Crystallization of macromolecules used to be rather empirical, and because of its unpredictability and frequent irreproducibility, it has long been considered as an “art” rather than science. It is only in the last 20 years that a real need has emerged to better understand and rationalize the crystallization of biological macromolecules. As the molecules involved in crystallization exhibited such contrarious behavior and were poorly characterized, conducting serious research into their crystallization was conceded as hopeless. Only when the demand for crystals by crystallographers and later molecular biologists could no longer be ignored, researchers pursued in earnest the study of macromolecular crystal growth (McPherson 1999).

Researchers working in the protein X-ray crystallography laboratory understand that the first requirement for protein structure determination is to grow suitable crystals.

Without crystals there is no X-ray structure determination of a protein!

The characteristics of crystals and biological macromolecules crystallization data have been compiled in the Biological Macromolecule Crystallization Database (BMCD) (Gilliland et al. 1994, Gilliland 1998). The BMCD contains crystal data and the crystallization conditions, which have been collected from the literature. The current version of the BMCD contains 43406 crystal entries (Tung and Gallagher, 2009) from macromolecules for which diffraction quality crystals have been obtained. These include proteins, protein-protein complexes, nucleic acid, nucleic acid:nucleic acid complexes, protein:nucleic acid complexes, and viruses. All crystallographic information about particular macromolecule including X-ray diffraction data and statistics information are available online in the Protein DataBank (RCSB PDB) database (Berman et al. 2000). The RCSB PDB provides a variety of tools and resources for studying the structures of biological macromolecules and their relationships to sequence, function, and disease.

**CRYSTALLOGENESIS KEYWORDS**

Macromolecules

All of the macromolecules are polymers of one of the precursor classes that include the amino acids, the ribonucleotides and deoxyribonucleotides, sugars of various sorts, fatty acids, etc. These small molecules are linked together in a sequence by complicated series of chemical reactions in the cell to form the macromolecules such as proteins, nucleic acids (RNA and DNA), polysaccharides and lipids. The structural complexity and physiological role of macromolecules are a function of the diversity of the precursors, the sequence in which they are joined together, the number of precursors in the polymer, and finally, the 3D form after polymer synthesis.

Macromolecules assume 3D structures that sequester and pack hydrophobic groups in their interior and leave hydrophilic groups exposed to solvent. Solvent molecules form solvent layers around macromolecules.

Targets of biological crystallogenesis

For biologists, studying crystal growth should be correlated with biological problems, and crystallization projects on macromolecular complexes, membrane proteins, and especially engineered proteins, are being developed.

For physicists, growing large monocrystals can be a goal in itself, and one might speculate that exploration of optical, electronic, mechanical, and other physical properties of crystalline arrays made from biomacromolecules or assemblies can lead to novel frontiers in the material science.

For chemists, usage of chemical and molecular biology tools could lead to designing molecular devices and other nanostructures mimicking macromolecular crystals in the future.

Crystals

Crystals are chemically well defined; often they are geometrical solids with regular faces and sharp edges. From the physical point of view, crystals are regular three-dimensional arrays of atoms, ions, molecules, or molecular assemblies. Ideal crystals can be pictured as infinite and perfect arrays in which the building blocks (the asymmetric units) are arranged according to well-defined symmetries into unit cells that are repeated in 3D by translations. Experimental (laboratory-grown) crystals have finite dimensions; their periodicity is never perfect, due to different kinds of local disorders or dislocations. The phenomenon underlying structural chemistry and biology is the ability of crystals to diffract X-rays, neutrons, or electrons.

Macromolecular crystals

Macromolecular crystals are rather small with poor mechanical properties and a high content of solvent filled channels that make up 20–80% of their volume. These crystals are always extremely fragile and sensitive to external conditions. For this reason macromolecular crystals

---

**Laboratory exercises**

**INTRODUCTION TO LAB EXERCISES - MACROMOLECULAR CRYSTALIZATION**

Ivana Kutá Smatanova

University of South Bohemia in Ceske Budejovice, Faculty of Fisheries and Protection of Waters, South Bohemian Research Center of Aquaculture and Biodiversity of Hydrocenoses and School of complex systems, Zamek 136, 373 33 Nove Hrady and Academy of Sciences of the Czech Republic, Institute of Nanobiology and Structural Biology GCRC, Zamek 136, 373 33 Nove Hrady

ivanaks@seznam.cz
have to be kept in the solvent-saturated environment, else dehydration will lead to crystal cracking and destruction.

**Crystallization and crystallization strategy**

Crystallization is a multiparametric process involving the three classical steps of nucleation, growth, and termination of growth. The tactics to crystallize proteins can be separated into two parts assuming that a significant quantity of pure homogeneous protein is available. First the initial condition must be established where the protein molecule can be displaced into a state of supersaturation followed by an equilibration process that favors minimal nucleation and optimal crystal growth. The methods of crystallizing proteins usually have to be applied over a broad set of conditions adjusting chemical variations such as pH, ionic strength, metal ions or detergents. Physical factors including temperature, gravity, surfaces, viscosity, dielectric properties or vibrations must also be considered during the crystallization process. Biochemical issues also come into play where the purity, modification or aggregations determine the fate of successful crystal growth. In 1999 McPherson (1999) reviewed a comprehensive list of factors effecting protein crystal growth (Table 1). Even when initial crystallization conditions were successfully determined and crystals were produced, their quality may not be sufficient for X-ray diffraction and thus not adequate for subsequent structure determination. Here optimization of the initial screening conditions must be performed to improve the quality of the crystal to enable X-ray analysis. This entails fine ‘tuning’ of any of the variable parameters described above aimed at optimizing the supersaturation state to produce a crystal of the highest quality. Because of the multiparametric nature of the crystallization process and the diversity of the individual properties of proteins, it is strongly recommended to collect as much information about the protein of interest as possible. Table 2 lists examples of questions essential to consider prior to any crystallization experiments.

**Crystallization experiments**

Generally, the protein crystallization experiments proceed in two steps. First step is test screening of the protein solubility considering the precipitants (precipitating agents) and other solution components. Usually in this step, insoluble protein is observed as an amorphous precipitate, which means that the precipitation conditions were too severe to allow crystal growth. In the second optimization step conditions, which gave rise to precipitates in the first step, are modified systematically to allow the advance to insoluble state that is required for the formation of crystal nuclei.

**Crystallization results**

For the examination of the crystallization trials a stereomicroscope is used. Crystals are usually easy to distinguish from amorphous precipitate. Diffractable crystals are typically single, transparent, of a defined form characterized by planar faces and free of cracks and any defects. Crystals are often birefringent, i.e. they appear dark and bright as they rotate under crossed polarizers in the stereomicroscope. Several methods to test whether crystals are protein or salt are now available. These include crush test, dehydration test, and dye binding test, gel electrophoresis and X-ray diffraction.

---

**Table 1. Parameters effecting the crystallization of macromolecules.**

<table>
<thead>
<tr>
<th>Physical</th>
<th>Chemical</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature</td>
<td>pH</td>
</tr>
<tr>
<td>Surfaces</td>
<td>Precipitant type</td>
</tr>
<tr>
<td>Methodology/approach to equilibrium</td>
<td>Precipitant concentration</td>
</tr>
<tr>
<td>Gravity</td>
<td>Ionic strength</td>
</tr>
<tr>
<td>Pressure</td>
<td>Specific ions</td>
</tr>
<tr>
<td>Time</td>
<td>Degree of supersaturation</td>
</tr>
<tr>
<td>Vibrations/sound/mechanical perturbations</td>
<td>Reductive/oxidative environment</td>
</tr>
<tr>
<td>Dielectric properties of the medium</td>
<td>Concentration of the macromolecules</td>
</tr>
<tr>
<td>Viscosity of the mediumrate of equilibration</td>
<td>Metal ions</td>
</tr>
<tr>
<td>Homogeneous or heterogenous nucleants</td>
<td>Crosslinkers/polyions</td>
</tr>
<tr>
<td>Rate of equilibrium</td>
<td>Detergents/surfactants/amphiphiles</td>
</tr>
<tr>
<td>Rate of crystallization</td>
<td>Non-macromolecular impurities</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Biochemical</th>
<th>Biological</th>
</tr>
</thead>
<tbody>
<tr>
<td>Purity of the macromolecule/impurities</td>
<td>Rarity of most biomacromolecules</td>
</tr>
<tr>
<td>Ligands, inhibitors, effectors</td>
<td>Biological sources and physiological state of organisms or cells</td>
</tr>
<tr>
<td>Aggregation state of the macromolecule</td>
<td>Bacterial contaminants</td>
</tr>
<tr>
<td>Post-translation modifications</td>
<td>Purity of macromolecules</td>
</tr>
<tr>
<td>Source of macromolecule</td>
<td>Macromolecular contaminants</td>
</tr>
<tr>
<td>Proteolysis/hydrolysis</td>
<td>(odd macromolecules or small molecules)</td>
</tr>
<tr>
<td>Chemical modifications</td>
<td>Sequence microheterogeneities</td>
</tr>
<tr>
<td>Genetic modifications</td>
<td>(fragmentation by proteases or nucleases)</td>
</tr>
<tr>
<td>Inherent symmetry of the macromolecule</td>
<td>Conformation microheterogeneities</td>
</tr>
<tr>
<td>Stability of the macromolecule</td>
<td>(flexible domains, oligomers, aggregation)</td>
</tr>
<tr>
<td>Isoelectric point</td>
<td>Batch effects (two batches are not identical)</td>
</tr>
<tr>
<td>History of the sample (denaturation, degradation)</td>
<td></td>
</tr>
</tbody>
</table>

---

© Krystalografická společnost
In the case of growing microcrystals, the seeding techniques can be used to grow the crystal. The seeds (microcrystals) are transferred to a new protein-precipitant drop using a streak seeding wand or a crystal transfer syringe, respectively. Seeds provide a template on which further molecules can assemble, and given the proper environment, time, and patience, the seed will enlarge into a crystal.

Experience and reproducibility are guides in making crystallization experiments. Don’t forget this and welcome in PX!


### Table 2. Essential questions to consider prior to successful crystallization.

<table>
<thead>
<tr>
<th><strong>Biology and production</strong></th>
<th><strong>Biochemistry</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>What is the biological origin of the protein of your interest?</td>
<td>How long does it take to purify one batch of the protein?</td>
</tr>
<tr>
<td>Has the gene encoding the protein been sequenced</td>
<td>How do you assess purity of the protein?</td>
</tr>
<tr>
<td>Has the protein been cloned?</td>
<td>(SDS-PAGE, HPLC, MASS)</td>
</tr>
<tr>
<td>How many mg/l of culture can you produce?</td>
<td>What are the principal characteristics of the protein?</td>
</tr>
<tr>
<td>How many mg of the protein can you obtain in standard purification?</td>
<td>(Mr, isoelectric point, disulfide bridges, hydrophobicity, etc.)</td>
</tr>
<tr>
<td></td>
<td>What are the friendly solvents?</td>
</tr>
<tr>
<td></td>
<td>Is the protein monomeric or oligomeric?</td>
</tr>
<tr>
<td></td>
<td>What is the stability of the protein vs time, T, or pH?</td>
</tr>
</tbody>
</table>

The most frequently used crystallization method is the vapor diffusion technique. In practice, sitting or hanging droplets (mixture of equal volumes of protein and crystallization solution) of up to a few microliters are placed over a reservoir solution. But also the (micro-)batch method is still being used.

Most important, the experimentation does not stop after the cover slide is placed over the reservoir, so to speak. The classical vapour diffusion experiment can be modified by several pre and post set-up techniques:

- A selection of pre set-up, vapour diffusion experiment alterations
  - 1. insertion of an oil barrier that will slow down the equilibration rate
  - 2. use of gels to, among other effects, slow down convection in the droplet
  - 3. use of capillaries in vapor diffusion mode to minimize handling of crystals
  - 4. microseed matrix seeding to outwit nucleation
  - 5. fluorescent dye-labelling of proteins
  - 6. addition of proteases for in situ, limited proteolysis

A selection of post set-up, vapour diffusion experiment alterations
  - 1. change reservoir precipitant concentration
  - 2. change temperature
  - 3. change pH
  - 4. seeding

During the hands-on practical exercises, the pre and post set-up alterations will be demonstrated and discussed in more detail. Also the standard microbatch trials in Terasaki plates (droplets under paraffin oil) can be manipulated by choosing different oils (mixture of silicon and paraffin oil).
CRYSTALLIZATION OF OWN PROTEINS USING COMMERCIAL SCREENING KITS

Lubica Urbániková

Institute of Molecular Biology, Slovak Academy of Sciences, Dubravska cesta 21, 845 51 Bratislava 45, Slovak Republic
lubica.urbanikova@savba.sk

Crystallization of proteins is influenced by many factors. Optimal conditions for crystal nucleation and growth cannot be predicted, they must be found experimentally. An exhaustive search of all variables will be time consuming, tedious and will need large amounts of protein. A highly effective approach to overcome these difficulties is the use of commercially available kits based on a sparse-matrix method. Crystallization kits were empirically derived based on known or published crystallization conditions of various proteins in the past and allow us to screen a large range of buffers, pH, additives and precipitants using small amounts of proteins. They become the most popular way of determining the preliminary conditions for obtaining protein crystals.

CRYSTALLIZATION OF MEMBRANE PROTEINS IN LIPIDIC MESOPHASES

Martin Caffrey

Membrane Structural and Functional Biology Group, Trinity College Dublin, Ireland
martincaffrey@tcd.ie

One of the primary impasses on the route that eventually leads to membrane protein structure through to activity and function is found at the crystal production stage. Diffraction quality crystals, with which an atomic resolution structure is sought, are particularly difficult to prepare currently when a membrane source is used. The reason for this lies partly in our limited ability to manipulate proteins with hydrophobic/amphipathic surfaces that are usually enveloped with membrane lipid. More often than not, the protein gets trapped as an intractable aggregate in its watery course from membrane to crystal. As a result, access to the structure and thus function of tens of thousands of membrane proteins is limited. The health consequences of this are great given the role membrane proteins play in disease; blindness and cystic fibrosis are examples. In contrast, a veritable cornucopia of soluble proteins have offered up their structure and valuable insight into function, reflecting the relative ease with which they are crystallized. There exists therefore a pressing need for new ways of producing crystals of membrane proteins. In this presentation, I will review the field of membrane protein crystallogenesis with reference to the latest update of the Membrane Protein Data Bank (www.mpdb.tcd.ie). Emphasis will be placed on crystallization approaches which make use of the lipidic systems. In my talk I will describe these methods and our progress in understanding how they work at a molecular level. The practicalities of implementing these methods in low- and high-throughput modes will be examined. A practical demonstration of the lipidic cubic phase or in meso method will be given at the FEBS Lab Exercises on Saturday, June 23.
OBSERVATION OF CRYSTAL GROWTH / SEEDING

Terese Bergfors

Dept. of Cell and Molecular Biology, Biomedical Center, Box 596, Uppsala University,
753 29 Uppsala, Sweden
terese@xray.bmc.uu.se

One of the most common questions in macromolecular crystallization is: “I have some small (or ugly, or bad) crystals now, what can I do to improve them?” One powerful optimization tool is seeding. In this technique, crystals grown previously are introduced into new drops. The “seed crystals” can be microcrystals, spherulites, or even other solid phases found by the initial screening experiments. In the new drops, these “seed crystals” become ready-made templates for the protein molecules to accumulate on. This eliminates the need for spontaneous nucleation, which is a kinetically limiting step in crystallogenesis.

There are many ways of performing seeding and this lab exercise will demonstrate two of them, streak-seeding (Stura, 1999) and jab-seeding (Mac Sweeney & D’Arcy, 2009). The exercises demonstrate seeding by manual methods, but seeding can also be done robotically, as will be demonstrated by Patrick Shaw Stewart. The purposes of the experiments are:

- To learn how to transfer seeds by easy, fast, and technically simple manual methods.
- To observe the effects of decreasing protein concentration on the seed growth.
- To observe the effects of diluting the number of seeds.
- To compare streak-seeding and jab-seeding as methods for microseed transfer.


Seeding Lab Exercises enclosed at the end of book of abstracts on page 48.

CRystallization in microfluidic chips

Claude Sauter

Architecture et Réactivité de l’ARN, Université de Strasbourg, IBMC, CNRS, 15 rue René Descartes, 67084 Strasbourg, France
c.sauter@ibmc-cnrs.unistra.fr

Microfluidic devices offer many advantages for crystal growth: they can easily handle nano-volumes of solutions and have been used to miniaturize crystallization methods such as batch, free interface diffusion, counter-diffusion or dialysis. Thus, they provide a very efficient parallelization of crystallization assays for high throughput screening applications. In addition, due to their small cross-section, microfluidic channels and chambers constitute convection-less environments that are a priori favorable to the growth of high quality crystals. Finally, some of these microfluidic chips are compatible with the in situ crystal analysis by X-ray diffraction. Different microsystems, their function and manufacturing process will be presented during the workshop. Participants will have the opportunity to perform crystallization assays and to grow crystals of model proteins in such devices.

User manual for XZ™ crystallization plates enclosed at the end of book of abstracts on page 63.
E6

CRystallization Under Oil - Unconventional Crystallization Techniques for Screening and Optimisation

Naomi E. Chayen and Lata Govada

Department of Bio-Molecular Medicine, Faculty of Medicine, Imperial College London, London SW7 2AZ, UK
l.govada@imperial.ac.uk

Dear Course Participants,
I have written down some of the methods that I will talk about in the workshop in the form of exercises, to give you step by step protocols of setting up the experiments.

The methods are mostly for optimization – in cases that you get poor crystals that need improvement. Some can also be applied for screening as mentioned in the specific protocols.

These methods have been successful in yielding diffraction quality crystals of a variety of proteins in cases where conventional methodology failed. I have used lysozyme and trypsin as models in some of the protocols in order to have examples and results that you can see within a short time, but of course, the aim is to use these techniques with your problem proteins.

There is no magic bullet to solve all the crystallization problems however having a portfolio of different techniques is very helpful.

GOOD LUCK!
Naomi

Exercises 1-7 enclosed at the end of book of abstracts
on page 51

E7

Additive Experiments, Microseeding, Grids for Optimization; Optimization Techniques Using Automatic Contact Dispensing

Patrick Shaw Stewart

Douglas Instruments Ltd, Douglas House, East Garston, Hungerford, Berkshire, RG17 7HD, UK
patrick@douglas.co.uk

Patrick Shaw Stewart will demonstrate several techniques for improving protein crystallization after finding initial crystallization conditions. These include additive experiments (which can also be used for initial screening), microseeding (including the MMS technique), microbatch-under-oil, and simple 2-d grids for optimization. Students can use model proteins or bring their own proteins.

One of the lab exercises will be microseeding with random crystallization screens. This method will often give additional hits and reduce the need for crystal optimization. To find out more please see the references and link below.

If you have proteins that you would like to crystallize by microseeding, please bring them. An appropriate amount to bring would be around 60 µl. We normally set up drops using 0.3 (protein) + 0.2 (reservoir) + 0.1 (seed stock). So 60 µl of protein would allow you to set up two 96-well plates.

If you’re short of protein, however, we can have a go with just 10 µl.

Please also bring crystals if you have any, to make the seed stock. If your crystals were grown by vapor diffusion, remove the reservoir solutions, cover the drops with pure paraffin oil, and reseal with tape. This will preserve them during your trip.

If your protein crystallizes quickly, we can set up crystals on the first day (with the robot) and harvest them to make the seed stock on the second day.

We don’t recommend transporting the seed stocks at room temperature because the seed crystals may dissolve.

We are providing several commercially available crystallization screens. However, if you know the condition your protein crystallizes in, we recommend to bring your crystallization buffer as well.


www.douglas.co.uk/mms.htm

© Krystalografická společnost
DYNAMIC LIGHT SCATTERING

Karsten Dierks¹, Arne Meyer¹, Howard Einspahr² and Christian Betzel³

¹XtalConcepts, Stellinger Steindamm 7, Hamburg, Germany
²67 Green Avenue, Lawrenceville, NJ 08648, USA
³Institute of Biochemistry and Molecular Biology c/o DESY, 22603 Hamburg, Germany
dierks@xtal-concepts.de

Dynamic Light Scattering (DLS) is an application to determine particle sizes in solution. It can be understood as a method to measure the velocity of the Brownian motion. This velocity is related to the particle size by the Stokes-Einstein equation, which links diffusion velocity, viscosity and temperature of the solvent with particle size (radius). In order to obtain correct values, therefore it is necessary to know these parameters.

Part of the instrument is a laser diode. The laser light is focused in the sample cell. The particles scatter the laser light in all directions. Because the particles are in constant motion, the intensity of the scattered light, as viewed from a certain direction, changes accordingly by alternating constructive and destructive interference. The velocity of the resulting intensity fluctuations depends on the Brownian motion and it is correlated with the hydrodynamic radius of the particle. By analyzing these intensity fluctuations, it is possible to determine the particle velocity and hence, via above equation, the desired particle radius. The scattered light is consequently collected and registered by a photo multiplier tube, counting single photons. The output signal of this tube is directed to an autocorrelator, which finally calculates the autocorrelation function (ACF).

TRACE FLUORESCENT LABELING FOR PROTEIN CRYSTALLIZATION SCREENING

Marc Pusey

iXpressGenes, Inc., 601 Genome Way, Huntsville, AL 35806, USA
marc.pusey@ixpressgenes.com

Fluorescence can be a powerful tool to aid in the crystallization of proteins. Fluorescent labeling methods require removal of unbound probe prior to use of the protein in the screening experiment. For the trace labeling approach the protein is covalently derivatized with a high quantum yield visible wavelength fluorescent probe. The final probe concentrations are typically < 0.25% of the protein molecules labeled, which has been shown to not affect the crystal nucleation or diffraction quality (Forsythe et al., 2006). The labeled protein is then used in a plate screening experiment in the usual manner. As the most densely packed state for the protein is the crystalline form, then crystals show under fluorescent illumination as the brightest objects in the well. Crystals are clearly seen even when buried in precipitate. This approach also finds ‘hidden’ leads, in the form of bright spots, with ~30% of those lead found being optimized to crystals in a single pass optimization trial. Visible fluorescence also enables selection of conditions that bypass interfering substances and greatly reduces materials considerations.


E10

PUBLICATION OF SCIENTIFIC RESULTS WITH EMPHASIS ON CRYSTALLIZATION COMMUNICATIONS

Howard Einspahr

IUCr Journals, 67 Green Avenue, Lawrenceville, New Jersey 08648 USA
hmeinspahr@yahoo.com

The demonstrator will be available (1) to consult on all matters related to drafting and publishing of crystallography papers and (2) to demonstrate new web pages designed to help authors. This can include actual editing of draft manuscripts brought by participants. Participants who would like help with drafts should visit the new web pages for authors at http://publbio.iucr.org. If you have not registered with the World Directory of Crystallographers, which is required for access, you may do so at http://www.iucr.org/people/wdc/help/online-services.

L11

PROTEIN CRYSTALLIZATION USING THE GCB©

José A. Gavira

Laboratorio de Estudios Cristalográficos, IACT, CSIC - U.Granada, PTCS., Avd. del Conocimiento s/n, 18100 Armilla (Granada), Spain
gavi@lec.csic.es

The Granada Crystallization Box Domino (GCB®) will help us to implement several experiments using the counterdiffusion technique (Protocols 1 & 2). In this technique the solutions of protein and precipitant agent are set to diffuse against each other resulting in a spatial-temporal gradient of supersaturation along the length of the capillary. Unlike conventional techniques, different supersaturation conditions leading to protein precipitation, nucleation and crystal growth can be obtained consecutively in a single experiment [1-3].

Counter-diffusion has proved its ability to increase protein crystal quality and it is regularly used for optimization after initial crystallization conditions have been found with other techniques. Using the pre-filled GCB Domino® we will test the potential of the counter-diffusion technique for initial crystallization screening in capillaries of 0.1 mm inner diameter (less than 400 nl for capillary length of 50 mm) and optimization in capillaries of 0.2 mm. We will do optimization of lysozyme or thaumatin crystals varying only protein concentration. The effect of protein concentration on crystal density and size should be observed.

Other implementation of the counterdiffusion techniques (crystallization in gel) and its applications will be commented.


Crystallization protocols 182 are enclosed at the end of book of abstracts on page 65.
Neutron Macromolecular Crystallography (NMC) is the prevalent method that accurately determines the positions of hydrogen atoms in macromolecules. There are two major limiting factors in determining protein structure with neutron diffraction; one is obtaining large crystal volumes and the other is the extended experimental duration. This lecture will demonstrate that crystallizing hyperthermophilic proteins by counter-diffusion crystallization can produce suitable protein crystals for NMC. Counter-diffusion crystallization in restricted geometry, such as in capillary tubes, allows diffusion to govern mass transport processes so that a spatial-temporal gradient of supersaturation is formed along its length. Consequently, proteins can crystallize in an optimized supersaturated condition where a single crystal can fill up the diameter of the capillary. We have employed the same process in capillary vessels having diameters exceeding 1mm while minimizing the ratio between buoyant and viscous forces. As a result, protein crystals suitable for neutron crystallography can be grown in this configuration with volumes greater than 1mm$^3$ while eliminating invasive crystal manipulation. Using crystals grown by this method, we have undergone neutron diffraction analysis of recombinant proteins from a hyperthermophilic archaeon. These proteins are excellent crystallization targets because of their thermal and mechanical stability and high propensity to crystallize. As an initial proof of principle, a novel inorganic pyrophosphatase (IPPase) was among the first of these proteins to be grown in large diameter capillaries resulting in a 9mm$^3$ volume crystal. The next generation neutron beamlines at the Spallation Neutron Source (SNS) along with an effective counter-diffusion crystallization procedure can improve structures of a diverse range of proteins of interest to the crystallographic community. Thus the crystallization of macromolecules can grow to volumes never before achieved, potentially leading to enormous increases in the number of neutron crystallographic protein structures.
CRYSTAL OBSERVATION, TESTING, HANDLING, MOUNTING AND CRYOCOOLING

Jiří Brynda, Petr Pachl

Institute of Molecular Genetics, Institute of Organic Chemistry and Biochemistry, Academy of Sciences of the Czech Republic, Prague
brynda@img.cas.cz

For the observation of crystallization experiments and crystal handling a stereomicroscope usually with a polarizing filter is used. Crystal can usually be distinguished from other matters in the droplet by its edges. Birefringence is also a good sign as it indicates an anisotropy in the material along the viewing axis. Birefringence also helps to distinguish small microcrystals or spherulites from amorphous precipitates. Absence of birefringence in protein crystals is a result of a case when a crystal is viewed along an axis, and in case of cubic space groups.

Protein crystals can be easily distinguished from salt crystals by staining with methylene blue or based on the difference in mechanical properties. Protein crystals are very sensitive and can be easily crushed with a needle. The protein content of the crystal can be analyzed by electrophoresis, but the final test obviously is the diffraction pattern...

Crystals of size and quality (monocrystal with size of at least 0.05mm) suitable for diffraction measurements need to be moved from the crystallization solution and mounted on the diffractometer. Protein crystals must always stay in mother liquor during the whole mounting process and the data collection. For X-ray measurement at room temperature the crystal is placed in the thin wall glass or quartz capillary surrounded with liquid vapors. In most cases, however, the cryocooling is the method preferred for crystal handling, diffraction data collection and storage. In addition to the benefits of eliminated radiation damage and increased resolution, the mounting of crystals is greatly simplified compared to the capillary mounting procedure.

Depending on the mechanical properties of the crystal fibers, spatulas or, most widely fiber loops are used. Using the nylon fiber loop, protein crystal is picked up by swiftly moving the loop alongside the crystal from the crystallization mother liquor.

The crystal is held within the loop by a surface tension and after equilibration in cryoprotective buffer must be cooled to cryogenic conditions as soon as possible. A simple and often effective approach is to flash cool the crystal in a goniostat nitrogen stream right on the X-ray device. This technique has the added advantage in leaving the crystal in position for immediate analysis and data collection. An alternative method, rapidly plunging the crystal into a liquid cryogen, also offers several advantages. It reduces the time between mounting the crystal and flash cooling, it produces higher cooling rate and results in more even cooling of both sides of loop-mounted sample. Crystal flash cooled in liquid nitrogen must be placed for data collection in the cold gas stream on a goniostat without any warming.

Once a crystal has been successfully cooled to cryogenic temperature it can be in principle stored for indefinite time. This allows cooling and characterizing crystals on a conventional source in the home laboratory and then storing them until synchrotron time becomes available. Dewars that can be used for transport, including shipment by airplane, are available.

Today you can buy the Crystal Handling Kit from Jena Bioscience (CO-150) which can help you to acquire skills in protein crystallization, crystal mounting and data collection.