervate are only those between $S_3P_2$ complexes, the relative concentrations of S and P in the condensed phase should not change or should not vary much even when experimental uncertainties are taken into account. As discussed above, the converging tie lines and the L-shaped phase boundary in Fig. 2c are direct consequences of the premise, based on the scenario illustrated in Fig. 2b, that the only LLPS-driving favorable interactions are those among stoichiometric $S_3P_2$ complexes. In that hypothetical scenario, tie lines would have negative as well as positive slopes (positive slopes when overall S and P concentrations are relatively low, negative slopes when either S or P concentration is high; see Fig. 2c). In contrast, the slopes of the experimental tie lines in Fig. 4a are all positive, spanning an approximate range from 0.24 to 9.1. It stands to reason, therefore, based on the fact that the trend inferred from experimental data in Fig. 4a is drastically different from that in Fig. 2c, that the data imply that favorable interactions among stoichiometric $S_3P_2$ complexes are not the only (or not at all) the LLPS-driving interactions in the assembly of the SynGAP/PSD-95 coacervate.
Theoretical analysis of experimental data suggests that nonstoichiometric auxiliary interactions are involved in the assembly of SynGAP/PSD-95 coacervates

To rationalize the SynGAP/PSD-95 phase properties in Fig. 4, it is useful to note that the trend of diverging tie lines in Fig. 4 c is similar to that predicted by the simple FH model for certain interaction strengths, such as that reported in Fig. 6 a of (29). Accordingly, we now apply FH approaches to assess alternate scenarios of SynGAP/PSD-95 in Fig. 6 model for certain interaction strengths, such as that reported in Fig. 4. Because SynGAP/PSD-95 LLPS must involve interaction between S and P (S or P does not individually phase separate), one may first consider a simple FH model with free energy in units of $k_B T M$ given by

\[
 f_1(\phi_S, \phi_P) = \phi_S \ln \phi_S + \phi_P \ln \phi_P + (1 - \phi_S - \phi_P) \ln (1 - \phi_S - \phi_P) - \chi_{SP} \phi_S \phi_P, \tag{27}
\]

where $\chi_{SP}$ is the favorable energetic (FH $\chi$) parameter characterizing S-P contacts as the only LLPS-driving interactions in this scenario (Fig. 5 a). An example of the binary phase diagrams predicted by this model is provided in Fig. 5 c. In contrast to the converging tie lines in Fig. 2 c, the tie lines in Fig. 5 c exhibit some degree of divergence from the protein-depleted to the protein-rich phase boundary, but the divergence is not as prominent as that in Fig. 4 a. Recognizing that the simple free energy function $f_1(\phi_S, \phi_P)$ in Eq. 27 does not provide any bias toward formation of $S_3P_2$ complexes, although such complexes are observed experimentally in dilute solution (42), it is reasonable to test whether incorporating favorability toward formation of $S_3P_2$ complexes in the model would result in predicted phase diagrams that conform better with the experimental trend. We do so by making the following modifications to the free energy $f(\phi_S, \phi_P)$ in Eq. 13 (which already contains favorable energy for $S_3P_2$ formation):

\[
 \tilde{\chi}_{SP} (\text{Eq. 13}) \rightarrow \chi_{SP} \phi_S \phi_P + \frac{1}{2} [\chi_{SS} \phi_S^2 + \chi_{PP} \phi_P^2] \tag{28a}
\]

and

\[
 \tilde{\chi}_{SP} = \chi_{SP} \phi_S \phi_P \exp(\tilde{\chi}_{SP}) (\text{Eq. 14c}) \rightarrow \chi_{SP} \phi_S \phi_P \exp(\chi_{SP}) \tag{28b}
\]

wherein Eq. 28a stipulates that individual $S_3$ trimers (not complexed with $P$ in the manner of the $S_3P_2$ complex) and individual $P$ molecules may participate in LLPS-driving interactions and/or that higher-order SynGAP/PSD-95 complexes in addition to $S_3P_2$ are present when protein concentrations are high, and these higher-order complexes also participate in LLPS-driving interactions because the two scenarios are accounted for by the same highly coarse-grained, mean-field formulation in Eq. 28a (Fig. 5 b). Moreover, unlike the stoichiometric complex-driven LLPS model free energy in Eq. 13, by dropping the $\exp(\tilde{\chi}_{SP})$ factor in Eq. 14c, Eq. 28b signifies that $S_3P_2$ formation is not coupled to the LLPS-driving interactions provided by the FH $\chi$ parameters (right-hand side of Eq. 28a) in the modified formulation described by Eq. 28. The modified formulation describes a system in
which S₃P₂ complexes are formed at low protein concentrations, whereas phase separation at higher protein concentrations is underpinned by a multitude of different S₃ and P interactions.

As exemplified by the two example phase diagrams in Fig. 5, d and e, the trend of tie-line divergence predicted by this modified formulation is more akin to the experimental results in Fig. 4 a than that predicted by the simple FH model in Fig. 5 c, although the differences between the predicted tie-line patterns among Fig. 5, c–e are not dramatic. Between the two models in Fig. 5, d and e with the same strongly favorable S-P interactions, it is instructive to note that the model in Fig. 5 e with favorable S-S and P-P interactions (consistent with experiment, these interactions are chosen to be insufficiently strong by themselves for S or P to phase separate individually) exhibits more tie-line divergence and thus captures slightly better the tie-line trend in Fig. 4 a than the model in Fig. 5 d with only favorable S-P interactions. In particular, the range of tie-line slopes, ([Scond] – [Sdil])/(Pcond) – [Pdil]), for the experimental results in Fig. 4 a (slope ~0.24–9.1, as mentioned above) are quite similar to that for the theoretical results in Fig. 5 e (slope ~0.28–12.0, calculated by multiplying each volume-fraction slope by a factor of 3). This model feature suggests that auxiliary S-S, P-P, and S-P interactions in addition to those associated with S₃P₂ assembly, rather than auxiliary S-P interactions alone, are likely in play in the phase separation of the experimental SynGAP/PSD-95 system as well. Nonetheless, the similarity of the tie lines in Fig. 5, c–e and the robustness of their overall pattern of divergence indicate that the experimentally observed phase properties are largely underpinned by auxiliary favorable S-P interactions.

CONCLUSIONS

It has become increasing apparent that the assembly and disassembly of biomolecular condensates are driven by stochastic, “fuzzy” interactions involving intrinsically disordered protein regions (see, e.g., (2,9,14,90)) as well as structurally specific interactions, including stoichiometric binding of folded domains in some instances (see, e.g., (8,33,37,91,92)). Deciphering how these interactions impart molecular recognition and how they may act in concert to achieve function is of fundamental importance to molecular biology (93,94), as has been emphasized in a recent study of the critical role of specific binding of the GIT1 and β-Pix GTPase regulatory enzymes in their complex coacervation (92). At a smaller length scale, a similar synergy between structurally specific interactions and stochastic, dynamic multivalent interactions has previously been indicated in the functional binding of IDPs in discrete, binary fuzzy complexes (95,96). In this context, the formation of the stoichiometric SynGAP/PSD-95 complex S₃P₂ in dilute solution (42) raises interesting questions regarding the role of these stoichiometric complexes in SynGAP/PSD-95 LLPS. Intuitively, a possible scenario would be that LLPS is driven solely by favorable interactions among S₃P₂ complexes. Theoretical considerations reveal that this scenario implies a peculiar form of binary phase diagrams with an L-shaped phase boundary and converging tie lines. However, subsequent experiments indicate that the LLPS properties of SynGAP/PSD-95 do not conform to this hypothetical scenario. Instead, our combined theoretical/experimental approach suggests strongly that auxiliary interactions beside those responsible for the formation of S₃P₂ are also involved in stabilizing the SynGAP/PSD-95 condensate. Our analysis has thus paved the way for investigating the nature of these auxiliary interactions. It appears that such LLPS-driving auxiliary interactions are readily available for certain folded proteins, as in the cases of lysozyme and crystallin (16). It would be instructive to compare SynGAP/PSD-95 with these simpler systems and also explore a possible involvement of disordered regions (97) in the formation of SynGAP/PSD-95 condensates. For instance, based on the interface between the third PDZ and SH3-GK domains in PSD-95 (98), the contacts leading to higher-order structures among PSD-95 molecules likely involve both hydrophobic and ionic interactions. Moreover, bioinformatics considerations (99) suggest that a stretch of chain sequence enriched in arginines, lysines, and glutamic acids between the coiled-coil and PBM domains of SynGAP is likely disordered. Such IDR s are potential participants in LLPS-driving interactions as well. In this regard, recently employed techniques for characterizing the roles of electrostatic and nonionic interactions in biomolecular condensates (100,101) would be useful. In any event, having demonstrated the capability of the relatively simple formulation developed here to gain fundamental insights into the mechanism of complex coacervation of SynGAP and PSD-95, we hope that our methodology will be useful for analyzing the role of stoichiometric complexes in the assembly of other biomolecular condensates as well.

AUTHOR CONTRIBUTIONS

Y.-H.L., M.Z., and H.S.C. designed research. Y.-H.L., H.W., and B.J. performed research. Y.-H.L. and H.W. analyzed data. Y.-H.L., H.W., M.Z., and H.S.C. wrote the article.

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