MS51.P12

Acta Cryst. (2011) A67, C554

Assembly and Solution Structure of the Retromer Complex

Suzanne Norwood, ^aDaniel Shaw, ^aNathan Cowieson^b James Swarbrick^c and Brett Collins, ^a ^aInstitute for Molecular Bioscience, University of Queensland, St Lucia, Queensland. ^bAustralian Synchrotron, Clayton, Victoria. ^cFaculty of Pharmacy and Pharmaceutical Sciences, Monash University, Parkville, Victoria (Australia). E-mail: s.norwood@uq.edu.au

Retromer is a peripheral membrane protein complex that plays a critical role in a broad range of physiological, developmental and pathological processes by mediating retrograde transport of proteins from endosomes to the *trans*-Golgi (TGN) network. Wnt signalling, toxin transport and amyloid production in Alzheimer's disease are just some of the processes known to be regulated by retromer-mediated trafficking.

Mammalian retromer consists of a core heterotrimeric cargorecognition subcomplex associated with a membrane-targeting dimer of sorting nexins. The core subcomplex consists of vacuolar protein sorting (VPS)26, VPS29 and VPS35 subunits that play different roles in complex stabilisation and in binding to transport cargo and regulatory proteins. The composition of the membrane-binding subcomplex is inconclusive but consists of a homo- or heterodimer of sorting nexins, SNX1, SNX2, SNX5 and SNX6. These BAR (Bin/Amphiphysin/Rvs)domain proteins can induce the formation of high curvature membrane tubules through the formation of a polymerised helical coat.

We have used small-angle X-ray scattering (SAXS), X-ray crystallography, nuclear magnetic resonance (NMR) and isothermal titration calorimetry to elucidate a qualitative and quantitative model of retromer assembly [1]. In our proposed model, VPS35 forms an extended, gently curved structure composed of alternating HEAT-like helical repeats. VPS26 and VPS29 bind to distal ends through N- and C-terminal regions of VPS35, respectively, to form a stable trimeric core assembly. Results from thermodynamics experiments have shown that VPS29 and VPS26 bind to VPS35 completely independently of each other, confirming that VPS35 plays the role of central scaffold and VPS29 and VPS26 do not form any contact with each other. Intriguingly, the core trimeric complex is able to form a symmetric dimer, which may have implications for functional interactions *in vivo*.

Solution structures of SNX1 and SNX2 homodimers have been determined using SAXS. The SNX dimer associates with cellular membranes enriched in phosphatidylinositol 3-phosphate and 3,5-diphosphate and is believed to polymerise into a helical protein coat. NMR studies have confirmed that VPS29 coordinates the binding of the core subcomplex to this membrane remodelling complex. This coupling of cargo binding and membrane tubulating events enable retromer to recruit cargo molecules into a tubular endosome–to-Golgi transport carrier.

[1] S. Norwood, D. Shaw, N. Cowieson, D. Owen, R. Teasdale, B. Collins, *Traffic* **2010**, *12(1)*, 56-71.

Keywords: saxs, retromer, sorting nexin

MS51.P13

Acta Cryst. (2011) A67, C554

 $On-axis\ single-crystal\ raman,\ fluorescence\ and\ UV/Vis\ microspectroscopy\ at\ the\ MX\ spectrolab\ of\ the\ swiss\ light\ source$

Martin R. Fuchs, Florian S.N. Dworkowski, Guillaume Pompidor, Vincent Thominet, Clemens Schulze-Briese, *Swiss Light Source, Paul Scherrer Institut, Villigen, Switzerland*. E-mail: martin.fuchs@psi.ch

By the in-situ combination of macromolecular crystallography (MX) with complementary optical spectroscopic techniques, the specificity of the information obtained from the crystallographic electron density maps can be greatly enhanced. Even at high resolution, e.g. the chemical state of co-factors and many structural ambiguities can only be resolved by the additional complementary information provided by UV/Vis absorption, fluorescence, Raman and Resonance Raman spectroscopy.

To exploit the strengths of these spectroscopic methods, a multimode micro-spectrophotometer (MS2) has been developed at the Spectroscopy Laboratory at beamline X10SA of the Swiss Light Source. An upgrade to the first installment of the spectrometer [1], which supported the UV/Vis absorption and fluorescence modes, the new version now is in regular user operation also for non-resonant and Resonance Raman operation. Raman spectroscopy in particular can provide specific information e.g. on ligand binding. Via difference Raman spectroscopy one can follow chemical changes such as ligand - active site interactions in soaking experiments. Resonant excitation enables monitoring of chromophore-related vibration bands.

The unique on-axis geometry of the instrument, with collinear alignment of both the X-ray and all optical axes, ensures the optimal overlap of the X-ray irradiated crystal volume and the area sampled by the spectrometer. Systematic errors from imperfect overlap of the sampled volumes can thus be avoided, which can become a problem in a more traditional off-axis alignment scheme. By utilizing exclusively reflective components for coupling and focusing elements, a broad spectral bandwidth down to 250 nm can be achieved. The MX SpectroLab, a separate off-line spectroscopy laboratory with a full goniometer setup is available for preparation of experiments on the beamline as well as for single crystal spectroscopy not requiring X-ray irradiation.

We present selected results obtained with the different modes of the spectrometer and will discuss the current instrument design as well as the final version of the spectrometer which is being designed to remain always online at the beamline.

[1] R.L. Owen, A.R. Pearson, A. Meents, P.Boehler, V. Thominet, C. Schulze-Briese *Journal of Synchrotron Radiation* **2009**, *16*, 173-182.

Keywords: spectroscopy, raman, UV/Vis absorption

MS51.P14

Acta Cryst. (2011) A67, C554-C555

Biophysical and structural characterization of the stress response protein CpxP

<u>Gina L. Thede</u>^a David C. Arthur,^a Ross A. Edwards,^a Daelynn R. Buelow,^b Tracy L. Raivio,^b J.N. Mark Glover,^a ^aDepartment of Biochemistry, and ^bDepartment of Biological Sciences, University of Alberta, Edmonton, Alberta, (Canada). E-mail: gthede@ualberta.ca

Bacteria have evolved mechanisms to sense and adapt to continually changing conditions, allowing them to survive in a wide range of environments. In Gram-negative bacteria, such as *Escherichia coli*, the Cpx two-component signal transduction pathway senses misfolded envelope proteins and subsequently activates the expression of periplasmic proteases and folding factors to maintain protein integrity in the envelope. The Cpx stress response system consists of a membrane-localized sensor histidine kinase CpxA, the response regulator CpxR, and the novel periplasmic accessory protein CpxP. CpxP is a key regulator of the Cpx response and has been found to inhibit the pathway, likely through interaction with CpxA, and is required for the degradation of some misfolded proteins by the periplasmic protease DegP.

Since CpxP has no homologues of known function, we have initially focused on its biophysical and structural characterization. Using multi-angle laser light scattering (MALLS), small-angle X-ray scattering (SAXS) analysis, and formaldehyde-mediated cross-linking experiments, we show that full-length *E. coli* CpxP is a dimer *in vivo* as well as in pathway inactivating (pH 5.8) and activating (pH 8.0) conditions *in vitro*. Far-UV circular dichroism (CD) was used to demonstrate that CpxP is mainly α -helical, while near-UV CD and SAXS revealed that the protein may undergo a small structural adjustment in response to a pathway-inducing stimulus (pH 8.0).

The crystal structure of CpxP, determined to 2.85 Å resolution, revealed an antiparallel dimer of intertwined α -helices with a highly basic concave surface. Each protomer consists of a long, hooked and bent hairpin fold with conserved LTXXQ motifs forming two diverging turns at one end. Three of six previously characterized *cpxP* loss-of-function mutations, M₅₉T, Q₅₅P, and Q₁₂₈H, likely result from a destabilization of the protein fold, whereas the R₆₀Q, D₆₁E, and D₆₁V mutations may alter interactions important for the signalling or proteolytic adaptor functions of CpxP.

Keywords: bacterial, biocrystallography, SAXS

MS51.P15

Acta Cryst. (2011) A67, C555

Structure of monomeric and dimeric Sgt1 protein from *Hordeum vulgare* in solution

Michal Taube,^a Joanna Pienkowska,^b Artur Jarmolowski,^c Maciej Kozak,^a ^aDepartment of Macromolecular Physics, Faculty of Physics, Department of Cell Biology, Institute of Experimental Biology, Faculty of Biology Department of Gene Expression, ^bInstitute of Molecular Biology and Biotechnology, Faculty of Biology, ^eA. Mickiewicz University, Poznan, (Poland). E-mail: mtaube@amu.edu.pl

Sgt1 (Suppressor of G2 allele of skp1) is a conserved eukaryotic protein that plays many important roles inside the cell [1]. Originally it was discovered as a component of yeast kinetochore assembly and member of SCF ubiquitin ligase complex. Sgt1 is also an interacting partner of Hsp90 molecular chaperone which is important for stability and folding of many key signaling proteins like kinases and steroid hormone receptors. In plants Sgt1 is involved in disease resistance to many pathogens and mutation of Sgt1 gene leads to lost of R protein triggered resistance in many cases. Sgt1 interacts with COP9 signalosome (involved in 26S proteasome protein degradation pathway) and SCF complexes and probably target resistance regulatory elements for degradation in plants. In humans Sgt1 positive regulate Nod1 innate immunity receptor pathway.

Sgt1 consist of five non-enzymatic domains: N-terminal *tetratricopeptide* repeat domain (TPR), middle CS domain, C-terminal Sgt1 specific domain (SGS) and two variable regions (VR1 and VR2) that interacts with many partner proteins[2]. CS domain share structural homology with p23 protein Hsp90 co-chaperone and also interacts with Hsp90. CS domain interacts with CHORD II domain of plant protein Rar1 which is involved in disease resistance. SGS domain interacts with Leucine-rich repeats protein like Barley R protein Mla1 and yeast adenylyl cyclase cdc35p. TRP domain interacts with *Arabidopsis* SRFR1. It is know that Sgt1 form a dimer in low ionic strength solutions and that dimerization is mediated by TPR domains.

Here we present the structure in solution of *Hordeum vulgare* Sgt1 in monomeric and dimeric form using small angle X-Ray scattering data measured at beamline X33 (EMBL c/o DESY, Hamburg) and homology modeling. Using MCR-ALS analysis [3] we were able to separate scattering curves from complex mixture of both species and model them using rigid body modeling. Sgt1 form an extended conformation in solution with disordered variable regions in both forms.

Our observation agrees with biological experiments which shows wide spectrum of Sgt1 interacting partners. Such conformation facilitates interaction between proteins. Dimerization may have regulatory role, which depends on physiological state of the cell.

 C. Azevedo, A. Sadanandom, K. Kitagawa, A. Freialdenhoven, K. Shirasu, P.Schulze-Lefert, *Science*, **2002**, 295, 2073-2076 [2] Y. Kadota, K. Shirasu, R.
Guerois, *Trends In Biochemical Sciences*, **2010**, *35*, 199-207 [3] J. Blobel,
P. Bernado, D.I. Svergun, R. Tauler, M.P ons, *Journal of American Chemical Society*, **2009**, *131*, 4378–4386

Keywords: small-angle X-ray scattering, Sgt1, plant disease resistance

MS51.P16

Acta Cryst. (2011) A67, C555

The structural biology knowledgebase – Structures, functions, methods and more

John Westbrook,^a Margaret Gabanyi,^a Yi-Ping Tao,^a Raship Shah,^a David Micallef,^a William McLaughlin,^b Torsten Schwede,^c Konstantin Arnold,^c Lorenza Bordoli,^c Paul Adams,^d Wladek Minor,^c Helen Berman,^a *aRutgers, the State University of New Jersey, Piscataway, NJ, (USA).* ^bThe Commonwealth Medical College, Scranton, PA, (USA). ^cUniversity of Basel, Basel, Switzerland, ^dLawrence Berkeley National Labs, Berkeley, CA, (USA). ^eUniversity of Virginia Medical School, Charlottesville, VA, (USA). E-mail: comments@sbkb.org

The Structural Biology Knowledgebase [1] (SBKB, URL: http:// sbkb.org) is a free online resource designed to combine all protocols and results of the structural genomics and structural biology efforts with information from the biological community in order to have a better understanding living systems and disease. We will present examples of how to navigate the SBKB and how to use its new interface and tools to enable biological research. For example, a protein sequence or PDB ID search will provide a list of protein structures from the Protein Data Bank, associated biological descriptions (annotations), homology models, structural genomics protein target information, experimental protocols, and the ability to order available DNA clones. Text searches find structures, annotations, publications, and technology reports created by the Protein Structure Initiative's high-throughput research efforts. Web tools that aid in bench top research, such as TargetTrack, the new target and protocol database (formerly TargetDB and PepcDB), and Sequence Comparison and Analysis tool for protein construct design, will also be demonstrated. Created in collaboration with the Nature Publishing Group, the Structural Biology Knowledgebase Gateway provides a research library, editorials about new research advances, news, and an events calendar also present a broader view of structural genomics and structural biology. The SBKB is funded by the Protein Structure Initiative/NIGMS.

 M.J. Gabanyi, P.D. Adams, K. Arnold, L. Bordoli, L.G. Carter, Flippen-J. Andersen, L. Gifford, J. Haas, A. Kouranov, W.A. McLaughlin, D.I. Micallef, W. Minor, R. Shah, T. Schwede, Y.P. Tao, J.D. Westbrook, M. Zimmerman, H.M. Berman *J Struct Funct Genomics*. 2011, in press.

Keywords: database, function, methodology

MS51.P17

Acta Cryst. (2011) A67, C555-C556

Combination of in-situ optical spectroscopy and macromolecular crystallography