BIOPHYSICS

Biomolecular condensates can both accelerate and suppress aggregation of α -synuclein

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Biomolecular condensates present in cells can fundamentally affect the aggregation of amyloidogenic proteins and play a role in the regulation of this process. While liquid-liquid phase separation of amyloidogenic proteins by themselves can act as an alternative nucleation pathway, interaction of partly disordered aggregation-prone proteins with preexisting condensates that act as localization centers could be a far more general mechanism of altering their aggregation behavior. Here, we show that so-called host biomolecular condensates can both accelerate and slow down amyloid formation. We study the amyloidogenic protein α -synuclein and two truncated α -synuclein variants in the presence of three types of condensates composed of nonaggregating peptides, RNA, or ATP. Our results demonstrate that condensates can markedly speed up amyloid formation when proteins localize to their interface. However, condensates can also significantly suppress aggregation by sequestering and stabilizing amyloidogenic proteins, thereby providing living cells with a possible protection mechanism against amyloid formation.

INTRODUCTION

With increasing life expectancy, neurodegenerative diseases involving pathological amyloid formation are becoming alarmingly common. Misfolded and aggregated proteins may accumulate during the lifetime of a cell when these are not refolded or cleared by protein quality control machinery (1). Such accumulation can hamper regular cell operations, eventually leading to cell death and, at the organ level, to various neurodegenerative diseases including Alzheimer's and Parkinson's diseases (2-4). For many years, researchers have studied the origins, mechanism, and toxicity of protein aggregation, to develop effective therapies, but many aspects of the mechanism of protein aggregation remain incompletely understood (5-8). The quest for new therapies is impeded by the fact that processes inside cells take place in a complex environment that is difficult to reproduce in vitro (9). By contrast, the vast majority of protein aggregation studies are carried out with synthetic peptides or protein fragments in dilute, homogeneous, and well-mixed solutions (10, 11).

One of the most notable features that distinguish the complex intracellular environment from the protein solutions often used in vitro is the presence of condensates formed by liquid-liquid phase separation (LLPS) of biomolecules into so-called membraneless or-ganelles (MLOs) (12–15). These compartments are usually liquid-like, highly concentrated droplets of proteins and nucleic acids. Examples of such organelles include nucleoli (16) and Cajal bodies (17) in the nucleus and stress granules in the cytoplasm (18). The main difference between MLOs and membrane-bound compartments is the lack of a physical barrier between the organelle and the surrounding solution. This results in the ability to exchange components with the environment, to undergo fusion, and to respond to environmental changes by rapid formation/dissolution (19).

Knowing that MLOs contain proteins at very high concentrations and that proteins that undergo LLPS and proteins that partition into Copyright © 2022 The Authors, some rights reserved; exclusive licensee American Association for the Advancement of Science. No claim to original U.S. Government Works. Distributed under a Creative Commons Attribution License 4.0 (CC BY).

liquid droplets often feature low-complexity domains (13, 20), a characteristic that is also common for amyloidogenic proteins, it becomes evident that the presence of biological condensates could drastically affect the aggregation process. For various phase-separating proteins, it has been suggested that prior condensation into liquid droplets can promote conformational changes within the disordered region, leading to the formation of gel-like structures or amyloid-like aggregates. Such a process has been observed, for instance, for hnRNPA1 (18), FUS (21), or Tau (22, 23). Recently, it has been shown that also α -synuclein (α Syn), one of the archetypical amyloid-forming proteins, can undergo LLPS under polyethylene glycol (PEG)-based crowded conditions and that the condensed a Syn droplets may facilitate aggregation (24, 25). However, it remains unclear whether LLPS of α Syn LLPS is also likely to happen in living cells, as α Syn is known to interact with many components inside the cell, including membranes, the cytoskeleton, and other proteins (26, 27), which may suppress the concentration of free α Syn and prevent the formation of homotypic aSyn condensates.

Nonetheless, there is also another, more general way by which LLPS can affect protein aggregation, which is also relevant for proteins that are present in cells at low concentrations. Condensates can concentrate guest biomolecules, including amyloidogenic proteins, by partitioning or interfacial adsorption, and provide a distinct chemical environment in which the stability and reactivity of biomolecules may be affected. This can alter the kinetics of protein aggregation in multiple ways (13, 28, 29). An enhanced local concentration of amyloidogenic proteins may result in acceleration of the aggregation process, according to the law of mass action (30). However, one has to take into account that the local environment of the condensed liquid may promote protein conformations that do not undergo aggregation as readily as ones dominating in the surrounding solution. This has been observed for amyloid- β (1-42) (31) and may occur also for other amyloidogenic proteins. Last, aggregating proteins could accumulate at the interface between condensate and the surrounding cytosol, potentially resulting in alternative, interfacial aggregation pathways, analogous to what has been observed for lipid vesicles and solid surfaces (32). In general, accumulation of amyloidogenic proteins at an interface can alter the kinetics

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of aggregation in two ways: An increased local concentration leads to faster aggregation, and an altered conformation of molecules bound to the interface can either stabilize free monomers or promote their transformation into fibrils.

Accumulation of a Syn at the interface has been reported to accelerate aggregation, e.g., for exosomes and SUVs (small unilamellar vesicles) (33). By contrast, aggregation was slowed down when SUVs were present in large excess over α Syn, and most protein monomers were trapped in a stable configuration at the surface of SUVs and there was no free monomeric α Syn in solution (34). Similar effects have been observed for SUVs and LUVs (large unilamellar vesicles) composed of mixtures of anionic, cationic, and neutral lipids (35). The effect of lipid membranes on α Syn is largely dependent not only on the lipid/protein ratio but also on the chemical structure of lipid and mutations in the protein chain and probably also on the size (and thus the curvature) of the vesicles or surface defects associated with curvature (36). It has been suggested that the presence of membranes can even induce fibril dissociation by stabilizing monomers and depleting the solution of free protein (37).

While there is ample evidence that biomolecular condensates can fundamentally alter protein distributions in vitro and in living cells by concentration, exclusion, or interfacial localization, a systematic investigation of the effects that preexisting condensates have on protein aggregation is lacking. Here, we study the consequences of "inert" model condensates (coacervates) on the aggregation of αSyn as a key amyloid-forming protein (supplemented with experiments using insulin). Our model condensates are inert in the sense that they are not composed of, or dependent on, the aggregating protein, and they do not undergo any form of liquid-to-solid transition themselves. The goal of using these model condensates is to investigate whether preexisting biomolecular condensates can have a generic effect on protein aggregation, by means of concentration, exclusion, or stabilization of disordered confirmations.

In the experiments, we use full-length α Syn (FL- α Syn) and two truncated variants to better understand which protein domains are responsible for specific behaviors. Three different coacervates were investigated as model condensates, and their selection was guided by well-defined coacervate models reported in literature of ribonucleoprotein (RNP) granules containing RNA and arginine-rich peptides (38-40), heterotypic condensates containing unstructured polypeptides (41), and active droplets containing small molecules (42, 43). We show that depending on the composition of the condensed phase, amyloidogenic protein can partition into the droplets, remain excluded, or accumulate at the interface. We find that FL-aSyn accumulates and aggregates preferentially inside two of the complex coacervate droplets and accumulates and aggregates at the interface of another type. Accumulation of FL- α Syn either inside or at the interface of coacervates always leads to enhanced aggregation compared to a homogeneous solution. Truncated variants of aSyn were typically excluded from the coacervate droplets and aggregated at comparable rate or slower than in homogeneous solution. The shortest variant, which only contains the β sheet-forming region and which normally has the fastest maximum rate of aggregation, is also accumulated inside two coacervates but aggregates notably slower than in solution. This demonstrates that sequestration of amyloidogenic proteins inside condensates can speed up aggregation by enhancing local concentrations in some condensates but slow it down in others due to a stabilization of the monomeric form of the protein.

in further experiments unless stated otherwise (40 µM for FL-αSyn and α Syn-108 and 160 μ M for NACore).

RESULTS

Properties of selected α Syn variants

We selected three α Syn variants with different net charge and length

of the intrinsically disordered region (Fig. 1A): wild-type FL-αSyn,

a truncated variant depleted of the negatively charged, disordered C-terminal domain (α Syn-108), and a relatively hydrophobic short

peptide from the non-amyloid-β component of the protein, which is

the part that is responsible for β -sheet formation in aggregates (NACore

peptide, αSyn-68-78). While having different physicochemical prop-

erties, these variants are all able to aggregate into amyloids (Fig. 1B),

and the kinetics of their aggregation can be described by a classical

nucleation and growth model (44), including primary and second-

ary nucleation (Fig. 1C). All variants form fibrillar aggregates, as con-

firmed by transmission electron microscopy (TEM) (Fig. 1D). For each variant, we determined the concentration at which complete ag-

gregation, defined as reaching plateau of thioflavin T (ThT) intensity, was reached in less than 5 days, and these concentrations were used

Partitioning of a Syn into coacervate droplets

We investigated the localization of selected α Syn variants in three coacervate systems, which may serve as very basic models of biological condensates: (i) (RRASL)₃ peptide/polyuridylic acid (RP3/polyU) (38, 45), (ii) poly-D,L-lysine/poly-D,L-glutamate (pLys/pGlu) (41, 46), and (iii) poly-L-lysine/adenosine triphosphate (pLys/ATP) (Fig. 2A) (42, 43). All these systems phase separate upon mixing (poly)cationic with a (poly)anionic components and form micrometer-size droplets that fuse into larger droplets over time, but remain liquid over the course of several days. In addition, all droplets have been shown to take up or exclude a wide range of biomolecules and complexes (38, 47-49), and are thus expected to influence the aggregation of aSyn. RP3/polyU coacervates (system i) have been suggested as a model for RNP granules, which are typically composed of RNA and arginine-rich peptides or proteins (50). Our model system (i) contains a relatively short cationic component (RP3 peptide) and a long anionic component (polyU RNA) and is thus expected to have a negative surface potential (51) and interact weakly with negatively charged guest molecules. pLys/pGlu coacervates (system ii) are composed of large, unstructured cationic and anionic peptides and have relevance for biomolecular condensates whose formation is known to be driven by proteins bearing charge patches, such as LAF-1 and Ddx4 (20, 52). Last, pLys/ATP coacervates (system iii), which contain the biologically relevant small molecule and hydrotrope ATP (53), are of relevance within the context of biomolecular condensates that bind ATP to tune their properties and composition (54, 55). Moreover, these coacervates have been used as active droplet mimics (43), which makes understanding their influence on protein aggregation and fibrillization also relevant from a protocell perspective. Unlike RP3/polyU coacervates, pLys/ATP coacervates contain a long cationic and a short anionic component, typically resulting in a positive surface potential when prepared at equal charge (56), and a strong interaction with negatively charged guest molecules. These three different model systems were selected to cover a variety of different condensate compositions and properties, for which we expected different interactions with the α Syn variants. Together, these model systems may yield generalizable, physicochemical insight into the influence of condensed aqueous droplets on protein aggregation.



Fig. 1. Amyloidogenic *α***Syn variants used in this study**. (**A**) Variants of *α*Syn used in the study and predicted disorder along the protein chain (PrDOS, in gray) (*75*) and distribution of charged residues (in color). For comparison of predicted disorder using different online tools, please see the Supplementary Materials and fig. S1. (**B**) Aggregation traces (normalized ThT fluorescence intensity) for various *α*Syn variants (recorded for 40 μM concentration of FL-*α*Syn and *α*Syn-108 and for 160 μM concentration of NACore), without coacervates. (**C**) Schematic depiction of the basic protein aggregation cycle model used in this study. (**D**) TEM images of fibrils formed by studied variants. Scale bars, 200 nm. (**E**) FL-*α*Syn conformation when bound to lipids [left, Protein Data Bank (PDB) ID: 1XQ8] and stacked in amyloid fibrils (right, PDB ID: 2NOA); relevant residues are indicated.

To determine whether a Syn and its variants partition into coacervate droplets or remain excluded, Alexa Fluor 647-labeled variants of α Syn were added to the coacervate emulsions and placed in a chambered glass slide to visualize them with confocal microscopy. Distinct partitioning could be observed for different combinations of labeled protein and coacervates (Fig. 2, B and C). FL-αSyn accumulated at the interface of the coacervate droplets and the solution phase, which was particularly visible for pLys/pGlu and pLys/ATP systems (Fig. 2D). For all coacervates, the average fluorescence inside the droplets (excluding the interface) was higher than in the surrounding dilute phase (Fig. 2C). The truncated variant αSyn-108 remained excluded from all coacervate droplets and particularly for pLys/pGlu system for which the ratio of concentration inside/outside was lowest (Fig. 2D). Last, the NACore fragment partitioned into RP3/ polyU droplets, and very weakly into pLys/ATP droplets, but remained excluded from coacervates formed by pLys/pGlu.

The tendency of $FL-\alpha Syn$ to localize to the interface of coacervate droplets may stem from the fact that its disordered chain includes

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both charged/hydrophilic and hydrophobic regions. Both large negatively charged RNAs and small hydrophobic dyes have been found to partition into pLys/ATP coacervates (42, 43). However, in an amphiphilic molecule, such as FL-aSyn, not all regions are preferentially taken up by the coacervate environment, resulting in a strong localization at the interface. Previous studies on partially unfolded proteins have also shown similar interfacial localization (57). Interfacial localization seems to be strongest for coacervates with relatively low molecular weight (MW) anionic components, such as ATP in pLys/ATP and pGlu in pLys/pGlu. Displacing these small anions with FL- α Syn in the coacervates results in a larger gain in entropy than displacing the large polyU in RP3/polyU coacervates. Some uptake of FL- α Syn inside the coacervates is possible for all coacervates tested (Fig. 2C) and can be explained by an overall favorable interaction between FL-aSyn and one of the components in the coacervates (57–60). The negatively charged C-terminal domain appears crucial for both the uptake and the interfacial localization: The truncated αSyn-108 was systematically excluded from the droplets. The NACore

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Fig. 2. Coacervate systems and interactions with α Syn variants. (A) Schematic depiction of coacervate systems used in the study. (B) Confocal microscopy images of coacervate systems with labeled α Syn variants [Alexa Fluor 647–labeled S9C-FL- α Syn and S9C- α Syn-108 and carboxytetramethylrhodamine (TAMRA)–labeled NACore]. Scale bar, 20 μ m. (C) Partition coefficient and transfer free energy (dilute phase-coacervate) determined from microscopy experiments. (D) Gray value profiles (angular averaging) of coacervate droplets from selected systems.

does not show any interfacial localization, presumably because it only contains the relatively hydrophobic core region of the protein. Instead, it is either sequestered and distributed homogeneously inside the coacervates or excluded, reflecting the favorable interactions between NACore and RP3, and the unfavorable interactions with pGlu most likely.

We note that the quantification of local concentrations by fluorescence (e.g., Fig. 2) is dependent on the assumption that the fluorescence intensity of a dye is proportional to its concentration under all different circumstances. High concentrations of labeled α Syn, particularly at the interface of the droplets, could potentially result in self-quenching of the dye and underestimating of the local α Syn concentration. However, the Alexa Fluor dyes we used here were selected for their resistance to self-quenching and photobleaching. Moreover, the experiments shown in Fig. 2, from which we extract local concentrations for later use, were always performed at very low concentrations of labeled α Syn (1 µM, which is orders of magnitude lower than typical aggregation conditions), and therefore,

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we assume that our estimates of the local α Syn concentrations inside the droplets was not affected by these effects.

Aggregation kinetics in the presence of coacervates

Upon phase separation, most coacervate forming material (peptides and RNA in our case) is condensed into droplets, which are in equilibrium with the surrounding diluted phase (supernatant). We used ThT assay (61, 62) to study the aggregation kinetics of the α Syn variants in the presence of the coacervate droplets. We verified that addition of the different α Syn variants does not affect the stability of the coacervate droplets (fig. S2). The supernatant usually contains very low but not negligible concentrations of the coacervate components. To distinguish the influence of coacervate droplets from the soluble components in the supernatant, we performed experiments in the presence of droplets and control experiments with only the supernatant (separated from coacervate droplets by centrifugation) for each coacervate system (an example for RP3/polyU is shown in Fig. 3A, and full traces are shown in fig. S3). In addition, a reference



Fig. 3. FL-*α***Syn aggregation in the presence of coacervates.** (**A**) Aggregation traces for FL-*α*Syn: without coacervates (reference), with RP3/polyU supernatant, and with RP3/polyU coacervates. (**B**) TEM images of aggregates formed in the presence of RP3/polyU coacervates. Scale bars, 1 µm (images on the left side) and 200 nm (images on the right side).

experiment was performed without any droplets or soluble coacervate components. We note that studies on α Syn aggregation use various names and conditions for the control experiments. In our study, the reference experiment performed without any coacervate material reflects the "standard" aggregation kinetics of α Syn variants, and it can be used to obtain information about the general influence of the coacervate components (both in solution and in droplets) on the aggregation kinetics. The control experiments are performed with the coacervate components in solution at saturation concentrations, and they allow separating the influence of the coacervate droplets from the influence of the coacervate components present in the diluted phase (in solution) by comparing them to the kinetics obtained in the presence of coacervate droplets.

Figure 3A shows a typical set of aggregation traces for one coacervate system and for one aSyn variant (FL-aSyn). It includes data for the reference sample without any coacervate material (blue traces), for the supernatant sample with diluted components showing faster aggregation (yellow traces), and for the sample with coacervate droplets (red traces). Before analyzing the kinetics of aggregation further, we used TEM to determine whether the amyloid fibrils formed in the presence of droplets appear similar to the fibrils formed in solution. Figure 3B shows TEM images of FL-aSyn fibrils present inside and at the interface of coacervate droplets. The fibrils at the interface appear to be aligned parallel to the interface, forming an apparent aggregation shell that is not completely disordered. Some fibrils protrude into the surrounding solution. These results indicate that the fibrils can nucleate and grow in coacervate droplets. Moreover, analysis of the fibrils shown in Fig. 3B revealed that there is no significant difference in thickness between these and fibrils formed in solution (fig. S4). Last, we also purified the incubated samples of α Syn with coacervates by dissolving the coacervates at high salt concentration, depositing the fibrils on a TEM grid and rinsing the grid with Milli-Q water. All combinations of a Syn variants and coacervates show the same fibril appearance as in samples without coacervates (Fig. 3 and fig. S4B).

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To elucidate the role of condensates on the kinetics of aggregation, we plotted the distributions of both the lag times (t_{lag} , which is predominantly determined by the primary nucleation rate) and the maximum slopes of the ThT-based aggregation curves (v_{max} , which is mostly determined by the elongation and secondary nucleation rate). As can be seen in Fig. 4, the presence of each of the coacervates resulted in faster aggregation for FL- α Syn with smaller spread of the nucleation times (fig. S5), although the localization of FL- α Syn in these coacervates is not identical: In the case of RP3/polyU, we observed a homogeneous distribution inside the droplets, while in the other two cases we observed a strong interfacial adsorption (Fig. 1B).

The presence of RP3/polyU droplets mostly affects the lag phase of α Syn aggregation. With these droplets, the lag phase was 4 times shorter than in controls with only supernatant and 10 times faster than in reference solution, indicating that the amyloid nucleation rate was enhanced by the droplets. On the other hand, the maximum αSyn aggregation rate in the presence of RP3/polyU droplets is comparable to the control samples with the RP3/polyU supernatant and to the reference sample without any coacervate material. Since these droplets concentrate FL- α Syn (Fig. 2), it was expected that the growth rate inside the droplets is enhanced, as we discuss below. The fact that we observe a comparable growth rate despite a higher local concentration indicates that FL-aSyn is less aggregation prone inside RP3/polyU coacervates. Possibly, one of the coacervate components can bind weakly to FL-aSyn monomers, oligomers, or short fibrils and slow down amyloid growth. Nevertheless, the presence of droplets accelerated aggregation overall, in the sense that the time to complete aggregation was reduced, due to a shorter lag phase.

Different behavior was observed for FL- α Syn aggregating in the presence of pLys/pGlu and pLys/ATP coacervates. FL- α Syn in both the supernatant and the coacervate sample showed a faster onset of aggregation. Unlike in the case of RP3/polyU, the growth phase in the presence of pLys/pGlu and pLys/ATP droplets was significantly faster than in the reference sample and also faster than in the presence of

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Fig. 4. Analysis of aggregation kinetics. (**A**) Distribution of the lag times (t_{lag}) for all protein variants and all coacervate systems (s, supernatant; c, coacervate) and for the reference sample. Symbols at the top indicate localization of the corresponding variant as determined using fluorescence microscopy. Differences between samples were tested for statistical significance (Student's *t* test) in coacervate droplet–supernatant control pairs. "ns" indicates values above 0.05, single asterisk indicates $\alpha < 0.05$, double asterisk indicates $\alpha < 0.001$, triple asterisk indicates $\alpha < 0.001$, and quadruple asterisk indicates $\alpha < 0.0001$. Violin plots were prepared using Gaussian kernels with bandwidth determined automatically using the Scott's method; density plots were cut at two bandwidth units past the extreme data points; violins are scaled to have the same area in supernatant-coacervate pairs. (**B**) Distribution of the maximum aggregation rates (v_{max}) for all protein variants and all coacervate systems [colors and symbols as in (A)].

supernatant. This may be linked to the fact that pLys/pGlu and pLys/ ATP systems seem to have a tendency to accumulate monomeric FL- α Syn at the interface of the droplets, thereby providing an alternative aggregation pathway, as we will discuss below. RP3/polyU system mostly accumulates FL- α Syn monomers inside and increases in concentration, and altered environment affects mostly the primary nucleation rate.

The same experiments were performed for the α Syn-108 variant and the NACore peptide (Fig. 4 and figs. S3 and S6). The aggregation behavior of the shorter variants was fundamentally different from FL- α Syn. While the samples incubated in the supernatant aggregated at comparable rate to the references for both α Syn-108 and NACore, the presence of droplets resulted in slower aggregation. Large spread of the aggregation parameters for α Syn-108 made it difficult to assess the significance of the effect, but for NACore, it was clear that the peptide in samples with coacervates aggregated significantly slower than peptide in both the supernatant and reference samples. The presence of all types of droplets resulted in lag times that were longer than in the corresponding supernatant sample, although the spread was typically very large, which made only the difference in the lag time of pLys/pGlu droplets statistically significant. The presence of RP3/polyU droplets also resulted in significantly slower maximum aggregation rates than in the supernatant, suggesting that the NACore peptides sequestered inside these droplets are less aggregation prone, for reasons we discuss below.

Last, to show that the observed divergent influence of coacervate droplets on protein localization and aggregation kinetics is not limited to α Syn (and its variants), we repeated these experiments (shown in Figs. 2 to 4) with insulin, which is known to form amyloid-like aggregates upon incubation (Supplementary Materials and figs. S7 to S10). We found that insulin could also be sequestered inside some coacervate droplets (RP3/polyU and pLys/ATP), excluded from others (pLys/pGlu 1:4 and 1:2), and accumulated at the interface in one case (pLys/pGlu 1:1) (fig. S10). We observed that aggregation of

insulin was enhanced in the presence of RP3/polyU and pLys/ATP coacervates, while for pLys/pGlu coacervates at all charge ratios aggregation was suppressed (figs. S8 and S9).

Spatiotemporal mapping of the aggregation process

To find out whether the faster and slower aggregation is related to the location where aggregation takes place, as the partitioning data (Fig. 2B) seem to suggest, and to follow the spatiotemporal distribution of aggregates in the presence of coacervates, we developed an intramolecular FL- α Syn fluorescence resonance energy transfer (FRET) probe, similar to the probe used before to study conformations of α Syn at a single molecule level (*63, 64*). The probe includes two fluorescent dyes close to the region responsible for β -sheet formation: Alexa Fluor 546 and Alexa Fluor 647 (Fig. 5A). Upon aggregation, the dyes become fixed close to each other, which results in enhanced energy transfer (fig. S11). Experiments were performed in a similar fashion to the partitioning experiments, but the images were collected for several days and the samples were incubated at 37°C. Collected fluorescence intensity images were used to create FRET efficiency maps, by calculating FRET efficiency for each pixel separately (Fig. 5B).

We observed distinct behavior for the different coacervates. In the case of RP3/polyU, the FRET signal increased throughout the entire coacervate droplet directly from the beginning and was slightly enhanced at the interface, while it remained constant and low outside the droplets (Fig. 5, C and D). After 60 hours of incubation, the FRET signal increased inside the droplets, which suggests that the aggregates are formed inside the droplets or at the interface and then move to the interior (Fig. 5B). For pLys/pGlu, the FRET efficiency was higher at the interface than in the surrounding solution, and also higher than inside the droplets, even after 60 hours. This suggests a more compact conformation and high concentration of FL- α Syn at the interface, potentially promoting faster nucleation into fibrils, which form at the surface of the droplets but do not move



Fig. 5. Aggregation monitored by FRET. (**A**) Positions of FRET labels in the FL- α Syn chain (PDB ID: 2NOA). (**B**) FRET maps of coacervate samples incubated with FRET-labeled FL- α Syn. Scale bar, 20 μ m. Insets show three times enlarged part of the image at 60 hours; the experiment was performed in the presence of the FRET probe and 10 μ M (RP3/polyU and pLys/ATP) or 40 μ M (pLys/pGlu) concentration of nonlabeled FL- α Syn. (**C**) Changes in FRET intensity in different areas of the coacervate systems over time. (**D**) FRET intensity radial profiles for coacervate droplets after 60 hours of incubation (distance from the droplet center normalized by the droplet diameter, angular averaging).

toward the interior. In the case of pLys/ATP, aggregation starts immediately, with practically no lag phase (as also observed in the kinetic experiments; Fig. 4). The highest FRET efficiency was observed inside the coacervate droplets, and the droplets seem to maturate over time, losing their liquid properties. Therefore, for the pLys/ATP system, we also expect that the aggregates nucleate within the coacervate droplets, despite the enhanced concentration of monomeric α Syn at the interface, as observed in the partitioning experiments (Fig. 2B). In FRET experiments, no accumulation of aggregated FL- α Syn could be observed at the interface of pLys/ATP coacervates. Presumably, even if aggregates nucleate at the interface, they immediately move to the interior of the droplets.

Similar observations were made using unlabeled $FL-\alpha$ Syn and ThT as a dye to stain the fibrils under the microscope (fig. S12A). In this case, the presence of protein aggregates is simply indicated by high ThT fluorescence intensity. Direct analysis of the fluorescence intensities is complicated in this case, because free ThT also partitions into coacervates (fig. S12B). Nevertheless, we could observe substantial increases in fluorescence intensity upon aggregation. In the presence of RP3/polyU and pLys/ATP droplets, aggregates were formed within the droplets, resulting in irregular solid-like particles. In the presence of pLys/pGlu coacervates, the highest fluorescence intensity was observed for the coacervate interface, suggesting that aggregation is promoted by droplet interface, as observed also for the FRET probe.

A kinetic model of protein aggregation accelerated or suppressed by condensates

Our microscopy experiments suggest that the presence of the coacervate droplets can affect the α Syn aggregation process either through partitioning of the protein into coacervate droplets or through α Syn/coacervate interface interactions. To prove that these interactions can also be the reason of differences observed in the kinetics of the aggregation process, we developed and fitted kinetic models to our experimental data (Fig. 5). Two separate models were developed to match the observations in Fig. 2: (i) α Syn is excluded from or sequestered by the droplets, and aggregation can take place both inside and outside the coacervate droplets (Fig. 5A); (ii) α Syn is localized at the coacervate interface, and heterogeneous nucleation followed by further aggregation can take place at the interface (Fig. 5B).

The kinetic models are based on the three-step nucleationelongation-secondary nucleation model proposed by Ferrone et al. (65). Formation of protein amyloid fibrils (a) is initiated by nucleation, (b) the increase of aggregate mass is mostly caused by elongation of fibrils, while (c) their presence also has an autocatalytic effect on formation of new fibrils through secondary nucleation. In the simple case of aggregation in a homogeneous solution, it has been shown that these three processes (with three rate constants) are sufficient to describe the aggregation kinetics of various amyloidogenic proteins (66). We developed two new models that describe fibril formation via the same three steps (a, b, and c), but now (i) taking place both outside and inside coacervate droplets or (ii) taking place both outside and at the interface of coacervate droplets. Transport equations for protein monomers ensure coupling between the aggregation loci. A detailed description of the differential equations for both models can be found in the Supplementary Materials.

In the first case of exclusion or sequestration, we assumed that α Syn monomers are freely exchanged between the dilute phase (which we call supernatant hereafter) that surrounds the coacervate

droplets, and that the exchange of α Syn between the supernatant and the condensed phase is much faster than the aggregation process itself. Consequently, local α Syn concentrations are always equilibrated (i.e., the partition coefficient, as determined in Fig. 2C, is constant). Aggregation of α Syn can occur both in the supernatant and in the coacervate droplets, and the rate constants of each step of the aggregation process (Fig. 1C) can be different in both phases, which makes our approach different from previous aggregation models. Kinetic rate constants for the data obtained for α Syn in the supernatant were determined by fitting a simple aggregation model (for a onephase system) and were further used as input for the supernatant phase when fitting parameters for the coacervate-containing samples (Fig. 5C), thereby reducing the number of fit parameters. Last, the fibrils are assumed to be immobile, in accordance with previous models (*30*): Once formed, they remain in the diluted or condensed phase.

Partitioning into the coacervate phase can accelerate aggregation because of increased local aSyn concentrations, but different rate constants for the aggregation process inside the coacervate droplets might mask this effect or further enhance it. In our experiments, we know the local concentration of a Syn from partitioning studies (Fig. 2C) and can therefore deduce the additional influence of the coacervate environment on the rate constants. We note that the fits can only be used to obtain an order-of-magnitude estimate of the rate constants, as it is difficult to determine the separate contributions of each step in the aggregation process from a fit without comparing data collected for different a Syn concentrations. Nevertheless, we found notable differences between the rate constants in the supernatant and the coacervate phase. The fitted aggregation curves for RP3/polyU suggest that the primary nucleation rate of FL- α Syn inside the droplets is higher than in the supernatant, while the elongation and secondary nucleation rates are slightly lower, implying that the RP3/polyU coacervate environment has a destabilizing effect on monomeric FL-αSyn, but a stabilizing effect on oligomers and fibrils, under the assumption that all steps in the aggregation process are not diffusion limited. For NACore, which was also sequestered in RP3/polyU coacervates, we found a similar trend, although the model is not able to capture the very sharp onset of aggregation in some cases (fig. S6B).

The truncated aSyn-108 was excluded from all types of coacervates, and hence, we did not observe any substantial difference in aggregation kinetics. Even in the presence of droplets, the aggregation process takes place predominantly in the supernatant, as there is hardly any α Syn-108 present inside the droplets. Therefore, we could not reliably determine the rate constants of aggregation for α Syn-108 in the coacervates. The same is true for NACore in the presence of pLys/pGlu coacervates and pLys/ATP coacervates where we observed an overall suppressed aggregation, manifested in a longer lag time, but the monomers did not show notable sequestration. On the contrary, they are excluded from pLys/pGlu coacervates and indifferent to pLys/ATP coacervates. Therefore, a simple three-step model as shown in Fig. 5A cannot explain the suppressed aggregation. It is possible that while free peptides were excluded, small oligomers, which form in early stages of the aggregation process, are sequestered by the droplets and prevented from further growth (see also Discussion).

In other cases, we observed strong accumulation of the α Syn variants at the interface of the coacervates, rather than inside, which suggests a different mechanism of aggregation. Therefore, we developed a kinetic model to describe binding of the protein to the interface

and further heterogeneous nucleation and elongation. The simplest model involving only binding (dependent on the number of binding sites and the binding constant) and heterogeneous (primary) nucleation was not able to capture the very rapid aggregation observed in some cases. When we also allowed elongation and secondary nucleation to occur at the interface, we could capture the rapid global aggregation (Fig. 5B). Note that this model assumes that binding to the interface is an equilibrium process, governed by thermodynamics. Although it is known that interfaces can give rise to kinetic trapping of proteins, we expect that the fluid, low-tension interface of a coacervate droplet (67) does not lead to kinetic trapping, which is supported by the fact that α Syn remained mobile at the interface, and we observed no difference in interfacial accumulation of α Syn up to 48 hours after incubation. For pLys/ATP coacervates, the primary nucleation rate constant at the interface is several orders of magnitude higher than in the supernatant, while other rate constants appear the same. This can explain the very rapid onset of aggregation with virtually no lag time for pLys/ATP. For pLys/pGlu coacervates, primary nucleation and elongation seem unchanged, but the secondary nucleation rate constant is significantly faster at the interface, which explains the very rapid increase in ThT fluorescence after a lag phase (Fig. 5B).

To confirm that the coacervate interface is crucial in enhancing the aggregation kinetics, we have performed additional experiments in which we changed the amount of available surface area. In the first experiment, we varied the amount of droplet-forming material. In the second experiment, we centrifuged the coacervate dispersions before adding α Syn, causing the droplets to fuse (and thus reducing the available surface area). In both experiments, we could observe that reducing the droplet surface area resulted in slower aggregation (fig. S13).

DISCUSSION

Our results show that condensates that are composed of nonaggregating material themselves can influence the aggregation of amyloidogenic proteins, such as αSyn and insulin, significantly and in a wide variety of ways. For FL-αSyn, we observed an increase in overall aggregation propensity for all systems. As suggested before by Weber and co-workers (30), this influence can be at least partially caused by higher local concentration of the aggregating protein inside or at the surface of coacervates. However, partition coefficients determined for the studied systems do not seem to explain the difference in the aggregation kinetics, unless we assume different aggregation rate constants inside the coacervate droplets and in the surrounding solution. In particular, we find that elongation and secondary nucleation rate constants are lower inside coacervates than in solution, indicating a partly stabilizing effect of the coacervate environment and a suppressing role in part of the aggregation process. Such differences may be expected, as the more crowded and hydrophobic coacervate environment (29, 68, 69), which is rich in functional groups that can interact with α Syn, affects the protein conformation and its tendency to aggregate. By contrast, our partitioning data combined with FRET microscopy indicate a different mechanism of accelerated aggregation for pLys/ ATP and pLys/pGlu coacervates, which both localized FL-αSyn at their interface, giving rise to surface-bound enhanced primary nucleation (pLys/ATP) and secondary nucleation (pLys/pGlu).

We have thus observed that the coacervate droplet interface can serve as a nucleation site for protein aggregation. Relatively high apparent kinetic rate constants determined for $FL-\alpha$ Syn at the pLys/

pGlu droplet interface may suggest that the coacervate droplets do not only serve as simple heterogeneous catalysis nucleation sites but also provide a distinct physicochemical environment or conformational arrangement, in which protein aggregation is substantially enhanced. It is interesting to note that very recently different behavior has been observed for FL- α Syn under conditions that promote phase separation of FL- α Syn itself (i.e., in the presence of PEG and at high concentrations). Under such conditions, FL- α Syn forms liquid droplets that undergo maturation (a transition into solid aggregates), and this transition was found to be initiated at the center of the droplets, suggesting that FL- α Syn droplets also provide a distinct environment in which the kinetic parameters of aggregation are altered, just like in the case of our pLys/pGlu droplets (70).

Furthermore, our results show that the influence of the coacervate droplets on aggregation kinetics depends on both the coacervate composition and the sequence/length of the aggregating protein. While aggregation of the full-length variant was accelerated in the presence of all coacervate systems, aggregation of the truncated variant, α Syn-108, was not significantly affected. This can be attributed to a different affinity of the full-length and the truncated α Syn variants to the coacervate material, and particularly to the positively charged components. The absence of the negatively charged C-terminal part in α Syn-108 makes this variant slightly positively charged at neutral pH [isoelectric point (pI) = 9.16], while FL- α Syn is strongly negatively charged (pI = 4.67). FL- α Syn has been shown before to aggregate faster in the presence of polycations in solution, and similar acceleration may occur inside coacervates or at their interface (71).

Another interesting observation is that pLys/pGlu and pLys/ATP affect the aggregation process differently, although they both contain pLys. The reason for this difference is the binding strength of the counterions present in these coacervate droplets. ATP has fewer negative charges and binds less strongly to pLys than pGlu, which is evidenced by the lower critical salt concentration of pLys/ATP droplets. As a result, FL- α Syn can displace ATP more easily than pGlu and bind more strongly to pLys. We hypothesize that stronger binding of the negatively charged tail of α Syn makes the protein more prone to aggregation, similar to previous reports (*72, 73*). In addition, the weaker interaction of ATP compared to pGlu leads to a lower viscosity inside the pLys/ATP condensates, which facilitates movement of aggregates and FL- α Syn bound to pLys inside the droplets.

Last, coacervate droplets are also able to slow down aggregation, which was most prominent for NACore peptide (Fig. 4) with RP3/ polyU and pLys/pGlu coacervates. This may be explained by sequestration of free peptides and small oligomers inside the coacervate (Fig. 2), in relatively stable conformation, not prone to rapid aggregation, an effect we also observed to a limited extent and were able to quantify for FL- α Syn in RP3/polyU coacervates (Fig. 6D). In combination with lowered monomer concentration is the surrounding solution, this can result in overall aggregation inhibition [similarly to the sequestration of amyloid- β (1–42) described before (31)]. Unexpectedly, in the case of pLys/pGlu droplets, where labeled NACore peptide remained excluded from the droplets, the aggregation was also slowed down. It is possible that while free peptides were excluded, small oligomers, which form in early stages of the aggregation process, are sequestered by the droplets and prevented from further growth. However, proving this is impracticable, because any action to separate the droplets from the supernatant will most likely disrupt such oligomers.

In conclusion, we show that preexisting liquid condensates can affect protein amyloid formation in vitro, both accelerating and slowing

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Fig. 6. Fitting of the aggregation models. (A) Schematic depiction of the aggregation-in-droplets model. (**B**) Schematic depiction of the interface-aggregation model. (**C**) Fits to the experimental data for RP3/polyU with FL- α Syn for the aggregation-in-droplets model and fits to the experimental data for pLys/pGlu and pLys/ATP with FL- α Syn for the interface-aggregation model. (**D**) Resulting aggregation kinetic rate constants, for the diluted-supernatant phase (s) and for the coacervate/interface phase (c/i), normalized by values for reference sample (without coacervate components). Violin plots were prepared analogously to plots in Fig. 4.

down the reactions. We expect that the same process can happen in living cells, which contain multiple MLOs, formed upon LLPS. By sequestering amyloidogenic proteins, such biological condensates may prevent protein aggregation, but it is also possible that they can function as heterogeneous nucleation sites. This provides a new perspective on the early stages of amyloid formation by α Syn (and protein aggregation in general) in the complex cellular environment.

MATERIALS AND METHODS

Reagents

Poly-L-lysine hydrobromide (MW = 15 to 30 kDa), ATP disodium salt, polyuridylic acid potassium salt, buffers, and ThT were purchased from Sigma-Aldrich. RP3 (RRASLRRASLRRASL-NH₂) and NACore (GAVVTGVTAVA) peptides were purchased from CASLO ApS

(Denmark). Labeled NACore peptide was synthesized on solid phase using the standard Fmoc peptide synthesis strategy. Poly-D,L-lysine hydrobromide (MW = ca. 21 kDa) and poly-D,L-glutamic acid sodium salt (MW = ca. 15 kDa) were purchased from Alamanda Polymers (USA). Alexa Fluor maleimides were purchased from Thermo Fisher Scientific. Poly-L-lysine grafted with PEG (PLL-g-PEG) was purchased from SuSoS AG (Switzerland). All aqueous solutions were filtered before use using Acrodisc 0.2- μ m nylon syringe filters (Sigma-Aldrich) or Pierce cellulose acetate filter spin cups with 0.45- μ m pore size (Thermo Fisher Scientific).

Protein preparation and labeling

Wild-type FL- α Syn, α Syn-108, and the cysteine mutants were expressed and purified as previously described (74). Purified proteins were stored at a concentration of ~200 μ M in 10 mM tris-HCl (pH 7.4)

at -80°C, supplemented with 1 mM dithiothreitol (DTT) for the cysteine mutants. Single labeled proteins were labeled according to the dye manufacturer procedures. For labeling of double-cysteine mutant (42C 90C), the first labeling step (with donor dye) was performed according to the dye manufacturer procedures, using 1:1 protein-to-dye ratio. Subsequently, the protein was incubated with prewashed Activated Thiol Sepharose 4B (Cytiva, USA) for 1 hour, rotating in the dark at 4°C. Next, the resin was washed with several volumes of 10 mM tris-HCl (pH 7.4), followed by elution of singleand double-labeled a Syn using buffer containing 25 mM DTT. Eluted fractions were pooled, concentrated to about 0.5 ml, and desalted. Triple excess of acceptor dye was added to the concentrated protein, and the solution was incubated for 1 hour at room temperature. Unbound dye was removed using Amicon Ultra-4/15 centrifugal filters with suitable molecular weight cut-off (MWCO). Protein solutions were filtered using Pierce cellulose acetate filter spin cups (Thermo Fisher Scientific) before every aggregation kinetic assay, and concentration was determined on the basis of absorbance ($\epsilon = 5600 \text{ M}^{-1} \text{ cm}^{-1}$ for wild-type α Syn and $\epsilon = 1400 \text{ M}^{-1} \text{ cm}^{-1}$ for α Syn-108).

Preparation of modified glass slides

All glass slides used for microscopy were modified according to the following procedure. First, the slide was washed thoroughly with Milli-Q water. Subsequently, the surface intended to be modified was cleaned with oxygen plasma, and a solution (0.01 mg/ml) of PLL-g-PEG in 10 mM Hepes buffer (pH 7.4) was applied on the glass immediately after the plasma treatment. Glass was incubated with the PLL-g-PEG solution for 2 hours at room temperature. Subsequently, it was rinsed three times with 10 mM Hepes buffer (pH 7.4) and three times with Milli-Q water and dried with pressurized air. Modified slides were stored at room temperature and used within 1 week.

Coacervate systems

Unless specified otherwise, the coacervate systems were prepared in 50 mM Hepes buffer (pH 7.4) with 100 mM NaCl and 100μ M EDTA. All coacervate systems were prepared at 1:1 positive to negative charge ratio and total concentration of coacervate-forming material (0.85 mg/ml) [RP3 (0.53 mg/ml) with polyU (0.32 mg/ml), pLys (0.49 mg/ml) with pGlu (0.36 mg/ml), and pLys (0.23 mg/ml) with ATP (0.62 mg/ml)].

Partitioning of labeled proteins

Localization of labeled proteins was studied using confocal microscopy. A Leica SP8x confocal microscope equipped with ×40 magnification water-immersion objective was used. Samples were placed in 18-well chambered glass coverslips (Ibidi GmbH, Germany), previously modified with PLL-g-PEG. Partition coefficients were determined by calculating ratio of fluorescence intensity in the condensed phase to fluorescence intensity in the outer phase (average intensity values from at least 10 droplets and from outer phase of similar area were used). Background signal of coacervate sample without labeled protein was subtracted separately for condensed and supernatant.

ThT aggregation kinetic assays

To estimate the aggregation kinetic parameters, we have performed standard ThT aggregation assays. Upon binding to β sheets, ThT fluorescence intensity increases by several orders of magnitude and the changes of fluorescence in the solutions of aggregating protein are proportional to the amount of aggregate formed ([*M*]).

Aggregation assays were performed under following conditions: 50 mM Hepes (pH 7.4), 100 mM NaCl, 100 μ M EDTA, 20 μ M ThT, and 40 μ M FL- α Syn or α Syn-108, or 160 μ M NACore. All aggregation assays were performed in nonbinding 384-well plates with black walls (Greiner Bio-One GmbH, Austria) at 37°C. To prevent evaporation, wells in the two outer rows were always filled with water and the plate was sealed with film. Measurements were performed using a Tecan Spark or Tecan Infinite M200 microplate reader. Fluorescence intensity was recorded every 12 min using bottom readout with continuous linear shaking in between. Excitation and emission wavelength range was controlled using monochromators for Tecan Infinite M200 (440 nm with 9-nm bandwidth and 480 nm with 20-nm bandwidth, respectively) or filters for Tecan Spark (430 nm with 20-nm bandwidth and 460 nm with 20-nm bandwidth, respectively).

To extract the basic kinetic parameters (t_{lag} and v_{max}) from the ThT fluorescence traces, we fitted simple aggregation model (as described in the Supplementary Materials) and used the maximum slope of the curve as v_{max} , and the intersection of line going through the max slope point and the baseline was used as t_{lag} (see inset in Fig. 3A).

Preparation of samples and TEM

Samples after the ThT aggregation kinetic assay in 384-well plates were used for electron microscopy experiments. To dissolve the coacervate material and separate the aSyn aggregates, sodium chloride solution was added to the selected wells, to a final concentration of 300 mM sodium chloride. After incubation for 5 min at room temperature, plate was centrifuged for 10 min at 1000 rcf. Subsequently, solution was gently collected from the selected wells and 50 µl of Milli-Q water was added. The plate was centrifuged again with the same settings, and again, solution was gently collected from the selected wells. Any precipitate from the selected wells was resuspended in 20 μ l of Milli-Q water, and subsequently, 2 μ l of the suspension was transferred onto a TEM grid (EM-Tec formvar carbon support film on copper, 300 square mesh, Micro to Nano, The Netherlands). Samples were blotted with filter paper, stained with 1.5 µl of 2% (w/w) sodium phosphotungstate solution (adjusted to pH 7.4), washed with 2 µl of water, and left to dry overnight. Imaging was performed using JEOL JEM-1400 FLASH.

Intramolecular FRET experiments

FRET experiment was performed using a Leica SP8x confocal microscope equipped with ×40 magnification water-immersion objective. Samples were placed in 18-well chambered glass coverslips (Ibidi GmbH, Germany), previously modified with PLL-g-PEG, and the whole setup was incubated at 37°C during the experiment. FRET probe was added at 0.01 ratio to the nonlabeled FL- α Syn (0.1 or 0.4 μ M and 10 or 40 μ M, respectively), and other components remained the same as for the ThT aggregation kinetic assay. Samples were excited at 488 nm, and the emission was recorded at 515 to 530 nm for the donor and 590 to 610 nm for the acceptor. Fluorescence intensity images were saved in 8-bit 512 × 512 pixel format. FRET value was calculated for each pixel using the following formula

 $FRET = \frac{acceptor channel intensity}{donor channel intensity + acceptor channel intensity}$

The FRET value was not determined if intensities for both channels were lower than 2, which is close to the detector dark count. A 512×512 array of FRET intensities was further converted into an

8-bit 512 × 512 image and visualized using custom hue/brightness two-dimensional color map. Hue corresponds to the FRET value scaled from 0.5 to 1 (pixels with values below 0.5 have the same hue as pixel with FRET equal 0.5). Brightness is proportional to the sum of fluorescence intensity for both channels, scaled from 0 to the value for the 95th percentile in the image collected after 60 hours (pixels with higher intensity have the same "max" brightness).

The FRET experiment in bulk (fig. S3) was performed using $0.4 \,\mu$ M concentration of the FRET probe and 40 μ M concentration of nonlabeled FL- α Syn and incubated at 37°C in an Eppendorf tube. Fluorescence spectra were measured using a JASCO FP-8300ST spectrofluorometer.

Statistical analysis

Microscopy images were analyzed using FIJI distribution of ImageJ. Error bars and error ranges of transfer energies and FRET plots were determined using SDs of pixel intensity values within selected range. Plots in Fig. 5C were prepared by manually selecting parts of the image. Plots in Figs. 2D and 5D were prepared using radial profile angle plugin for ImageJ. Violin plots were prepared according to the description under Fig. 4. Fitting of the aggregation kinetic models in Fig. 6C was performed using basinhopping function from scipy.optimize library in python.

SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at https://science.org/doi/10.1126/ sciadv.abq6495

View/request a protocol for this paper from Bio-protocol.

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