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Recombinant silk protein condensates show widely different properties depending on the sample background†

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There is an increasing understanding that condensation is a crucial intermediate step in the assembly of biological materials and for a multitude of cellular processes. To apply and to understand these mechanisms, *in vitro* biophysical characterisation techniques are central. The formation and biophysical properties of protein condensates depend on a multitude of factors, such as protein concentration, pH, temperature, salt concentration, and presence of other biomolecules as well as protein purification and storage conditions. Here we show how critical the procedures for preparing protein samples for *in vitro* studies are. We compare two purification methods of the recombinant spider silk protein CBM-AQ12-CBM and study the effect of background molecules, such as DNA, on the formation and properties of the condensates. We characterize the condensates using aggregation induced emitters (AIEs), coalescence studies, and micropipette aspiration. The condensed sample containing background molecules exhibit a lower threshold concentration for condensate formation accompanied by a lower surface tension and longer coalescence time when compared to the pure protein condensates. Furthermore, the partitioning of small AIEs is enhanced in the presence of background molecules. Our results highlight that the purification method and remaining background molecules strongly affect the biophysical properties of spider silk condensates. Using the acquired knowledge about spider silk protein purification we derive guidelines for reproducible condensate formation that will foster the use of spider silk proteins as adhesives or carriers for biomedical applications.

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1. Introduction

The need for sustainable and environmentally friendly biomaterials is growing. Protein-based materials are a promising solution to address this need.^{1,2} At the same time, the role of condensates in biological systems has gained increasing attention since the discovery of membrane-less organelles.^{3,4}

Especially in the field of biological materials, studies on biomolecular condensates have enhanced the understanding of underlying molecular mechanisms.^{5,6} Among others, condensates can be observed in the formation of mussel byssus threads and the formation of the squid beak.^{6,7} Condensation, also called coacervation, is also proposed to be an important step in the assembly of fibers from spider silk protein. Already inside the silk glands of spiders, it is assumed that condensation occurs due to increased protein concentration and a change in the pH and salt conditions.⁸ Likewise, the artificial production of spider silk fibers from recombinantly produced protein has been linked to condensation, which likely needs to occur before fiber pulling.^{9,10} Recombinant spider silk proteins have been largely studied in the context of materials and biomedical applications.^{11–14} Condensates are a crucial building block in the assembly process of these artificial biomaterials.^{9,15,16} This highlights the requirement for reproducible and large-scale protein condensation in the production process of protein-based materials. However, while providing a viable approach to the manufacturing of sustainable and

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† Electronic supplementary information (ESI) available: Fig. S1: protein sequence for CBM-AQ12-CBM; Fig. S2: SDS-PAGE of CBM-AQ12-CBM and BG lysate; Fig. S3: fluorescence microscopy images with free eGFP; Fig. S4: AIE structures; Fig. S5: inverse capillary velocity (ICV) analysis; Fig. S6: FTIR analysis; Fig. S7: light microscopy images with ssDNA from salmon testes; Table S1: overview of the values for the micropipette aspiration; Video S1: micropipette aspiration of IMAC silk; Video S2: HT silk coalescence; Video S3: IMAC silk + BG lysate coalescence; Video S4: IMAC silk coalescence. See DOI: <https://doi.org/10.1039/d4tb01422g>



biocompatible materials, the use of biomolecular condensates is challenging.

Biomolecular condensates are supramolecular assemblies that result from liquid–liquid phase separation (LLPS). This creates a dense condensate phase, containing a high biomolecule concentration. The surrounding dilute phase contains a low amount of the biomolecule. Weak interactions between the proteins are the driving force for condensation.^{17,18} Typically, condensates behave in a liquid-like manner, which can be characterized by their tendency to coalesce. Proteins that undergo self-condensation often contain highly repetitive, intrinsically disordered regions (IDRs).^{19,20} For spider silk proteins in condensates, it has been shown that their secondary structure is rich in α -helices which can transform into β -sheets when fibers are pulled from the solution.^{9,21} Furthermore, condensates provide a confined space *e.g.*, for adhesives in the mussel's byssus threads.²² At the same time, condensates show an exceptionally low surface tension which seems to be beneficial for substrate infiltration.^{23,24}

Condensates are metastable and known to be highly influenced by sample history and preparation.^{25,26} For example, pH, salt conditions, and temperature affect the concentration at which LLPS occurs.^{25,27,28} Additionally, salts (*e.g.* Na_2HPO_4) affect the state of condensates by transforming previous liquid-like condensates, that can coalesce, into solid-like condensates.^{9,27} The significance of these aspects becomes especially evident when comparing biomolecular condensates *in vivo* and *in vitro*. Naturally, condensates are influenced by a vast number of other components, that are present inside cells.^{29,30} For example, RNA and DNA are known to be able to form complex condensates with certain proteins.^{31,32} By varying RNA and protein ratios, hollow, vesicle-like condensates can be generated.³³ However, it is challenging to recreate these conditions when utilizing protein condensates for materials applications. In these cases, starting from a more simplified, pure system followed by a step by step understanding of the influence of specific background molecules is desirable. RNA and DNA are also common background molecules during recombinant protein production.^{34,35} The purification process of proteins can influence all these variables. By alternating the purification process, different purities and backgrounds can be achieved. However, the use of condensates in large-scale production of biomaterials requires careful balancing of factors such as protein purity, yield, and production costs, as well as factors influencing the biophysical properties of condensates.

This work sheds light on the importance of the purification protocol, and the presence of background molecules for the reproducible condensation of the recombinant spider silk fusion protein CBM-AQ12-CBM. The pure silk protein allows us to engineer a more simplified *in vitro* model system. The presence of the background molecules in the silk protein resembles crowded environments, similar to those observed *in vivo*. We qualitatively analyze the effect of background molecules on the condensation of spider silk proteins by concentrating experiments and light microscopy. Utilizing coalescence studies and micropipette aspiration of condensates,

we demonstrate that spider silk condensates exhibit different biophysical properties, depending on the presence of background molecules in the protein solution. Background molecules such as DNA and RNA reduce the concentration at which LLPS occurs by 75%. Silk protein condensates that are formed in the presence of background molecules frequently show bursting behavior and have a reduced surface tension, indicating the presence of a shell-like assembly at the condensate's surface. In contrast, we did not observe bursting in the simplified model system of pure silk protein condensates, which exhibit a higher surface tension, indicating the absence of shell-like structures. Furthermore, in the absence of background molecules the silk protein condensates are more selective to the uptake of small molecules such as aggregation induced emitters (AIEs). This work highlights the importance of sample history, purification, and preparation when characterizing the condensation properties of proteins. Our example focuses on silk protein assembly, but similar mechanisms are likely to apply to IDR containing proteins more generally.

2. Results & discussion

Protocol for reproducible silk protein production and condensation

We established a protocol to produce the spider silk protein CBM-AQ12-CBM (Fig. S1, ESI[†]) with reproducible concentration and properties (Fig. 1). Consistent production results enable controlled condensation. In short, the protocol involves recombinant protein production in *E. coli*, cell lysis by sonication (Fig. S2, ESI[†]), protein purification and desalting to deionized water, and a final concentrating step with centrifugal concentrators. Depending on the purification method used, we either have background molecules present imitating a crowded *in vivo* environment or create a simplified *in vitro* model system. The purification method developed here always involves a heat treatment for 30 min at 70 °C. The silk protein CBM-AQ12-CBM is stable at this temperature and does not denature. Thus, heat treatment is a fast and easy purification step, especially for large amounts of silk protein. However, after heat treatment, background molecules, such as DNA, RNA and other heat stable proteins are present in the protein solution. After heat treatment and desalting additional His-tag immobilized metal affinity chromatography (IMAC) can be carried out to receive a pure spider silk protein solution.

Quantification of background molecules in silk protein solution

Heat treatment (HT) purification for 30 min at 70 °C results in a slightly yellowish-colored heat treated protein solution (called HT silk), while HT followed by IMAC purification leads to a clear, transparent protein solution (called IMAC silk) (Fig. 2(a) and (b)). Sodium dodecyl sulfate–polyacrylamide gel (SDS–PAGE) of purified protein samples before and after desalting shows a strong band around 85 kDa corresponding to the CBM-AQ12-CBM spider silk protein (Fig. 2(c)). Several faint bands



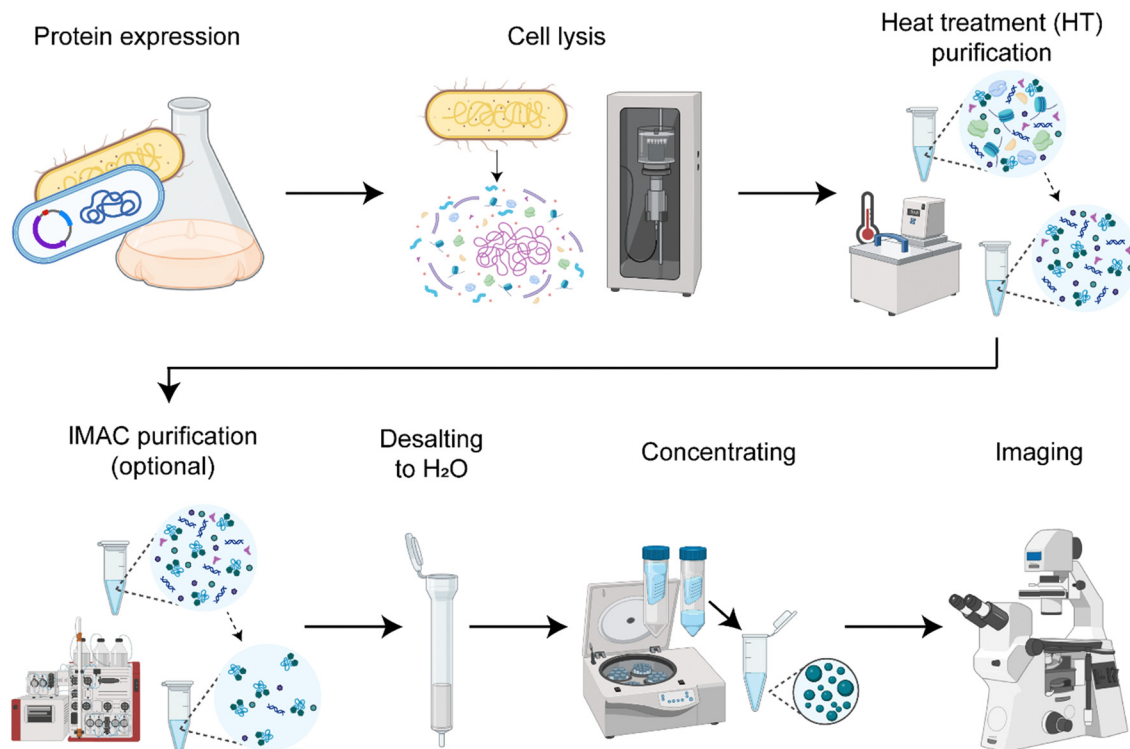


Fig. 1 Protocol for the recombinant production, purification and concentrating of the spider silk protein CBM-AQ12-CBM. Expression takes place in *E. coli* and cell lysis is done by sonicating. Protein purification involves a 30 min heat treatment (HT) step at 70 °C and optional His-tag immobilized metal affinity chromatography (IMAC). Proteins are desalted into water, concentrated with Vivaspin centrifugal concentrators (30 kDa cutoff, Sartorius) until LLPS occurs and imaged with light microscopy.

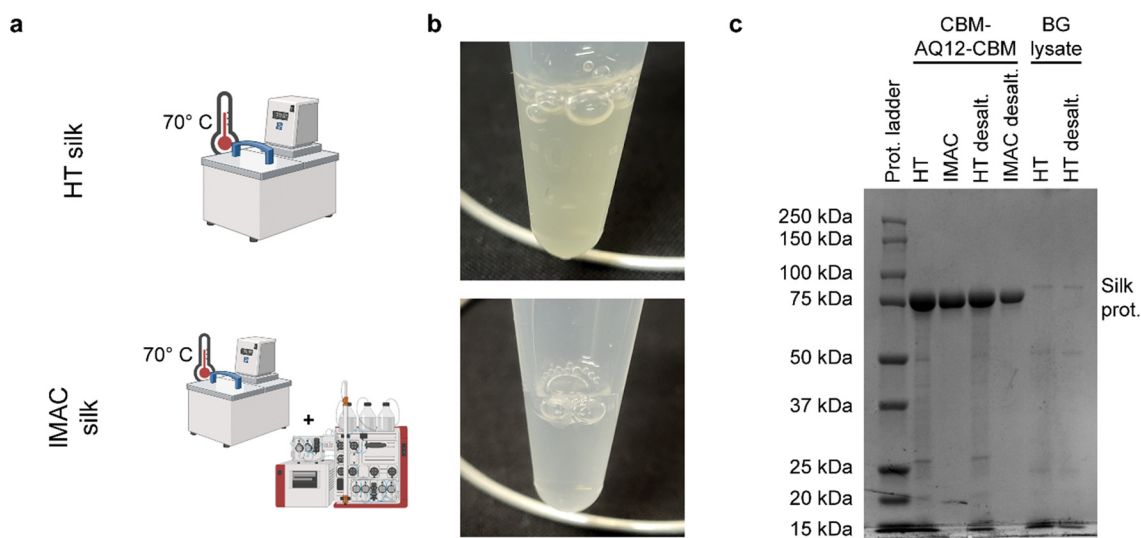


Fig. 2 Purification of the spider silk protein CBM-AQ12-CBM. (a) Protein purification involving heat treatment (HT) for 30 min at 70 °C or HT combined with His-tag immobilized metal affinity chromatography (IMAC). (b) The HT silk solution shows a slightly yellowish color while the IMAC silk solution is clear and transparent. (c) SDS-PAGE of CBM-AQ12-CBM and BG lysate before and after desalting to water. The HT sample shows several other faint bands in addition to the silk protein (85 kDa) indicating other heat stable proteins. No additional bands are observed after IMAC purification. Bands indicating other heat stable proteins are also observed in the BG lysate.

indicating the presence of smaller proteins are visible in the HT sample and are not removed by desalting. In contrast, no

additional bands are visible after IMAC. Furthermore, a control, free from silk protein, was prepared to obtain the components



in the purification background (called BG lysate). This sample also shows multiple faint bands, indicating the presence of a variety of proteins.

The type of background molecules, as well as the yield of silk proteins, were further investigated by amino acid analysis and DNA/RNA extraction. Amino acid analysis of HT silk, IMAC silk, and BG lysate was performed to determine the concentration of CBM-AQ12-CBM and to estimate the total protein amount, including other heat stable proteins. For this, the concentration of CBM-AQ12-CBM was calculated based on the protein sequence and the number of amino acids measured during amino acid analysis. This value was compared to the total amount of amino acids to determine the amount of protein in the BG lysate. However, due to the degradation of specific amino acids (*e.g.*, cysteine and methionine) during the hydrolysis process required for amino acid analysis, this is only an estimation. A whole genome extraction of the differently purified protein samples was carried out with and without RNase to estimate the amount of RNA and DNA present in the samples.

HT silk shows the largest solid fraction with a dry mass of $1.4 \pm 0.1\%$ (mean \pm standard error of the mean, $n = 4$), while IMAC silk has a dry mass of 0.4%. The dry mass of BG lysate is $0.8 \pm 0.2\%$ ($n = 2$) (Fig. 3(b)). HT silk contains a significant number of various background molecules which include DNA, RNA, and other components (Fig. 3(a)). Other components might be, *e.g.*, sugars. The fraction of these background molecules is estimated to be around $4.5 \pm 1.8\%$ of DNA, $5.3 \pm 5.6\%$ of RNA, and $45.8 \pm 9.2\%$ of other components, respectively ($n = 4$). It was not possible to identify other heat stable proteins in the HT silk based on the composition of amino acids given by the amino acid analysis. However, SDS-PAGE gives an estimation of their size and relative amount (Fig. 2(c)). This becomes more evident from the solid fraction composition of the BG lysate which consists of around $10.9 \pm 6.4\%$ DNA, $11.2 \pm 6.4\%$ RNA, $16.9 \pm 1.4\%$ other heat stable proteins,

and $61 \pm 11.4\%$ other components ($n = 2$). No background components of any kind were detected in the IMAC silk. In addition to the presence of background molecules, HT silk and IMAC silk differ in their average yield of silk protein with 560 and 260 mg L^{-1} of culture, respectively. The pH shows no significant variation between the different samples and is around 5.1–5.3 after the final desalting step. The conductivity is highest in the BG lysate with $320 \pm 105 \mu\text{S cm}^{-1}$ ($n = 2$) and lowest in IMAC silk with $56.9 \mu\text{S cm}^{-1}$ ($n = 1$). The conductivity of HT silk is around $201 \pm 65 \mu\text{S cm}^{-1}$ ($n = 4$).

When comparing the dry mass composition of HT silk and the IMAC silk it becomes clear that there is a multitude of various background molecules present in HT silk, such as DNA and RNA. Both are known to affect protein condensation and might also interact with the spider silk protein. The composition of the background becomes clearer from the dry mass composition of the BG lysate which essentially shows the background molecule composition without the spider silk protein. The complex composition of the background hinders the prediction of possible interactions between the background molecules and the silk protein. In contrast, the absence of detectable background molecules in IMAC silk eases the understanding of molecular interaction between silk proteins without the influence of other molecules and thus creates a simplified model system. Reducing the complexity of the system, batch-to-batch variation and unwanted molecular interactions enables reproducible protein condensation.

Background molecules lower the critical concentration for LLPS of silk proteins

HT silk and IMAC silk were prepared as described above (see the methods section for details). A third sample was prepared by mixing IMAC silk with BG lysate (called IMAC silk + BG lysate). Each sample was concentrated in centrifugal concentrators (30 kDa cutoff) containing 6 ml of 2 mg ml^{-1} protein



Fig. 3 Composition of the protein samples after purification. (a) Dry mass composition of HT silk, IMAC silk and BG lysate showing a significant number of various background molecules including DNA and RNA for HT silk and BG lysate. No background molecules were detected for IMAC silk. (b) Comparison of production yield, pH, conductivity, and dry mass of HT silk, IMAC silk and BG lysate showing a clear difference in the yield, conductivity, and dry mass between HT and IMAC purification while BG lysate shows similar properties to HT silk.





Fig. 4 Light microscopy images with phase contrast of concentrated samples of HT silk, IMAC silk and IMAC silk + BG lysate. Samples with ~ 30 , 60 , 120 and 180 mg ml^{-1} were imaged. HT silk and IMAC silk + BG lysate form condensates at $\sim 30 \text{ mg ml}^{-1}$ protein concentration, while IMAC silk forms condensates at $\sim 120 \text{ mg ml}^{-1}$. In HT silk and IMAC silk + BG lysate the presence of an inverted phase can be observed at 60 and 120 mg ml^{-1} . It was not possible to concentrate these samples until 180 mg ml^{-1} . IMAC silk also has condensates at 180 mg ml^{-1} .

solution. The concentrating progress was regularly monitored. When a desired final concentration (final volume) was reached, the concentrated protein solution was collected and imaged using optical microscopy (Fig. 4). For the HT silk and IMAC silk + BG lysate, LLPS occurred at a protein concentration of 30 mg ml^{-1} . When further concentrating these samples to 60 and 120 mg ml^{-1} , the coexistence of condensates in dilute phase (normal phase) and regions with dense phase volume fractions higher than 50% (inverted phase) was observed in the same sample. It was not possible to concentrate these samples to 180 mg ml^{-1} . The IMAC silk showed no presence of condensates at protein concentrations of 30 and 60 mg ml^{-1} , but at 120 mg ml^{-1} . In contrast to the first two samples, IMAC silk was possible to be concentrated to 180 mg ml^{-1} , resulting in LLPS. Thus, the presence of background molecules reduces the critical concentration for LLPS by 75% . In all samples coalescence of condensates was observed, indicating liquid-like properties. Upon dilution of condensed samples we observe that condensates disappear, indicating that condensate formation is reversible. The presence of an inverted phase was identified by the partition of silk protein precipitates, which form due to the metastability of the system, into the dense phase, as well as by adding free green

fluorescent protein (eGFP) which partitions into the dilute phase (Fig. S3, ESI[†]).

Scanning electron microscopy (SEM) shows filamentous structures originating from silk proteins

Differences in the supramolecular interactions of the silk protein in condensed samples and additional interactions with the background molecules were investigated using SEM. Concentrated samples of HT silk, IMAC silk + BG lysate, IMAC silk and only BG lysate were vitrified with liquid ethane ($-180 \text{ }^\circ\text{C}$) and freeze dried. All samples, except BG lysate, contained condensates. Freeze dried samples were cracked, transferred to carbon tape coated stubs and sputter coated with 6 nm Au/Pd . The consistency of the samples after freeze-drying varied between the different samples. Samples containing silk protein required a force to break into pieces, while the BG lysate behaved like a powder. Samples were imaged with the Sigma VP scanning electron microscope (Zeiss) using a SE2 detector (Fig. 5). The BG lysate sample shows a spongy structure that can arise during sample preparation. However, in contrast to the BG lysate, all samples containing silk protein show thin filament-like structures. These are most prominent in HT silk and IMAC silk + BG lysate. Additionally, a layered structure is





Fig. 5 Scanning electron microscope images (Zeiss Sigma VP, SE2 detector, 6 nm Au/Pd coating) of freeze-dried HT silk (34 mg ml^{-1}), IMAC silk + BG lysate (30 mg ml^{-1}), IMAC silk (142 mg ml^{-1}) and BG lysate (prepared in a similar way to that of IMAC silk). All samples show the presence of a bicontinuous network. Samples containing silk protein also show filamentous structures.

very prominent in IMAC silk + BG lysate. The filamentous structures are observed to stretch between the layers. Structures that might correlate to condensates were only observed in the IMAC silk. The filamentous structures likely originate from the silk protein and might be the reason for the stronger material properties that prevent the freeze-dried samples from breaking down into a powder. Similar layered structures, as observed especially in the IMAC silk + BG lysate, were previously described for hydrogels of the silk protein eADF4.³⁶ However, in this case the effect of cross-linking was investigated and the layered structure was assumed to be a result of the chemical sample fixation which differs from the vitrification done in this study.

Background molecules affect the uptake of AIEs by silk condensates

AIEs are small organic molecules that show increased photo-emission in an aggregated state, triggered by the restriction of molecular vibrations and rotations in the dense environment of the condensates.^{37,38} The partition of two different AIEs, 4,4',4'',4'''-(ethene-1,1,2,2-tetrayl)tetrabenzoic acid (TPE4-COOH) and 4,4',4'',4'''-(ethene-1,1,2,2-tetrayl)tetrakis(benzene-4,1-diyl)tetrakis(1-hexylpyridin-1-ium) bromine (TPE4PH) (Fig. S4, ESI[†]), was studied to investigate differences in the biophysical properties of the condensates (Fig. 6).³⁹ Free eGFP was used to verify the identity of the dilute phase in the sample. TPE4COOH partitions preferably into the condensates for all three types of condensates. However, fluorescence is still detectable also in the dilute phase. The AIE fluorescence emission contrast between the dense and dilute phase is higher for the HT silk and the IMAC silk + BG lysate compared to the

IMAC silk. TPE4PH only partitions into the condensates of samples that contain background molecules *i.e.*, HT silk and IMAC silk + BG lysate. TPE4PH stays in the dilute phase for IMAC silk. Thus, the selective uptake of small molecules, such as AIEs, by the spider silk condensates studied here depends on the presence of background molecules. In the case of TPE4COOH the presence of these molecules increases partitioning into the condensate. The partitioning effect is even stronger for TPE4PH which only enters into the condensates in the presence of background molecules. The increased uptake of other molecules into condensates as well as the decrease of the condensation threshold have been previously described for crowding agents.⁴⁰ Therefore, the background molecules present in the HT silk and the IMAC silk + BG lysate might act as crowding agents. We further discuss this hypothesis in the last paragraph of this section.

Micropipette aspiration shows that the presence of background molecules affects the surface tension of condensates

Micropipette aspiration was used to determine the biophysical properties of the IMAC silk condensates. The biophysical properties of the HT silk condensates were determined using micropipette aspiration as described before.²⁴ The IMAC silk condensates were aspirated with 20 Pa applied pressure ΔP with a micrometer-sized glass pipette (Fig. 7(a) and Video S1, ESI[†]) followed by the retraction of the tongue when $\Delta P = 0$ Pa (Fig. 7(b) and Video S1, ESI[†]). The aspiration and retraction length were measured (Fig. 7(c)) and plotted against time, to obtain the aspiration rate \dot{L}_a and retraction rate \dot{L}_r from the slope of the corresponding curves (Fig. 7(d) and (e)). The aspiration and retraction rates were used to calculate the bulk





Fig. 6 Partition behavior of the aggregation induced emitters TPE4COOH and TPE4PH into silk protein condensates. Bright field images are shown in the first column. Images with free eGFP were taken to validate the correct phase. TPE4COOH partitions into all condensates. TPE4PH only partitions into the HT silk and IMAC silk + BG lysate, but not into IMAC silk condensates.

viscosity η_b and the critical pressure ΔP_c of aspiration (see eqn (2) and (4) in the methods section). The bulk viscosity η_b of the IMAC condensates was 5.4 ± 0.7 Pa s (mean \pm standard error of the mean, $n = 9$) (Fig. 7(f) and Table S1, ESI[†]), while the effective bulk viscosity η_{eff} (eqn (5)) of the HT silk condensates was only 2.2 ± 0.4 Pa s ($n = 17$) (original data published previously in Tunn *et al.*²⁴). The surface tension of the IMAC silk condensates was obtained from the Young Laplace law (eqn (1)). The mean surface tension γ of the IMAC silk condensates was 52.2 ± 3.2 $\mu\text{N m}^{-1}$ ($n = 9$). The surface tension of the HT silk condensates was more than two times lower (19.1 ± 2.2 $\mu\text{N m}^{-1}$ ($n = 17$)) than the surface tension of the IMAC silk condensates.²⁴ Interestingly, some of the HT silk condensates burst upon aspiration. Therefore, we assume that the HT silk condensates form a shell-like assembly at the interface between the dense and dilute phase with a surface viscosity η_s of 53.2 ± 9.3 $\mu\text{N s m}^{-1}$ ($n = 17$).²⁴ We did not observe any bursting of the IMAC silk condensates during micropipette aspiration or under any other experimental conditions tested. Therefore, we assume that the IMAC silk condensates do not have a shell-like assembly and report the viscosity of the bulk condensate η_b . The reduced surface tension of HT silk condensates can be explained by the assembly of background molecules on the interface. In this case the background molecules would act as a

surfactant. Furthermore, the assembly on the interface could explain the bursting of condensates observed with HT silk.

The coalescence time of condensates is affected by background molecules

Light microscopy videos were acquired to study the coalescence of IMAC silk condensates, HT silk condensates and IMAC silk + BG lysate condensates (Videos S2–S4, ESI[†]). The final condensate diameter after the coalescence and the total time of the coalescence event were determined with ZEN microscopy software (Zeiss) (Fig. 8(a)). The final condensate diameter was plotted against the coalescence time (Fig. 8(b)). Observed IMAC silk condensates and IMAC silk + BG lysate condensates were generally smaller than those from HT silk. The coalescence time increased with the condensate size in all samples. Coalescence of the observed IMAC silk condensates is fastest with less than 1 s. Coalescence of the observed HT silk condensates was slower compared to IMAC silk condensates with a similar size range. In contrast, IMAC silk + BG lysate condensates show a much slower coalescence time of up to 14 s for condensates in the same size range.

Various aspects can explain the increased coalescence time in the presence of background molecules. One possibility is the higher total amount of macromolecules in samples with





Fig. 7 Micropipette aspiration of IMAC silk condensates. (a) Aspiration of a condensate. Aspiration direction marked with the arrow. (b) Retraction of the tongue at $\Delta P = 0$ Pa. Arrow points at the retraction direction. (c) Scheme of the micropipette aspiration with pipette radius R_p , the condensate radius R_0 and the length L of the aspirated or retracting condensate, respectively. (d) Aspiration curve with linear fit $L_a = \dot{L}_a t + n$ of a condensate. (e) Retraction curve with linear fit: $L_r = \dot{L}_r t + n'$ of a condensate. (f) Box plots of the surface tension (obtained from eqn (1)) and the bulk viscosity (obtained from eqn (2)). IQR = inter quartile range. Number of datasets: 9.



Fig. 8 Coalescence studies of silk protein condensates. (a) Coalescence of two IMAC silk condensates. (b) The coalescence time of condensates is dependent on the final droplet diameter for HT silk, IMAC silk + BG lysate and IMAC silk. A larger final condensate diameter results in a slower coalescence time for all samples. IMAC silk condensates show the fastest coalescence, while IMAC silk + BG lysate condensates show the slowest coalescence.

background such as HT silk and IMAC silk + BG lysate that can lead to increased viscosity of the dilute phase. The change in the aspect ratio over time was fitted using an exponential decay function (see details in the methods section) to obtain the characteristic relaxation time scale of the coalescence of the condensates (Fig. S5a–c, ESI†). The characteristic relaxation

time was then plotted against the characteristic length to obtain the inverse capillary velocity (ICV) of the condensates (Fig. S5d, ESI†). The ICV was $0.01 \text{ s } \mu\text{m}^{-1}$ for IMAC silk ($n = 11$), $0.07 \text{ s } \mu\text{m}^{-1}$ for HT silk ($n = 10$) and $0.31 \text{ s } \mu\text{m}^{-1}$ for IMAC silk + BG lysate ($n = 5$). Using the surface tension values obtained from micropipette aspiration (HT silk and IMAC silk) the bulk



viscosity $\eta_{b,c}$ of the condensates can be determined from the ICV. The $\eta_{b,c}$ of the IMAC silk was 0.5 Pa s and 1.4 Pa s for the HT silk condensates. These bulk viscosities are 2–10 \times lower than the bulk viscosities obtained from micropipette aspiration with 5.4 Pa s for IMAC silk and 2.2 Pa s for HT silk condensates. The difference of the viscosity obtained from the aspiration data from that of the coalescence data may be due to viscous dissipation, the geometry of the micropipette setup and the ICV analysis, which does not take the viscosity of the dilute phase into account.

Possible interaction mechanisms of the background molecules with the silk-protein

The presented results indicate that the background molecules have a strong effect on the protein concentration required for condensation, the partitioning of dye molecules and the biophysical properties of the spider silk condensates. However, through Fourier-transform-infrared spectroscopy (FTIR) of IMAC silk and HT silk it is not possible to detect a change in protein secondary structure (Fig. S6, ESI †). Fig. 9 illustrates possible mechanisms of the interaction of background molecules with condensate forming proteins in general. Probable mechanisms relevant for the system studied here are discussed in the following paragraphs.

Background molecules can affect protein condensation in various ways. One common phenomenon is molecular crowding. Crowding agents such as dextran and polyethylene glycol (PEG) are known to induce LLPS by increasing the effective concentration of the protein.⁴¹ This results in LLPS occurring at

lower protein concentrations. It is possible that some components of the background molecules act as crowding agents. Evidence pointing towards the crowding hypothesis is that uptake of TPE4PH into the HT silk and the IMAC silk + BG lysate condensates is enhanced compared to IMAC silk condensates where it is excluded. It has been shown previously that crowding agents can promote the uptake of molecules into condensates.⁴⁰ However, due to the complex composition it is difficult to predict which molecules present in the background could have a crowding effect.

Another possible interaction between background molecules and condensating proteins is cooperative condensation, *i.e.* complex coacervation.⁴² In particular, DNA and RNA are known to form complexes with intracellular proteins that contain RNA or DNA binding domains.^{33,43–45} Generally, negatively charged RNA or DNA can interact with positive charges of proteins to form complex condensates. The spider silk protein CBM-AQ12-CBM has a mainly neutral AQ12 middle part, and slightly negatively charged CBM terminal domains at pH values higher than the isoelectric point. This indicates a low probability of complex condensation with negatively charged DNA and RNA. However, other weak molecular interactions such as H-bonding and hydrophobic interactions between DNA/RNA and the intrinsically disordered middle part could occur.⁴⁶ Cooperative effects could also result in a lower concentration needed for LLPS since the number of possible interactions and the concentration of macromolecules participating in the condensation is increased.

Furthermore, the presence of background molecules affects the biophysical properties of the condensates, such as surface tension and viscosity. Micropipette aspiration revealed that the surface tension is two times higher for the IMAC silk condensates compared to the HT silk that contains background molecules. The lower surface tension of the HT silk condensates might contribute to the condensation at lower protein concentrations observed for the HT silk. In the HT silk the background molecules could act in a similar way as surfactant and assemble at the interface of condensates (interfacial segregation). Additionally, if a surface tension reduction takes place by background molecules assembling at the interface, these molecules are likely to interact with the condensating proteins at the surface of the condensate. This could reduce the fluctuations of the proteins at the interface leading to the formation of a thin protein shell. The existence of a thin protein shell might also explain the longer fusion times observed for the HT silk condensates as well as the bursting of the HT silk condensates studied by Tunn *et al.*²⁴ The formation of condensates with a shell layer has been previously found for several intracellular complex condensates containing proteins and DNA/RNA.³⁸ However, the presence of the background molecules also increases the overall viscosity of the system, which could lead to increased fusion times of the HT silk and the IMAC silk + BG lysate condensates.

Another possible action mechanism of the background molecules is the stabilization of the mixture of these molecules and the proteins. For example, background molecules could



Fig. 9 Interaction mechanisms of background molecules with spider silk proteins. For simplicity, all background molecules (DNA, RNA, other proteins, other components) are represented in orange. The spider silk protein is represented as blue dots (CBM) connected by the green IDR (AQ12). Molecules are not drawn to scale.



screen charges on the surface of proteins and thus inhibit molecular interactions between the proteins. This would reduce the likelihood of LLPS occurring.⁴⁷ Since we observe condensation of the spider silk protein at lower protein concentrations in the presence of background molecules, mixture stabilization seems unlikely in our system.

We assume that any kind of fragmented DNA or RNA could have a similar effect on the protein condensation as we observe with HT silk and IMAC silk + BG lysate. This is due to our sample preparation involving cell lysis with a lysis buffer that includes DNase, which likely results in HT silk and BG lysate mainly containing fragmented DNA. Additionally, RNases present in the environment will likely cut the remaining RNA in the HT silk and BG lysate into fragments. We further tested this by adding ssDNA from salmon testes (deoxyribonucleic acid sodium salt from salmon testes, Sigma-Aldrich) to the IMAC silk (Fig. S7, ESI†). Sonicated and thus fragmented DNA enhances condensation of IMAC silk while long DNA that was not sonicated shows no effect. However, adding sonicated ssDNA to IMAC silk seems to easily gel the condensates.

In the case of the IMAC silk that does not contain any detectable background molecules, we assume that single component condensation takes place. The IMAC silk condensates formed are simple condensates containing only CBM-AQ12-CBM. The concentration of the IMAC silk needed to form condensates is four times higher than for the HT silk that contains background molecules. The results of viscosity measurements of the IMAC silk varies 10-fold between the coalescence (0.5 Pa s) and micropipette techniques (5.4 Pa s) but less than 2-fold for the HT silk condensates (1.4 and 2.2 Pa s). A lower sample viscosity could explain the fast coalescence of the IMAC silk condensates determined by coalescence analysis. In contrast, a higher viscosity of the IMAC silk could be explained by the overall higher silk protein concentration. The viscosity of the HT silk condensates is influenced by the presence of background molecules.

3. Conclusions

The formation of condensates of recombinant silk proteins was strongly affected by the presence of background co-solutes. Several properties of the condensates—such as surface tension, viscosity, uptake of dyes—were also strongly affected. The presence of background molecules creates a complex environment where individual contributions are difficult to pinpoint. We find several implications of this strong context dependency. One is practical—when producing biosynthetic materials, the assembly steps become difficult to control if there are variations in parameters that are unknown, posing a risk, for example to batch to batch variations. We cannot expect any system to be 100% pure, so the effect of co-solutes will always be present, just the extent of their effect is varied. On the other hand, in natural systems materials such as silks are not a single component, and we should expect that interactions between components affect strongly how materials are formed.⁴⁸ We

should therefore expect that designs for biosynthetic materials could benefit from involving the use of multicomponent systems. Furthermore, silk condensates are a promising material in various biomedical applications. Condensates in general are possible candidates for drug delivery systems, bio-adhesives, and for tissue engineering and repair.^{49–53} Acquiring a deeper understanding of mechanisms taking place at the condensate interface is crucial for applications. Additionally, silk proteins have high biocompatibility and low cytotoxicity.^{54–56} Here condensation was achieved more easily with background components, and as we demonstrated earlier proteins assembled in this way can form highly functional materials.^{9,11} A deeper understanding of the interactions involved will be needed.

In our interpretation, the background molecules might act as a molecular crowder, as components partition into the condensates (cooperative condensation) or form an interfacial layer on the surface of the protein condensate (interfacial segregation). Combinations of these are also possible. Further research will focus on determining the identity of the background molecules affecting the condensate formation in the HT sample and their action mechanisms. IMAC silk as a simplified model system will allow investigation of the influence of various macromolecules on LLPS and condensate properties in a controlled way.⁵⁷ This will enable us to decipher conditions that benefit the condensate properties for material formation.

An interesting question will be to understand how widely our results for silk-IDRs can apply to other types of IDRs.⁵⁸ The formation of biomolecular condensates through IDRs is of wide interest in biological research.^{59,60} As a methodology to understand their functions, comparing them *in vivo* and *in vitro* is an essential approach.^{61,62} Not only should we think that background components in *in vitro* studies affect properties of IDRs, but also that unavoidably the *in vivo* environments have a strong presence of background.⁶³ The evaluation of this background should be included in any study on the condensation of IDRs.

4. Methods

Protein expression and purification

The silk fusion protein CBM-AQ12-CBM (85 kDa) was used in this study. This protein has a triblock structure, with an AQ12 block flanked on both sides by a cellulose binding module (CBM).⁹ The AQ12 block is an engineered silk protein sequence made up of 12 repeats derived from the major ampulla gland silk fibroin 3 from *Araneus diadematus*.⁶⁴ At the C-terminus of the silk-protein a polyhistidine (6× His) has been added, to facilitate affinity purification. The cloning procedure has been described previously.⁹ The silk protein was expressed in *E. coli* BL21 (DE3) (ThermoFisher Scientific) with a pEt-28a (+) (kanR) (Novagen) vector at 30 °C using EnPresso media (EnPresso B 500, EnPresso GmbH) in accordance with the protocol of the manufacturer. Full-baffled plastic flasks were used for the expression (Tunair). 24 h after the induction, the cells were



harvested (11 949 rcf for 15 min) and resuspended in 50 mM tris/HCl lysis buffer (pH 7.4) containing 100 mM NaCl, 3 mM MgCl₂, 1 mg ml⁻¹ lysozyme (EC 3.2.1.17, Merck), 20 µg ml⁻¹ DNase I (EC 3.1.21.1, Merck) and protease inhibitor tablet (1 tablet/50 ml buffer, SIGMAFAST™ Protease Inhibitor Cocktail Tablets, EDTA-Free, Merck). For resuspension, 3 ml of the lysis buffer was added for every gram of the pellet. After 1 h incubation at room temperature (RT) the cells were lysed by sonication (40% amplitude, 3 × 1 min with 2 s pulse time) using a Q500 sonicator (Qsonica) with 1/2" sonicator tip 4406 (Ramcon). The majority of non-target proteins was removed through heat treatment purification (30 min, 70 °C) and centrifugation (2 × 15 min at 3200 rcf, RT).

E. coli cells containing an empty pEt-28a (+) (kanR) (Novagen) vector instead of the vector coding for the silk protein, were used to prepare the BG lysate as a control. The same expression and lysis procedures were performed on the control cells.

Part of the HT silk was further purified by His-tag immobilized metal affinity chromatography (IMAC) (ÄKTA-pure, Cytiva) using HisTrap FF columns (Cytiva) and binding buffer (500 mM NaCl, 20 mM Imidazole, pH 7.4) and elution buffer (500 mM NaCl, 500 mM Imidazole, pH 7.4). All silk protein and the control BG lysate were desalted using Econo-Pac 10DG columns (Bio-Rad) into deionized water. The whole volume of one production batch for each sample was mixed in plastic tubes to achieve consistent protein concentration and solution composition. Conductivity (Jenway 4520 conductivity meter) and pH (pH-indicator strips 4.0–7.0, Merck) were measured for all samples. The protein solution was stored at –80 °C after flash freezing with liquid nitrogen.

SDS-PAGE

SDS-PAGE electrophoresis was performed for samples obtained at different stages of the expression and purification. Pre-cast 10% Mini-Protean TGX gels (Bio-Rad) and Precision Plus Protein Standard Dual Color (Bio-Rad) were used. SDS-PAGE was performed at constant 110 V ~ 60 min and staining was done with Coomassie brilliant blue R-250. The destaining solution contained 48% acetic acid and 40% ethanol. The gel was imaged with a Gel Doc XR+ Gel Documentation System (Bio-Rad).

Amino acid analysis

Amino acid analysis was carried out to determine the concentration of CBM-AQ12-CBM as well as the total amount of protein. The analysis was done as previously described.^{11,24} In short, samples were hydrolyzed in 6 M HCl (Merck), containing 0.1% phenol (Sigma-Aldrich). L-Norleucine (Sigma-Aldrich) was added to all samples as an internal standard. After acid evaporation, samples were resuspended in a citric acid buffer (Sykam GmbH). The samples were measured with S433 amino acid analyzer (Sykam GmbH) using a 570 nm and 440 nm UV detector. The retention time of individual amino acids was determined with an external standard (Sykam GmbH). The concentration of CBM-AQ12-CBM was quantified with the

internal standard and the proteins sequence. Specifically, alanine, glycine, glutamine, and glutamic acid were used for concentration determination, due to these amino acids being most abundant in CBM-AQ12-CBM. The total protein amount was determined by taking all the detected amino acids into account. However, since certain amino acids degrade during hydrolysis, this only provides an estimation of the total protein amount.

DNA/RNA quantification

DNA and RNA present as the background in protein solutions and BG lysate were quantified with the Wizard Genomic DNA Purification Kit (Promega). Since the samples were already lysed and purified no additional lysis was performed. Instead, 600 µl of the sample were added to a microcentrifuge tube. For each sample (HT silk, IMAC silk, BG lysate) two tubes were prepared. In one tube of each sample, 3 µl of RNase solution were added and the tubes were inverted 2–5 times. After 15 min incubation at 37 °C, 200 µl of protein precipitation buffer was added to each microcentrifuge tube. Tubes were vortexed for 20 s followed by a 5 min incubation on ice and centrifuging for 3 min at 16 000 rcf. The supernatant was transferred to clean tubes containing 600 µl of RT isopropanol and the tubes were gently inverted a few times. Samples were centrifuged for 2 min at 16 000 rcf and the pellet was washed with RT 70% ethanol. The pellet was air dried for 15 min and samples were rehydrated in 40 µl deionized water overnight at 8 °C. Absorption was measured from 200 nm to 350 nm with an EON microplate reader (Biotek). Absorbance at 260 nm was used to calculate DNA and RNA concentrations. Samples prepared with RNase were used to calculate the amount of DNA while samples without RNase were used to calculate the DNA + RNA amount. To determine the amount of RNA, the amount of DNA was subtracted from the DNA + RNA amount. Samples only containing DNA were calculated with an extinction coefficient of 50, while samples containing a mixture of DNA and RNA were calculated with an extinction coefficient of 45.

Dry mass

The dry mass of protein solutions and BG lysate was determined after freeze-drying (Christ Alpha 2–4) and weighing the residual pellet.

Concentrating

HT and IMAC CBM-AQ12-CBM, as well as BG lysate, were thawed and centrifuged at RT for 15 min at 3200 rcf. Three samples were prepared, 2 mg ml⁻¹ HT silk, diluted with deionized water, 2 mg ml⁻¹ IMAC silk, diluted with deionized water, and 2 mg ml⁻¹ IMAC silk, diluted with BG lysate (ratio 1:1). For each sample, four centrifugal concentrators with a 30 kDa cutoff (Vivaspin, Sartorius) were each filled with 6 ml of the prepared protein solution. Samples were concentrated in RT at 1200 rcf until the desired protein concentrations of 30, 60, 120, and 180 mg ml⁻¹ were achieved for each sample. The remaining volume of the concentrated protein solution was



used as an estimate of the protein concentration before collection of the sample from the concentrator tube. The final concentration of the samples was determined based on the weight of the collected solution assuming a density of 1 g cm^{-3} . Concentrated samples were used for further studies on the same day as they were concentrated.

Imaging to study the effect of protein concentration on LLPS of silk proteins

Samples were imaged with an AxioVert.A1 inverted light microscope (Zeiss) after concentration and collection of the sample. Before pipetting the samples onto the glass slides, the samples were gently mixed with a pipette tip by stirring. $3 \mu\text{l}$ of sample were imaged between two coverslips. Imaging was done with a $40\times/0.6$ Ph2 objective and phase 2 contrast ring. The presence of an inverted phase was studied by mixing $4.5 \mu\text{l}$ of concentrated protein solution with $0.5 \mu\text{l}$ free eGFP (Filter set 38, Zeiss). Free eGFP partitions into the dilute phase but not into the dense phase.

Imaging to study the partition behavior of aggregation induced emitters in silk protein condensates

Condensated samples ($4.5 \mu\text{l}$) (prepared as previously described) were mixed with either $4,4',4'',4'''$ -(1,1,2,2-ethenetetrayl)tetrabenzoic acid (TPE4COOH) or $4,4',4'',4'''$ -(ethene-1,1,2,2-tetrayltetrakis(benzene-4,1-diyl))tetrakis(1-hexylpyridin-1-ium) bromine (TPE4PH) ($0.5 \mu\text{l}$) so that the final AIE concentration was $55 \mu\text{M}$ or $40 \mu\text{M}$, respectively. The presence of the correct phase was verified by the addition of free eGFP. Images were taken with a Nikon Eclipse Ti inverted microscope (Nikon), ORCA-flash 4.0 LT digital camera (Hamamatsu, Japan), $60\times/1.4$ oil immersion objective lens, $1.5\times$ tube lens and micromanager software (version 2.0.0). Bright field images were taken for all samples. Samples containing free eGFP were imaged with 5% 470 nm light (LDI Laser Diode Illuminator, 89 North) and emission light was collected between 485–535 nm. Samples containing TPE4COOH were excited with 50% 405 nm light (LDI Laser Diode Illuminator, 89 North) and the emitted light was detected from 410–460 nm. Samples containing TPE4PH were imaged with 20% 445 nm (LDI Laser Diode Illuminator, 89 North) with emission light collection in the range of 450–505 nm. Data analyzation was carried out in Fiji (ImageJ 1.54f).

Micropipette aspiration

Micropipette aspiration of the IMAC silk condensates was performed in *ca.* 2 mm high measurement chambers made by spacing 2 coverslips with 6 layers of parafilm. The chamber was filled with 5–10 μl condensate solution and sealed with a small amount of oil (Immorsol 518F, Carl Zeiss) to prevent evaporation. The experiments were conducted at room temperature using pulled (PN-31, Narishige) borosilicate capillaries (WPI, 1 mm/0.5 mm outer/inner diameter). The radius of the pipettes R_p is typically 10–20 μm . To enable horizontal insertion of the micropipette into the observation chamber the pipettes were bent with a microforge (MF-900, Narishige). A piezo

electric pressure controller (OB1 Mk3, Elveflow) connected to the micropipette was used to fill the micropipette with water before the experiment and control the pressure during experiments. Before the aspiration of the condensates a small amount of dilute phase was aspirated, and the zero pressure of the micropipette was determined. At first the micropipette was brought into contact with a condensate of radius R_0 . The aspiration was conducted by applying a constant negative suction pressure ΔP . After releasing the pressure ($\Delta P = 0 \text{ Pa}$) the relaxation of the condensate was recorded. Videos of the aspiration and retraction with a frame rate of about 10 fps were acquired in bright field mode using a Nikon Eclipse Ti inverted microscope (Nikon) equipped with an ORCA-flash 4.0 LT digital camera (Hamamatsu, Japan). The microscope was operated using micromanager (version 2.0.0). The data was analyzed in Fiji (ImageJ 1.54f) with the manual tracking plugin to measure the aspiration and retraction length. The measure function of Fiji was used to determine the diameter of the micropipette and the condensate. Linear fits and calculations were performed in Microsoft Excel. The surface tension γ of the condensate was obtained from the critical pressure ΔP_c using the Young-Laplace law:¹

$$\Delta P_c = 2\gamma \left(\frac{1}{R_p} - \frac{1}{R_0} \right) \quad (1)$$

with the pipette radius R_p and the condensate radius R_0 . The balance of the aspiration force and the friction force of the condensate in the pipette leads to a viscous flow at constant velocity yielding an aspiration rate \dot{L}_a (eqn (2)).

$$\dot{L}_a = \frac{1}{3\pi\eta_b} R_p (\Delta P - \Delta P_c) \quad (2)$$

with applied pressure ΔP , critical pressure ΔP_c and the bulk viscosity of the condensate η_b . The viscosity was calculated from the aspiration rate (eqn (2)). The retraction rate \dot{L}_r is defined as

$$\dot{L}_r = \frac{1}{3\pi\eta_b} R_p \Delta P_c \quad (3)$$

and was used to calculate the critical pressure. The critical pressure was either determined by stepwise increasing the suction pressure up to a value where the condensate would enter the micropipette or calculated by

$$\Delta P_c = \frac{\Delta P \dot{L}_r}{\dot{L}_a + \dot{L}_r} \quad (4)$$

For the full derivation of all equations please see Guevorkian *et al.*² The values reported in the main text are the mean \pm standard error of the mean (SEM) with the number of datasets in brackets. All data are presented in Table S1 (ESI[†]).

For comparison with the biophysical properties of the HT silk condensates the data published in Tunn *et al.* were used.^{24,65} The surface tension of the HT silk condensates was calculated before. The effective viscosity η_{eff} of the HT



condensates is

$$\eta_{\text{eff}} = \frac{4\eta_s}{3\pi R_p} \quad (5)$$

where η_s is the surface viscosity reported in Tunn *et al.*²⁴ The effective viscosity gives an estimate of the bulk viscosity of the HT silk condensates.

Analysis of coalescence of silk protein condensates

The samples were prepared and concentrated until LLPS occurred, as previously described. 5 μl of sample were imaged in a chamber, constructed of 2 coverslips and a 50 μm thick spacer tape (iSpacer, SunJinLab), to prevent evaporation. Videos of fusion events were taken with an Axio Observer.Z1 (Zeiss) inverted light microscope (20 \times /0.5 Ph2 objective, 1.6 \times tube lens, phase 2 contrast) and Prime BSI sCMOS camera (Photometris). The frame rate was approximately 54 fps. The final condensate diameter after coalescence and coalescence time were determined with ZEN microscopy software (Zen lite 3.9, Zeiss). For further analysis the coalescing condensates were approximated by an ellipse. The aspect ratio ($\text{AR} = l_{\text{long}}/l_{\text{short}}$) of the ellipse over the coalescence time was determined as described previously and fitted with

$$\text{AR}(t) = 1 + (\text{AR}_0 - 1)e^{-t/\tau} \quad (6)$$

where AR_0 is the aspect ratio at $t = 0$ and τ is the characteristic relaxation time of the condensate.⁶⁶ The obtained characteristic relaxation times were plotted against the characteristic lengths of the condensates (calculated as a geometric mean with $l = \sqrt{(l_{\text{long},0} - l_{\text{short},0}) \cdot l_{\text{long},0}}$). The inverse capillary velocity (ICV) was obtained from the slope of a linear fit to the data:

$$\text{ICV} = \frac{\eta_{\text{b,c}}}{\gamma} \quad (7)$$

The surface tension γ obtained from micropipette aspiration allowed us to calculate the viscosity $\eta_{\text{b,c}}$ of the condensates.

Scanning electron microscopy

Samples of HT silk, IMAC silk, and IMAC silk + BG lysate were concentrated until LLPS occurred, as previously described. Additionally, a control free of silk protein was prepared by mixing BG lysate with water, and concentrated similar to IMAC silk + BG lysate. A small amount of sample was transferred into a microcentrifuge tube and vitrified in liquid ethane (-180°C), followed by freeze-drying (Christ Alpha 2–4). Freeze-dried samples were fractured and transferred to stubs coated with carbon tape. The samples were sputter coated with 6 nm Au/Pd and imaged with a Sigma VP scanning electron microscope (Zeiss), SE2 detector at 1.5 kV.

Author contributions

Design and conceptualization of the study: J. T., M. B. L., J.-A. G., A. S.-V. (AIEs), E. A.-P. (AIEs). Experimental part: J. T. (protein expression and purification, concentrating and

imaging, amino acid analysis, quantification of background molecules, SEM, coalescence studies, data analysis), I. T. (micropipette aspiration, AIEs), M. S. (protein expression and purification), T. V. (amino acid analysis, SEM and data analysis), M. M. (plasmid for BG lysate), G. B. (micropipette aspiration), A. S.-V. (synthesis of AIEs). Writing – original draft: J. T., I. T. (micropipette aspiration and parts of discussion), M. S. (parts of methods). Writing – finalizing: M. B. L.

Data availability

All newly generated data are presented in the manuscript and the ESL.† The raw data are available on Zenodo.org: DOI: <https://doi.org/10.5281/zenodo.12529262>.

Conflicts of interest

The authors declare no conflicts of interest.

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