

## Poster Sessions

The handling of cryo-cooled protein crystals has been automated by robotics systems, where samples are kept in liquid-nitrogen storage, to be loaded on the goniometer for the experiment and retrieved afterwards. Crystals are automatically centered in the X-ray beam by means of an automated goniometer head and software that applies algorithms that determine the crystal position in three-dimensional space from images taken with high resolution digital microscopes. For the steps that follow, we report on the development of XPRESSO, a new crystal screening and data collection system for macromolecular samples.

The screening process starts out by taking short series of X-ray diffraction images from which the general quality of the crystal is judged by the resolution limit, the mosaicity, the ability to find the unit cell, and the presence of ice rings. User input limits for the unit cell help distinguish between the actual sample and unwanted crystals, such as from buffers or salts co-crystallized with a protein.

For the data collection a strategy is determined based on the screening results. It takes into account the exposure time, sample to detector distance, scan width, and resolution limit, among others. The data is integrated in parallel to the data acquisition, followed by data scaling. Space group determination is the final step. Results are provided as HTML reports, including Matthews coefficient probabilities.

**Keywords:** automation, protein, software

### MS72.P06

*Acta Cryst.* (2011) A67, C660

#### New Tools for Biological Crystallography in the Home Lab.

Marianna Biadene,<sup>a</sup> Matthew Benning,<sup>b</sup> <sup>a</sup>*Bruker AXS GmbH, Karlsruhe, Germany* <sup>b</sup>*Bruker AXS, Madison, WI, (USA)*. E-mail: Marianna.Biadene@bruker-axs.de

Advances in crystallographic hardware and software have enabled structural biologists to investigate more challenging projects. Recent developments have greatly increased the capabilities of in-house diffraction systems providing increased productivity for synchrotron trips and home-lab studies.

We have made recent improvements in source and detector technology which have significantly improved the capability of home-lab systems for both screening and data collection. Developments include next generation microfocus sources which exhibit significantly higher intensity as well as enhanced beam stability. Combined with a new sensor-based active pixel detector, these systems provide a significant improvement in overall performance while offering extremely low maintenance and cost of ownership. A new feature in our PROTEUM software, XPRESSO, offers a completely automated data acquisition and analysis pipeline for macromolecular crystallography. New developments in hardware and software will be discussed.

**Keywords:** microfocus source, detector, automated data acquisition

### MS72.P07

*Acta Cryst.* (2011) A67, C660

#### Graphical user interface for automated crystallography data reduction.

Duncan Sneddon, Graeme Winter, Alun Ashton. *Diamond Light Source Ltd. Diamond House, Harwell Science and Innovation Centre,*

*Didcot. (UK). OX11 0DE.* E-mail: Duncan.sneddon@diamond.ac.uk

With the advances in throughput at synchrotron facility sites capable of yielding up to 15 data sets an hour the automatic reduction of single crystal diffraction data is becoming a routine component on synchrotron beamlines. For the past three years at Diamond Light Source, packages such as fast\_dp (in house development) and xia2[1] have been employed to carry out data reduction immediately after data collection and are automatically triggered without user interaction. Here we describe how our first implementation which was geared towards processing single sweeps of data with no user interaction has been integrated with a graphical user interface that provides finer control to these programs and the ability to process multiple or partial sweeps of data.

[1] G. Winter, *J. Appl. Cryst.* **2010**, 43, 186-190.

**Keywords:** data processing, automation, GUI

### MS72.P08

*Acta Cryst.* (2011) A67, C660

#### Readiness evaluation method for X-ray diffraction data collection systems

Wei Ding, Zhi-Jie Liu, *National Laboratory of Biomacromolecules, Institute of Biophysics, Chinese Academy of Sciences, Beijing, (China)*. E-mail: zjliu@ibp.ac.cn

High quality diffraction data is critical for structure determinations. For macromolecular crystallography, there are many factors that may compromise the final data quality. In addition to crystal quality and data collection strategy, the instrumentation factors, such as x-ray beam quality, goniometry and the quality of detectors, are important for data quality too. Since the data collection systems are composed of many electronic and mechanical units and they have to work synergically at their at least normal performance. But this may not always be true.

In order to develop a quick and simple method to access the overall performance of the X-ray data collection system, we proposed a protocol to test X-ray diffraction facilities' readiness. In this protocol, cubic insulin crystals are used to quickly collect anomalous data at long wavelength such 1.54Å. The weak anomalous signals from the 3 disulfide bonds are used to indicate the accuracy of the whole data collection system. This method has been tested at different data collection systems including both rotating anode based home labs and synchrotron beam lines. The detailed results and analysis will be presented.

**Keywords:** data collection, readiness

### MS72.P09

*Acta Cryst.* (2011) A67, C660-C661

#### New version of CrysAlisPro optimized for automatic macromolecular data collection and processing

Tadeusz Skarzynski, Mathias Meyer and Przemyslaw Stec, *X-ray Diffraction Department, Agilent Technologies, Yarnton, Oxfordshire (UK)*. E-mail: tadeusz.skarzynski@agilent.com

While the majority of macromolecular X-ray data are currently collected using highly-efficient beam lines on an ever increasing number of synchrotrons, there is still a need for low-maintenance, reliable systems for in-house experiments. In addition to crystal screening and optimization of x-ray experiments before a successful synchrotron trip,