Location: H14

BP 10: Crystallization, nucleation and self-assembly (joint session CPP/BP)

Time: Tuesday 9:30–10:30

BP 10.1 Tue 9:30 H14

Investigation of the Short-Time Diffusive Dynamics During Salt-Induced Protein Crystallization Using Neutron Spectroscopy — •CHRISTIAN BECK^{1,2}, MARCO GRIMALDO¹, FELIX ROOSEN-RUNGE³, FAJUN ZHANG², FRANK SCHREIBER², and TILO SEYDEL¹ — ¹Institut Laue Langevin, Grenoble, France — ²University of Tübingen, Germany — ³Lund University, Lund, Sweden

Protein crystals are needed to obtain high-resolution protein structures, and therefore understanding different processes/pathways leading to their formation is of fundamental biophysical and medical interest. Previous studies investigating the kinetics of crystallization in situ using static methods (SAXS and microscopy) provided evidence for non-classical crystallization pathways in the presence of multivalent salts [1,2]. Using dissolved β -lactoglobulin proteins as a model system, we studied the ZnCl₂-induced crystallization. Here, we employ quasi-elastic neutron backscattering (NBS) and neutron spin-echo (NSE) spectroscopy to access the kinetics of the nanosecond diffusive dynamics of proteins during crystallization on a nanometer length scale. NBS provides information on the changes of the center-of-mass diffusion, internal diffusive dynamics and on the fraction of immobile proteins associated with the crystals. Accessing coherent scattering with NSE, we probe different scattering vectors q to disentangle the different diffusive contributions of proteins in crystals or aggregates, and in the liquid phase, respectively.

[1] A. Sauter et al. ACS Cryst. Growth Des. 14 (2014) 6357

[2] A. Sauter et al. Faraday Discuss. 179 (2015) 41

BP 10.2 Tue 9:45 H14

Protein crystallization near liquid-liquid phase separation — KLIM PETROV, JAN HANSEN, •FLORIAN PLATTEN, and STEFAN U. EGELHAAF — Heinrich Heine University Duesseldorf

The crystallization of protein (lysozyme) solutions is studied as a function of protein and salt concentration at ambient conditions. In addition to tetragonal crystals at low salt concentrations (far away from phase separation), needle-like and kinetically roughened crystals occur in the vicinity of the binodal. The crystallization induction time and the growth rate are inferred from optical microscopy and linked to the solubility and protein-protein interactions. Based on these data, the different states of the protein solution are linked to different driving forces for crystallization.

BP 10.3 Tue 10:00 H14 Does liquid-liquid phase separation enhance protein crystallization? — •Ralph Maier¹, Andrea Sauter¹, Georg Zocher¹, Stefano Da Vela¹, Olga Matsarskaia¹, Ralf Schweins³, Michael Sztucki⁴, Fajun Zhang¹, Thilo Stehle^{1,2}, and Frank $\begin{array}{l} {\rm Schreiber}^1 - {}^1{\rm Universit\"at} \ {\rm T\"ubingen}, \ {\rm Germany} - {}^2{\rm Vanderbilt} \ {\rm Universit\largey} \ {\rm School} \ {\rm of} \ {\rm Medicine}, \ {\rm Nashville}, \ {\rm USA} - {}^3{\rm ILL}, \ {\rm Grenoble}, \ {\rm France} - {}^4{\rm ESRF}, \ {\rm Grenoble}, \ {\rm France} \end{array}$

Solutions of the protein human serum albumin (HSA) exhibiting a reentrant phase behavior with a metastable liquid-liquid phase separation (LLPS) inside the condensed regime in the presence of trivalent salts [1] were studied, focussing on the effects of the metastable dense liquid phase on the crystallization pathways. Optical microscopy and small angle X-ray and neutron scattering were used to follow protein crystallization and to explore the role of metastable LLPS. No evidence of nucleation inside the dense liquid phase was observed. On the contrary, heterogeneous nucleation at the walls of the glass container dominates. This suggests that the existence of a metastable LLPS is not a sufficient condition for a two-step nucleation. The unstable or metastable dense liquid phases serve as a reservoir for crystal growth. Furthermore, the crystallographic analysis of the resulting crystals shows that crystals with different morphology grown under different conditions share the same structure and the metal ions create two bridging contacts within the unit cell which stabilize the unit cell. [1] Matsarskaia et al., J. Phys. Chem. B, 120, 7731 (2016)

BP 10.4 Tue 10:15 H14

Using x-ray scattering to understand the formation of unexpected structures in organic thin films — JENNY LEBERT¹, EVA M. KRATZER^{1,2}, AXEL BOURDICK³, MIHAEL CORIC¹, STEPHAN GEKLE³, and •EVA M. HERZIG^{1,2} — ¹Herzig Group, MSE Technische Universität München, Lichtenbergstr. 2a, 85748 Garching, Germany — ²Dynamik und Strukturbildung - Herzig Group, Universität Bayreuth, Universitätsstr. 30, 95447 Bayreuth, Germany — ³Biofluid Simulation and Modeling, Universität Bayreuth, Universitätsstr. 30, 95447 Bayreuth, Germany

The morphology plays an important role for the performance of organic, semi-conducting thin films. Understanding the self-assembly processes that occur during the drying of the photoactive films, will allow us to make progress in controlled nanomorphology tuning. We have therefore developed tools to investigate thin film formation processes using synchrotron radiation to resolve structure and structural developments [1,2]. We have now investigated an in-situ polymerization method for polythiophene to examine how much we can influence the morphology during film formation with such an approach. GI-WAXS measurements, molecular dynamics simulations, and spectroscopic analysis suggest the presence of polythiophene in a novel and stable crystal structure with an enhanced intermolecular interaction [3]. [1] S. Pröller et al. Rev. Sci. Instrum. 2017, 88(6): 066101. [2] S. Pröller et al. Adv. Energy Mater. 2016, 6(1): 1501580. [3] J. Lebert et al. ACS Omega 2018, 6: 6388-6394.