ENCODING SELECTIVITY OF A TRANSMEMBRANE CHANNEL

Robert M. Stroud*, Peter Nollert, Larry J.W. Miercke, William E.C. Harries, Joe O’Connell
S-964 UCSF Medical School
stroud@msg.ucsf.edu

Membrane channel proteins of the aquaporin family are integral plasma membrane proteins that are highly selective for permeation of specific small molecules, with absolute exclusion of ions including protons or OH- anions and charged solutes, and without dissipating the electrochemical potential across the cell membrane. There are 11 human family members. The family includes channels that conduct water, called aquaporins, and channels that conduct glycerol or other linear polyalcohols (aquaglyceroporins), and ion conducting channels. The E. coli glycerol facilitator (GlpF) was cloned, expressed, purified, and crystallized with its primary permeant substrate glycerol. The crystal structure of GlpF, determined by multiple isomorphous replacement at 2.2 Å resolution illustrate the basis for its selectivity. Glycerol molecules G1, G2, G3 line up in an amphipathic channel in single file. In the narrow selectivity filter of the channel, the glycerol alkyl backbone is wedged against a hydrophobic corner and successive OH groups form hydrogen bonds with a pair of acceptor, and donor atoms. These hydrogen bonds serve to orient the CHX-OH groups within the narrower regions of the channel, and explain the stereo, and enantio selectivity measured for this channel. Two conserved -Asn Pro Ala- motifs meet in the center of the membrane and form a conserved interface between two gene-duplicated segments that each encode three and one half membrane spanning helices around the channel. This structure elucidates the mechanism of selective permeability for linear carbohydrates and suggests how ions and water are excluded. A functional assay was developed, and shows that the channel is both stereo- and enantio-selective. The conservation of sequence shows how the water channels of the same family conduct water and exclude glycerol.

The nature of the channel directly shows how it rejects conduction of protons. This is measured in assays, and tested in collaboration with the Shulten group using molecular mechanics. A comparison of features of the channel itself, and of the space through the center of the tetrameric arrangement of proteins suggests that it is the ion channel in AQP6. Mutations made in the sequence to mimic aquaporins AQP1, and AQP6 lead us to distinguish the basis for water vs. glycerol selectivity, and provide us with a high resolution, tunable structure of an ion channel.

This functional, and structural analysis provides arguably the most detailed bases for understanding of the selectivity of channels for permeant ions, molecules, and organic molecules.
REFERENCES

Submit your manuscripts at
http://www.hindawi.com