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porin- OmpPst1 of *Providencia stuartii* is investigated in its susceptibility for two carbapenem antibiotics imipenem and meropenem. Electrophysiology measurements revealed kinetic parameters of antibiotic binding to the channel. Activity of antibiotics against bacterium determined by microbiological assays correlates with the results obtained from liposome swelling assay and planar lipid bilayer measurements. In addition, the effect of divalent and trivalent cations on antibiotic affinity to the channel is investigated. To follow the exact translocation pathway, molecular modeling provides atomistic details on the interaction of specific side chains of the two antibiotics with the channel residues and the position of affinity sites. Further, mutation of these specific sites in the channel and recording ion current fluctuation in presence of antibiotics reveal the rate limiting interaction for its translocation. Here, we have concluded on the relation between permeation of the antibiotics and resistance acquired by *Providencia stuartii*. Moreover, employing a multidisciplinary approach from MD simulations to protein engineering helps to get a molecular picture. This information further might give insights for rational drug design, for effective uptake of antibiotics through porins- check point of the cell.

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2816-Pos Board B508

Antibiotic Transport through Porins

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The emergence and dissemination of coordinated resistance mechanism (MDR) underline the importance to understand the rate limiting steps in antibiotic action. Here OmpF porin from *E. coli* is used as an example to demonstrate what limits the penetration of carbapenems. Bacteriological kinetic killing assays, temperature dependent ion conductance measurements, and all-atom computer simulations were combined to study interaction and translocation of clinically relevant β -lactam antibiotics through wild type OmpF and two mutants D113A and D121A, where the key residues at the constriction region have been substituted. Expression of these various OmpF mutants in an otherwise porin-null bacterial strain revealed an increase of bacterial susceptibility for the mutants. High-resolution conductance measurements and modulating the temperature indicates lower energy barriers for mutant porin correlating with the microbiological assays. All atom modeling provided a most probable pathway able to identify the relevant side-chain interactions. This combined approach allows identifying rate limiting interaction and suggests possible modification to enhance antibiotic penetration.

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2817-Pos Board B509

Characterization of a Cyanobacterial Outer Membrane Protein: An *E. Coli* Tole Homologue from *Synechocystis* Sp. Pcc 6803

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E. coli TolC (tolerance to colicins) represents an interesting class of outer membrane (OM) proteins, as it has an α -helical periplasmic tunnel and β -barrel membrane region, providing a conduit for export of metabolites and xenobiotics from cell interior to exterior, and import of colicin E1 (1). A TolC homologue, Slr1270 from *Synechocystis* 6803, cloned and expressed in *E. coli*, has > 40% similarity and ~16% identity to *E. coli* and *Pseudomonas* counterparts, and has a similar domain organization. Homology modeling using *Pseudomonas* OprM as template modeled 93% of Slr1270 sequence. The 1581bp slr1270 gene was cloned and overexpressed in *E. coli*. Protein from inclusion bodies, refolded through step-wise dialysis showed major bands at ~55kDa and >150 kDa on SDS-PAGE corresponding to the monomer and trimer respectively and ~300 kDa on CN-PAGE. Purified protein displays a far-UV CD spectrum characteristic of *E. coli* TolC with > 50% α -helix, and

formed channels in planar lipid bilayers with a characteristic single channel conductance of ~50 pS in 0.1M NaCl. The intact protein mass spectrum (LC-MS) with a major peak at 54,489 probably represents a mixture of two species, the TolC product with 1-40 removed and an intact 6-His tag (calculated mass 54,457.1 Da), and a product with 1-38 removed and a 5-His tag (calculated mass 54,490.2 Da) after a single carboxypeptidase event. The small peak at 54,638 Da probably corresponds to TolC product with 1-38 removed and an intact 6-His tag (calculated mass 54,627.3 Da). Peptides 39-76 and 41-76 were recovered from trypsin digests confirming N-terminal heterogeneity. (1) Zakharov, S. D. et al. 2012. Pathways of Colicin Import: Utilization of BtuB, OmpF Porin, and the TolC Drug Export Protein, *Biochem. Soc. Trans.*, 40, 1463-1468.

2818-Pos Board B510

Electrophysiological Analysis of PapC Mutants Provides Insights into the Mechanism of Plug Displacement

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The PapC usher is a twin beta barrel pore of the outer membrane of uropathogenic *E. coli* used for the assembly of the P pilus, a key virulence factor in bacterial colonization of human kidney cells. The usher catalyzes the translocation and ordered addition of folded pilus subunits delivered by the chaperone PapD to the growing pilus. Each PapC monomer is a 24-stranded beta barrel, flanked by N- and C-terminal globular domains and occluded by a large plug domain (PD). An alpha helix and the beta 5-6 hairpin loop are additional structural components that may play a role in controlling plug dynamics. Indeed, structural studies have revealed that the PD is released to the periplasmic side during pilus biogenesis, but the exact mechanism for PD displacement has not yet been elucidated. Several key residues that may be critical for plug stabilization have been proposed. They cluster in regions at the interface of the plug, the barrel, the alpha helix and the hairpin, and appear to belong to either electrostatic or allosteric networks. In order to assess the roles of these residues in plug displacement, we have used planar lipid bilayer and patch-clamp electrophysiology to compare the activity of the wildtype channel with that of mutant channels with either single or multiple alanine substitutions at these sites. Many mutants showed an increased propensity at plug displacement, as witnessed by openings with a conductance similar to that of the previously characterized plugless mutant (~ 3 nS). Others displayed quieter than WT activity. In addition, evidence of modal gating was observed in WT and some mutants. Together, these mutants provide insight into the molecular mechanism of PD displacement for pilus assembly and translocation through the PapC usher.

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Recombinant Production of Human Aquaporin-1 to an Exceptional High Membrane Density in *Saccharomyces Cerevisiae*

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Due to the lipid nature of cellular membranes preventing transport of most solutes between the cytosol and the extracellular environment as well as from the cytosol to the interior of organelles, cellular homeostasis relies on integral membrane proteins allowing selective trans membrane movement of solutes. Aquaporins constitute a family of physiologically very important integral membrane proteins that are found in all three kingdoms, eubacteria, archaea and eukaryotes. As protein channels, they facilitate passive transport of water across cell membranes. In the present study the yeast *Saccharomyces cerevisiae* was exploited as a host for heterologous expression of human aquaporins. Aquaporin cDNA was expressed from a galactose inducible promoter situated on a plasmid with an adjustable copy number. Human aquaporin was C-terminally tagged with yeast-enhanced GFP to quantify functional expression, to determine sub-cellular localization, to estimate *in vivo* folding efficiency and to ease establishment of a purification protocol. We found functional human aquaporin-1 to constitute up to 8.5 percent of total membrane protein content after expression at 15°C in a yeast host over producing the Gal4p transcription factor and growth in amino acid supplemented minimal medium. In-gel fluorescence combined with western blotting showed that low accumulation of correctly folded recombinant Aquaporin-1 at 30°C was due to *in vivo* mal-folding. Reduction of the expression temperature to 15°C almost completely

prevented Aquaporin1 mal-folding. Bioimaging of live yeast cells revealed that recombinant Aquaporin-1 accumulated in the yeast plasma membrane. A detergent screen for solubilization revealed that CYMAL-5 was superior in solubilizing recombinant Aquaporin-1 and generated a monodisperse protein preparation. A single Ni-affinity chromatography step was used to obtain almost pure Aquaporin-1.

2820-Pos Board B512

Aquaporin Trafficking as a Specific Regulatory Mechanism to Adjust Membrane Water Permeability

Gabriela Amodeo, Yaneff Agustin, Jozefkowicz Cintia, Marquez Mercedes, Vitali Victoria, Scochera Florencia, Alleva Karina. IBBEA DBBE, Universidad de Buenos Aires, Buenos Aires, Argentina. Plant aquaporins regulate water fluxes across membranes by enhancing membrane water permeability (P_f). In particular, the plant plasma membrane holds PIPs, one of the largest groups of aquaporins. PIPs are divided in two clusters (PIP1 and PIP2) that disclose intriguing aspects: i) the potential of modulating P_f by PIP1-PIP2 co-expression, distinguished for each PIP showing differential capacity to reach the PM and ii), the faculty to reduce water permeation through the pore after cytosolic acidification, as a consequence of a gating process. Our working hypothesis is that cytosolic pH (gating) and PIP co-expression (trafficking) enhance plasticity to the membrane water transport capacity as a consequence of a PIP1-PIP2 cooperative interaction. Thus, PIP1 cellular trafficking and its effect in water permeability emerge as playing a key role as a regulatory mechanism. To analyze this interaction we used PIP1-PIP2 pairs from different species (*Fragaria ananassa* and *Beta vulgaris*). Our experimental approach included i) designing mutants to alter the PIP-PIP2 interaction by means of site directed mutagenesis; ii) tracking aquaporin localization -at internal structures or expressed at the level of the PM-; and iii) analyzing water transport capacity in control and inhibited (medium acidification) conditions by means of measuring P_f in *Xenopus* oocytes. Our finding support evidences in agreement with the concept that PIP2 and PIP1 interact to form functional heterooligomeric assemblies, and thus the composition of these PIP assemblies determines their functional properties. As PIP1 alone is not able to reach the plasma membrane its contribution to enhance water permeability is associated to its translocation and interaction with a PIP2. This regulatory mechanism seems to be present in different vascular plants. This information is integrated in a proposal for water transport pathways including the organs where this PIPs are present.

2821-Pos Board B513

Selectivity Filter Scanning of the Human Voltage Gated Proton Channel HhV1

Deri Morgan¹, Boris Musset^{1,2}, Vladimir V. Cherny¹, Susan M.E. Smith³, Kethika Kulleperuma^{4,5}, Sindhu Rajan⁶, Régis Pomès^{4,5}, Thomas E. DeCoursey¹.
¹Molecular Biophysics & Physiology, Rush University, Chicago, IL, USA, ²ICS-4 Zelluläre Biophysik, Institute of Complex Systems, Jülich, Germany, ³Department of Biology and Physics, Kennesaw State University, Kennesaw, GA, USA, ⁴Molecular Structure and Function, Hospital for Sick Children, Toronto, ON, Canada, ⁵Department of Biochemistry, University of Toronto, Toronto, ON, Canada, ⁶Medicine, University of Chicago, Chicago, IL, USA. Extraordinary selectivity is crucial to all proton conducting molecules, including the human voltage gated proton channel, hH_v1, because [H⁺] is minuscule. Here we use "selectivity filter scanning" to elucidate the molecular requirements for proton specific conduction in hH_v1. Asp¹¹², in the middle of the S1 transmembrane helix, is essential to the WT channel selectivity [Musset et al., 2011. *Nature* 480:273-277]. We neutralized Asp¹¹² by mutating it to Ala (D112A), then introduced Asp at each position along S1 from 108 to 118, searching for "second site suppressor" activity. All mutants except for D112A/V109D lacked even the anion conduction exhibited by D112A. Proton specific conduction was restored with Asp or Glu at position 116. The D112V/V116D channel resembled WT in selectivity, kinetics, and ΔpH dependent gating. Both R211H and R211H/D112V/V116D were inhibited by internally applied Zn²⁺ when the channel was open, indicating similar S4 accessibility. At position 109 Asp allowed anion permeation in combination with D112A, but did not rescue function in the nonconducting D112V mutant, indicating that selectivity is established external to the constriction at Phe¹⁵⁰. The three positions (109, 112, 116) that permit conduction all line the pore in our homology model, delineating the conduction pathway. Evidently, proton selective conduction requires a carboxyl group to face the pore at a constriction in the external vestibule. Molecular dynamics studies indicate reorganization of ionic networks in response to mutations and suggest that the distribution of charged

groups in the external vestibule modulates charge selectivity. That the selectivity filter functions in a new location helps define local environmental features that produce proton selective conduction.

2822-Pos Board B514

The Permeation Pathway Mechanism in Ciona Intestinal Hv Channel

Ester Otarola¹, David E. Baez-Nieto¹, Gustavo Contreras¹, Osvaldo Yañez², Karen Castillo¹, Peter Larsson³, Ramon Latorre¹, Carlos Gonzalez¹.
¹CINV, Valparaíso, Chile, ²Centro de Bioinformática y Simulación Molecular, Universidad de Talca, Talca, Chile, ³Department of Physiology and Biophysics, University of Miami, Miami, FL, USA. Voltage-gated proton (HV) channels are expressed by different cells type including immune cells, microglia, among others. In other voltage-gated cation channels, the pore-forming domains are S5, S6, and the P loop connecting S5 and S6. Since Hv channels do not contain a S5-S6 region, other parts of the channel must form the pore domain. Interestingly, Hv channel present three gating charges in the S4 followed by an asparagine, N264, highly conserved among all Hv channel. Using non-stationary fluctuation analysis we establish the conductance for the dimer and the monomer in 200 and 100 fS, respectively. Mutations at position S191 (S2) and N264 (S4) modified the unitary conductance of Hv channel. Furthermore, mutants S191E/D and N264R removed the H⁺ current remaining the S4 functioning unaltered, according to voltage-clamp fluorometry experiments. The introduction of an arginine or lysine at position 264 or negative residue at position S191, drastically reduced or abolished the proton currents. Interestingly, mutations of the analogue position S191 of Ciona VSOP, in the voltage sensor domain of a non-conductor Shaker K⁺ channel produce fully functional voltage gated H⁺ channels. In the same way, mutation resembling the position N264 of Ciona Hv in the VSD of the voltage-dependent phosphate Ci-VSP spontaneously entail the formation of a voltage-gated H⁺ channel. Thus, S191 and N264 form the molecular determinants of permeation pathway in Hv channel. Supported by Fondecyt grants ACT 1104 and Fondecyt 1120802 to CG and 1110430 to R.L.

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Mechanism of Non-Selectivity in NAK Channel

Van Ngo, Haina Wu, Stephan Haas, Robert Farley. University of Southern California, Los Angeles, CA, USA. The amino acid sequence of the non-selectivity filter of the bacterial NaK channel (TVGDG) is similar to that of the K-selective KcsA channel (TVGYG), yet NaK conducts both Na⁺ and K⁺ equally well. To investigate the non-selectivity mechanism of NaK, we performed non-equilibrium molecular dynamics simulations using step-wise pulling protocols and Jarzynski's Equality. Two ions in the crystal structure of NaK were kept at site S3 ($z \approx 14.0$ Å) in the filter and in the small vestibule above S3, and one ion for pulling was placed at $z = 0$ Å below the filter entrance. Two sets of simulations with either NaCl or KCl were compared to quantify how the filter dehydrates and conducts the different ions along the z-axis toward the extracellular surface. The simulations show that the entry of Na⁺ into the filter is favored by about 0.5-2 kcal/mol over K⁺ due to easier dehydration of Na⁺. This difference is attributed to the collapse of S3 by 11% in volume on Na⁺, resulting in a higher negative charge density of carbonyl oxygen atoms around Na⁺ than K⁺. When ions enter the filter and displace the ion at S3 (called Na3 or K3) into the small vestibule, the pulled K⁺ in S4 (below S3) is more dehydrated than the pulled Na⁺. K3 and the other K⁺ and water molecules in the filter become significantly more symmetric around the z-axis than Na⁺ and water molecules. This symmetric distribution of K⁺ ions and water favors the movement of K⁺ above S3, offsetting the slight ion selectivity at the filter entrance. We hypothesize that the non-selectivity filter of the NaK channel favors Na⁺ over K⁺ below S3, but becomes more selective for K⁺ than Na⁺ above S3, thus resulting into the non-selectivity as experimentally observed.

2824-Pos Board B516

Statistics of Simulated Ion Channels

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¹Department of Physics, California Polytechnic State University, San Luis Obispo, CA, USA, ²Carmel High School, Carmel, IN, USA, ³Department of Physics, Indiana University Purdue University Indianapolis, Indianapolis, IN, USA. The statistical analysis of ionic currents through biological ion channels is straightforward when single channels are recorded. However, analysis can be intractable when more than one channel is present due to signal overlap. We present a statistical analysis of simulated ion channel recordings when the number of channels is small. Through numerical analysis, we display relationships