

amplified by PCR introducing in all of them the same double restriction sites present in a library of expression vectors, which will add different tags (MBP, GB1, His-tag at the N- or C-terminus) to each construct. All constructs are subsequently tested for bacterial overexpression using different strains and at different culture conditions. The last step consists of a solubility screening for each construct tested at a specific strain and expression condition, by using different buffers designed according to a semi-rational approach. The results from different analysis will be presented.

Keywords: methods, expression, solubility

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Community nominated projects at the Joint Center for Structural Genomics

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The Joint Center for Structural Genomics (JCSG), funded through the NIH NIGMS Protein Structure Initiative (PSI), has implemented a high-throughput structural biology pipeline, which has delivered more than 1100 structures over the last 10 years. Numerous technological advances have been incorporated to allow efficient, parallel processing of a large numbers of targets from both bacteria and eukaryotes. Until recently, the primary focus of the PSI was to increase structural coverage of protein sequence space, by determining structures from large protein families that lacked any structural representative. However, in 2010, the PSI entered its third phase (PSI-Biology), where the emphasis shifted to make high-throughput structure determination more broadly available to the biological research community. NIGMS has funded several High-throughput Enabled Structural Biology Centers to focus on specific biological topics, as part of the PSI-Biology Network. In addition, individual researchers are able to propose “community-nominated” targets to the PSI through the PSI Structural Biology Knowledgebase (<http://cnt.sbkf.org/CNT/targetlogin.jsp>).

Here we will present specific examples of “community-nominated” projects at the JCSG, highlighting the diversity and scope of these biological projects. These projects are sometimes initiated as a follow-up to structures previously solved by the JCSG, where we team up with biologists to perform biochemical studies aimed at elucidating the functional role of these novel proteins. Frequently, the scope of the project is then expanded to target other related proteins. In other cases, biologists come to us with exciting new projects, targeting structures of novel and biologically significant proteins or protein families, proteins involved in interesting pathways, essential proteins but still of unknown function identified through microarray studies, or protein-protein/protein-DNA complexes. Results from some of these projects will be outlined.

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Keywords: high-throughput structural biology, collaborative biological research, community targets

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X-ray fluorescence tool for rapid in-house evaluation of heavy atom derivatives

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The MIR phasing method still plays a crucial role in the structure determination of a novel protein. However, to acquire adequate heavy atom derivatives, one must repeatedly go through the tedious procedure, transferring crystals to various heavy atom solutions with different concentration for different soaking time ranging from minutes to days [1]. Moreover, the result of soaking can only be obtained by checking a difference Patterson map that requires a full data collection. If one can assess the presence of heavy atoms in a protein crystal prior to a data collection, it will greatly reduce the amount of work required to obtain heavy atom derivatives.

To meet this need, we developed an X-ray fluorescence (XRF) tool that can be used to detect heavy atoms introduced by soaking. This XRF tool is composed of an SDD, an XYZ stage, counting circuits and software, and can be added to an existing in-house single crystal system.

We performed some experiments to evaluate this tool. The tool was placed on a Rigaku MicroMax 007HF rotating anode X-ray generator equipped with a VariMax Mo multilayer optic. Some lysozyme crystals were soaked in K_2PtCl_4 solution and subjected to both XRF and X-ray diffraction (XRD) measurements. We found that those crystals showing clear XRF spectra of Pt tend to have discrete peaks corresponding to Pt's on Harker sections and have higher occupancy when refined against ΔF 's derived from the XRD measurement.

This XRF tool can be used not only to eliminate unwanted “Native” crystals in-house prior to beam time, but also to estimate amount of heavy atoms introduced in a protein crystal by soaking.

[1] T.L. Blundell, L.N. Johnson, *Protein Crystallography, London: Academic Press*. 1976, 183-239.

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The Joint Center for Structural Genomics (JCSG; www.jcsg.org) is one of the four PSI:Biological Center for High-Throughput (HT) Structure Determination. Our mission in this 3rd phase of the Protein Structure Initiative (PSI) is to extend HT Structural Biology (HTSB) to a broader biological community. To meet these goals, each of the HT centers has been partnered with Biological Centers that present challenging new opportunities for HTSB, in addition to our internal biomedical-theme projects. The JCSG is in its 12th year of operation and provides a robust and flexible HTSB platform that is applicable to a large variety of targets from both bacteria and eukaryotes. Our main approach involves processing large numbers of targets via an extensive combination of bioinformatics and biophysical analyses to efficiently characterize each target in order to optimize its path through our pipeline. In close collaboration with our PSI:Biological Partnerships and