

564-Pos**Effects of Tolaasin Inhibitory Factors on Tolaasin Peptide Channel Evaluated by Competition with Zn²⁺**

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Tolaasin is a 1.9 kDa bacterial lipodepsipeptide toxin and forms membrane pores causing brown blotch disease on the cultivated mushrooms. Molecular multimerization of tolaasin is required to form membrane channel. Previously, we showed that various chemicals inhibit the multimerization of tolaasins and thus prevent from brown blotch disease. These chemicals named tolaasin-inhibitory factor (TIF) were applicable to control the disease. Various candidates were chosen from different food additives and they were successful for disease control at 1-100 μ M. Interestingly, after tolaasin molecules formed channels on the erythrocyte membrane and hemolysis started, further hemolyses were stopped as soon as TIF's were added. The added TIF was not washed by centrifugation and addition of fresh HBS solution. These result imply that TIF's are able to inhibit the preformed tolaasin channels. TIF may inhibit tolaasin-induced hemolysis either by plugging the channel pores or by the dissociation of tolaasin multimers. Zn²⁺ is a potent tolaasin inhibitor known to bind to tolaasin channel. In order to characterize the TIF-induced inhibition, competition effect of Zn²⁺ and TIF on tolaasin channel activity was investigated. When Zn²⁺ and TIF-9 were added simultaneously, no additive effects were observed at various concentration combinations. However, in the combinations of Zn²⁺ plus either TIF-11 or TIF-16, no additive effect was measured and the inhibitory effect of Zn²⁺ was reduced in the presence of low concentration of Zn²⁺. In these experiments, the inhibition of hemolysis was dominated by Zn²⁺ concentration. Based on these results, we suggest that TIF has high affinity to tolaasin channel than Zn²⁺ although Zn²⁺ is a strong inhibitor.

565-Pos**Effects of Tolaasin Inhibitory Factors on Tolaasin-Induced Blotch Formation and Hemolysis**

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Tolaasin, 1.9 kDa peptide toxin, is produced by *Pseudomonas tolaasii*. It forms pores on the membrane and thus destroys cellular structure, causing brown blotch disease on the cultivated mushroom. Previously, we showed that the tolaasin-induced pore formation required the molecular multimerization and the multimerization of tolaasin was blocked by the treatment of various tolaasin-inhibitory factors (TIF). This was successful to prevent from causing brown blotch disease. Various TIF's blocked effectively the tolaasin-induced hemolysis at 1-100 μ M. The tolaasin-induced hemolysis was dependent on the temperature and pH; however, TIF always inhibited hemolysis. In the pitting test using mushrooms of *Agaricus bisporus*, tolaasin treatment caused brown blotches on the surface of mushroom and the addition of TIF at 0.1-10 mM completely blocked the blotch formation by tolaasin. Since TIF's were selected from various food additives, the effect and mechanism will be different on the tolaasin-induced hemolysis. In this study, we have measured the synergic effect of various TIF's on the inhibition of tolaasin toxicity. When TIF-9 and TIF-11 were added, tolaasin-induced hemolysis was effectively inhibited. In the presence of both TIF's, the inhibition was increased to 2.5 times compared to the inhibition obtained by any one of them. Similar synergic effect was also measured by the additions of TIF-9 and TIF-16. The inhibition was also increased more than 3 times. However, TIF-10 and TIF-11 did not show any synergic effect. These results showed that there are synergic effects among various TIF's and the effect was dependent on the type and concentration of TIF.

566-Pos**A Biochemical Characterization of the Major Peptides from the Venom of the Giant Neotropical Hunting Ant *Dinoponera Australis***Stephen R. Johnson^{1,2}, Julio A. Copello¹.¹Southern Illinois University - School of Medicine, Springfield, IL, USA,²Carbon Dynamics Institute, LLC, Springfield, IL, USA.

Venom from the "false tocadira" *Dinoponera australis*, a giant Neotropical hunting ant, paralyzes small invertebrate prey and induces a myriad of systemic effects in large vertebrates. HPLC/DAD/MS analyses revealed that the venom has over 75 unique proteinaceous components with a large diversity of properties ranging in size, hydrophobicity, and overall abundance. The six most abundant peptides, demonstrative of this diversity and hereafter referred to as *Dinoponeratoxins*, were *de novo* sequenced by exact mass precursor ion selection and Edman degradation. The smallest peptide characterized, Da-1039, is hydrophilic and has similarities to vasoactive peptides like kinin and bombesin. The two largest and most abundant peptides, Da-3105 and Da-3177, have a 92.9% identity in a 28 residue overlap and share ~50 of their sequence with ponerin G2 (an antimicrobial from another ponerine

ant *Pachycondyla goeldii*). One peptide, Da-1585, is a hydrophilic cleavage product of an amphipathic peptide, Da-2501. The most hydrophobic peptide, Da-1837, is amidated (a PTM observed in one half of the major peptides) and shares homology with poneratoxin, a sodium channel modifier found in the bullet ant *Paraponera clavata*. This study is the first examination of potential pharmacophores from venom of the genus *Dinoponera* (Order: Hymenoptera).

567-Pos**Phloretin Affects the Voltage Gating of Alpha-Hemolysin Channel**

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We studied the effect of dipole modifiers on the channel forming activity of *Staphylococcus aureus* alpha-hemolysin in phosphocholine bilayers bathed in 1 M KCl (pH 7.5). We manipulate bilayer dipole potential (V_d) adding phloretin or phloridzin to reduce (V_d), and 6-ketocholestanol and RH 421 to increase V_d . In the absence of dipole modifiers, an alpha-hemolysin pore fluctuates between high (~1 nS) and low (~0.1nS) conductance states at transmembrane voltage $V \geq |100|$ mV. Addition of 20 μ M phloretin after the channel formation induces transition of the channel to low-conductance state at $V \geq |25|$ mV. Adding phloretin before channel incorporation does not influence its voltage gating. Additions of phloridzin (20 μ M) and RH 421 (10 μ M) in the membrane bathing solution before or after channel formation, or addition of 6-ketocholestanol (50 mol.%) into the membrane composition do not affect the channel gating. We conclude that variation in V_d induced by dipole modifiers does not influence alpha-hemolysin pore gating and the effect of phloretin is likely to be attribute to specific interaction. It is also likely that the binding site for phloretin becomes accessible after alpha-toxin incorporation into the membrane and subsequent pore formation. We suggest that phloretin binding reduces the energy barrier for conformation transition of alpha-hemolysin pore to low-conductance state. The nature of specific interaction between phloretin and alpha-hemolysin channel is discussed. The work is supported in part by RFBR (project 09-04-00883), the Program of Presidium of the RAS "Molecular and Cell Biology", the Grant of Administration of St.-Petersburg for young scientists and State Contract (FAE III372).

568-Pos**Peptide-Gating of Hynacs Shares Features with Proton-Gating of ASICs**Stefan Dürrnagel¹, Andjelko Golubovic², Stefan Gründer¹.¹Department of Physiology, RWTH Aachen University, Germany,²Department of Physiology, University of Würzburg, Germany.

Recently we have identified five new ion channel subunits (HyNaC1-5) from *Hydra magnipapillata* that belong to the ENaC/DEG family of ion channels. We could show that the ion channel made out of HyNaC2/3/5 is gated directly by two endogenous neuropeptides, the HydraRFamides I and II. Prominent human members of the ENaC/DEG family are the proton-gated ASICs, which are involved in pain sensation, learning, memory and salt taste. Here we present our results showing that peptide gating of HyNaCs shares features with proton gating of ASICs. We found that similar to ASICs, HyNaCs were opened by low Ca²⁺ concentrations with a concentration of ~15 μ M leading to half maximal activation. Moreover, the apparent peptide affinity was increased in solutions containing 10 μ M Ca²⁺ compared to 2 mM. Application of HydraRFamides in low Ca²⁺ solution lead to a potentiation of currents indicating that divalent cations block the open channel. In ASICs, Asp₄₃₂ is involved in this open channel block by Ca²⁺. Introducing this mutation in HyNaCs almost completely abolished Ca²⁺ induced currents and the apparent peptide affinity became independent of Ca²⁺ concentration. This mutation also diminished the potentiating effect of low Ca²⁺ on peptide activated currents. These results show that Ca²⁺ is an important modulator of peptide-gating of HyNaCs, like it is on proton-gating of ASICs, and that the Ca²⁺ binding site is conserved between HyNaCs and ASICs.

Protons, which gate ASICs, had a dual effect on HyNaC2/3/5 currents. First, application of pH 5 inhibited open channels. Second, in solutions nominally free of divalent cations strongly increased current amplitude after washout of protons uncovered an activation of the channel by low pH. In summary these results indicate that the gating mechanism is similar between the evolutionary old HyNaCs and ASICs.

569-Pos**Mapping the β -Scorpion Toxin Receptor Site on Voltage-Gated Sodium Channels**Zhongli Zhang¹, Izhar Karbat², Lior Cohen², Todd Scheuer¹, Dalia Gordon²,Michel Gurevitz², William A. Catterall¹.¹University of Washington, Seattle, WA, USA, ²Tel-Aviv University,

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Voltage-gated sodium channels are molecular targets of β -scorpion toxins, which enhance excitability by shifting the voltage dependence of activation