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Amyloids and protein aggregation

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A general discovery in protein science in the past few decades has been the finding that a number of unrelated proteins and peptides all have a marked propensity to form amyloid fibrils *in vivo* and *in vitro*. These structures have become known as the pathological hallmark of some of the most prevalent neurodegenerative diseases. More recently, the process of amyloid formation has been demystified through a number of key mechanistic findings, some of which are highlighted in this themed collection.

1. Equilibrium and solubility

1.1 Phase transitions

Amyloid formation from a solution of a peptide or protein is a phase transition which results in highly ordered fibrils in a solid phase that co-exist with a dilute solution phase consisting mainly of monomers and a minority population of oligomers. At equilibrium, the chemical potential of the peptide in solution, μ_m , is the same as for peptides in the fibrillar aggregates, μ_f .^{1,2}

$$\mu_m = \mu_m^0 + RT \ln [m] = \mu_f = \mu_f^0$$

Crucially, the chemical potential of the monomer is concentration dependent, while that of the fibril is not; this situation, therefore, defines quantitatively the equilibrium in terms of the solubility, which is the highest concentration at which the peptide can exist as monomers for infinite time. Above the solubility, the concentration of monomers in the fibril phase grows with monomer concentration while the equilibrium concentration of monomers in solution tends towards a constant. In addition to the solid amyloid phase, dense liquid phases of proteins can form

through liquid–liquid phase separation (LLPS,^{2–4}). It is becoming increasingly clear that in many cases these functional liquid condensates can promote the formation of pathological amyloid fibrils through undergoing a liquid to solid transition in disease.

1.2 Reversibility

Amyloid formation is a non-covalent polymerization reaction and thus in principle reversible without the requirement to break bonds. During the aggregation process, both the forward and backward processes take place, but their relative rates change over time as an equilibrium is being established. This reversibility is readily demonstrated by diluting an equilibrated system, after which some of the fibrils dissolve to establish a new equilibrium defined by the lower total concentration of peptide. Fibrils may also dissolve if brought to solution conditions at which they are less stable than under conditions at which they were formed.

The review by Buell (<https://doi.org/10.1039/D1SC06782F>) provides a comprehensive summary of the thermodynamics of amyloid formation, highlighting the commonalities and difference between unimolecular protein folding and protein self-assembly. Buell further outlines several examples of methodologies to study amyloid thermodynamics as well as recent findings regarding amyloid stability.

2. Experimental challenges and development

If the monomer concentration is brought above the solubility limit, *e.g.* through a temperature jump, the solution becomes supersaturated and the disappearance of monomers and formation of fibrils can be monitored over time. Using defined and systematic variations in the starting state it is possible to acquire reproducible data that can form the basis for deriving an aggregation mechanism. The sensitivity of nucleation and growth steps to a range of perturbations including sequence inhomogeneity, additional solution components, surface area and roughness of sample containers, air–water interfaces, *etc.* makes it essential to minimize and gain control over these factors. While initial attempts were focused on pure buffer systems, recent advances have enabled mechanistic insights into aggregation processes *in vivo* (Sinnige, <https://doi.org/10.1039/D2SC01278B>).

3. Kinetics and mechanisms

The thermodynamics of amyloid formation can readily be understood as a simple free energy balance between the dilute solution and the aggregated phase as outlined above. By contrast, it has become clear that the kinetics of this process are remarkably intricate and consist of a number of processes that typically occur

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