- [4] Robinson A.J., Overy C., Kunji E.R.S. (2006) Proc. Natl. Acad. Sci. U.S.A. 105: 17766–17771.
- [5] Robinson A.J., Kunji E.R.S. (2006) Proc. Natl. Acad. Sci. U.S.A. 103: 2617–2622.
- [6] Kunji E.R.S., Robinson A.J., (2006) Biochim. Biophys. Acta 1757: 1237–1248.
- [7] Nelson D.R., Felix C.M., Swanson J.M. (1998) J. Mol. Biol. 277: 285–308.

doi:10.1016/j.bbabio.2010.04.141

# 3L.4 Structure based study of the functionality of NhaA in pH and $\mathrm{Na^{+}}\ \mathrm{homeostasis}$

Etana Padan

Biological Chemistry, Institute of Life Sciences, Hebrew University, Jerusalem 91904, Israel E-mail: etana@vms.huji.ac.il

 $Na^+/H^+$  antiporters are essential for homeostasis of  $Na^+$ .  $H^+$  and volume and are critical to cell viability. The crystal structure of Escherichia coli NhaA determined at pH 4 has provided insights into the mechanism of activity of a pH-regulated  $Na^+/H^+$  antiporter [1,2]. The structural fold of NhaA is novel; six of the twelve transmembranes (TM) form an inverted repeat of which two TMs are interrupted in the middle of the membrane forming a unique electrostatic organization, important for activity. Two funnels, a deep cytoplasm-facing and a shallow periplasm-facing are separated by a barrier. The functional unit of NhaA is a monomer [3] with a "pH sensor" separated from the active site. Novel structure-based experimental and computational approaches demonstrate that amino acid residues in TM II contribute to the cation pathway of NhaA and its unique pH activation between pH 6 and 8.5 [4,5]: 1) the highly conserved residues of TM II are located on one side of the helix facing either the cytoplasmic or periplasmic funnels of NhaA. 2) Cys replacements of the conserved residues and measuring their antiporter activity in everted membrane vesicles identified new functional important residues. 3) Several of the Cys replacements were significantly alkylated by a membrane permeant probe implying the presence of water-filled cavities in NhaA. 4) Several Cys replacements were modified by MTSES and/or MTSET, membrane impermeant, negatively and positively charged reagents, respectively, that could reach the Cvs replacements only via water filled funnel(s). Remarkably, MTSES but not MTSET repaired the mutant D65C implying the importance of Asp65 negative charge for pH activation of NhaA. The crystal structure of NhaA allowed to model the eukaryotic NHE1 [6] and NHA2 [7].

#### References

- Hunte C., Screpanti M., Venturi M., Rimon A., Padan E., Michel H. (2005) *Nature* 534: 1197–1202.
- [2] Padan E. (2008) Trends Biochem. Sci. 33: 435-443.
- [3] Herz K., Rimon A., Jeschke G., Padan E. (2009) J. Biol. Chem. 284: 6337–6347.
- [4] Olkhova E., Hunte C., Screpanti E., Padan E., Michel H. (2006) Proc. Natl. Acad. Sci. U.S.A. 103: 2629–2634.
- [5] Herz K., Rimon A., Olkhova E., Kozachkov L., Padan E. (2007) J. Biol. Chem. 285: 2211–2220.
- [6] Landau M., Herz K., Padan E., Ben-Tal N. (2007) J. Biol. Chem. 282 37854–37863.
- [7] Schushan M, Xiang M, Bogomiakov P, Padan E, Rao R, Ben-Tal N. (2010) J. Mol. Biol. **396** 1181-1196.

## doi:10.1016/j.bbabio.2010.04.142

### **3L.5** Novel secretory pathway pumps

Lisbeth Rosager Poulsen, Rosa Laura López-Marqués, Danny Mollerup Sørensen, Thomas Pomorski, Morten Buch Pedersen, Michael Gjedde Palmgren Centre for Membrane Pumps in Cells and Disease—PUMPKIN, Danish National Research Foundation, Department of Plant Biology and Biotechnology, University of Copenhagen, DK-1871 Frederiksberg C, Denmark

*E-mail:* palmgren@life.ku.dk

The super family of P-type ATPases constitutes a large class of membrane proteins involved in the active transport of ions and lipids across biological membranes. Among the prominent members are the Na<sup>+</sup>/K<sup>+</sup>-ATPase, the Ca<sup>2+</sup>-ATPase of sarcoplasmic reticulum and the plasma membrane H<sup>+</sup>-ATPases of plants and fungi. The family can be divided phylogenetically into five distinct subfamilies (P1-P5), each divided into additional subgroups (A, B etc.). The importance of these biological pumps is underlined by the fact that its members are found in all forms of life, from bacteria to man and are involved in fundamental physiological processes, ranging from ion homeostasis and signal transduction to heavy metal and lipid transport across membranes. P4 ATPases constitute the largest P-type ATPase subfamily in eukaryotes but are absent from prokaryotes. They are found in all membranes of the secretory pathway, except the endoplasmic reticulum. P4 ATPases have been associated with flipping of lipids across membranes, a process likely to be the initial event in vesicle budding in the secretory pathway. However, our understanding of lipid translocation, vesiculation and the involvement of P4-type ATPases in these processes is just beginning to emerge and further biochemical characterization of P4-ATPases is required in order to clarify whether these transporters indeed are capable of directly catalyzing transmembrane phospholipid flipping. The B-subunit of P4-ATPases shows unexpected similarities between the  $\beta$ - and  $\gamma$ -subunits of the Na<sup>+</sup>/K<sup>+</sup>-ATPase. It is likely that these proteins provide a similar solution to similar problems, and might have adopted similar structures to accomplish these tasks. P5 ATPases remains the least characterized group of P-type ATPases. They evolved at the branching point between eukaryotic and prokaryotic organisms and thus are associated with the event of compartmentalization in eukaryotes. Localization studies indicate that they reside in internal membrane systems, a hallmark of eukaryotic cells. As no P4-ATPases have been identified in the endoplasmic reticulum, where P5-ATPases are present, it remains an intriguing possibility that in this compartment P5A-ATPases are functional homologues of P4-ATPases.

doi:10.1016/j.bbabio.2010.04.143

## 3P.1 Probing the conformation of the yeast ADP/ATP carrier by fluorescent probes

Valerie L. Brandt, Edmund R.S. Kunji

MRC Mitochondrial Biology Unit, Wellcome Trust/MRC Building, UK E-mail: vlb@mrc-mbu.cam.ac.uk

The mitochondrial ADP/ATP carrier exchanges cytosolic ADP for ATP synthesised in the mitochondrial matrix. During the transport cycle the carrier opens the central substrate binding site to the intermembrane space in the cytoplasmic state and to the mitochondrial matrix in the matrix state [1,2]. The charged residues of the PX [DE]XX[RK] motifs form a salt bridge network on the matrix side of the cavity, when the carrier is in the cytoplasmic state [3,4], whereas the charged residues of the [FY][DE]XX[RK] motifs, which are present on the cytoplasmic side of the cavity, could form a salt bridge network when the carrier is in the matrix state [2]. A structure of the