Society of General Physiologists
70th Annual Meeting and Symposium
September 6-11, 2016
Marine Biological Laboratory
Woods Hole, MA

Genetic & Animal Models for Ion Channel Function in Physiology and Disease

Organizers: Andrea Meredith (University of Maryland School of Medicine) and Mark Nelson (University of Vermont)
The Society of General Physiologists would like to acknowledge the sponsors who have helped make this meeting such a prominent gathering of researchers.

2016 SGP Symposium
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SGP was founded in 1946 "to promote and disseminate knowledge and interest in the subject of general physiology" Our international membership is made up of physiologists who work in academia, government, and industry. We have become known for promoting and nurturing research related to membrane transport and ion channels; membrane structure, regulation, and dynamics; and cellular contractility and molecular motors.

The major activity of the Society is its annual symposium at the Marine Biological Laboratory in Woods Hole, which members can attend at a discounted rate. SGP symposia cover the forefront of physiological research and are small enough to maximize discussion and interaction.

Membership in the Society is open to any individual actively interested in the field of general physiology. Categories of membership include regular, young investigator, and emeritus. The Membership year runs annually from January 1st to December 31st.

Visit www.sgpweb.org for more information and to join or renew today!

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Save the Date!

The Society of General Physiologists & The Journal of General Physiology

5th Annual Mixer

Feb 12th 2017
New Orleans, Louisiana
(Sunday of Biophysics Meeting)

Please join us and connect with fellow physiologists, learn about SGP membership benefits, & the upcoming 2017 SGP Symposium

More information will be forthcoming
Important Meeting Site Information

**Registration:** Meeting registration takes place on the first floor of the Swope Center from 2:00 to 9:00 p.m. on Wednesday, September 7. Check-in is at the main desk in Swope Lobby (opposite the door) and your registration packets are at the first table on your left. If you arrive during the night, instructions are posted at Swope Center on how to contact MBL Security who will have your room key.

**Lectures:** Will be held in the Lillie Auditorium, Lillie Building (on MBL Street). Coffee breaks will be set up in the foyer outside of the auditorium, with more mingling and seating space just outside. Timing is tight, so it’s important to be on time for lectures and to return quickly from the 20-minute breaks.

**Poster Instructions:** Poster boards are located on the second floor of Swope Center. Posters may be up to 30” wide x 60” high (portrait-style). Thumbtacks are provided. Poster numbers and titles are listed in the abstracts section of this program. Posters will be presented in two sessions.

- **Poster Session A:** Posters may be displayed after 8:00 a.m. on Thursday, and must come down by Friday noon.
- **Poster Session B:** Posters may be displayed after 3:00 p.m. on Friday afternoon, and must come down by Saturday evening.

**Mixers:** All mixers will be held in the MBL Club at 100 Water Street.

**Meals:** Conference meals are included in your registration, even if you are staying offsite. All meals are served in Swope Dining Hall overlooking Eel Pond. There are three meals a day starting with Dinner on day of arrival and ending with Lunch on the day of departure. There will be breakfast and a boxed lunch on the last day. There are selections for those with dietary restrictions including gluten-free, vegetarian, vegan, and nut- or sugar-free desserts.

*N.B. The dining hall will stop serving lunch at 1:30 p.m.*
**MBL’s Cards:** Every participant is issued an MBL ID card (proximity card) at registration. This card, *to be kept on you at all times*, will provide 24-hour access to Swope, and your dormitory building. Many buildings, including the dormitories, have automatically locking doors. Although Swope will be open until 9 p.m., other buildings will always be locked and you will need your proximity card to gain access. If you should have any questions, both MBL and SGP personnel will be available to answer them during the course of the meeting.

**Internet Access:** MBL has wireless internet service throughout its campus. To use the system simply browse for wireless networks on any wireless enabled device and select "MBL-Guest". Open a web browser and you will be automatically redirected to a login page. Enter the login credentials: user name = mblguest, password = mblguest. In addition, there are 6 terminals from which you can check your email or do a web search in Swope Center. For other computer activity, the Computer Room is located in the MBL Library, Lillie 204.

**Parking:** If you drive to Woods Hole, remember that *parking is by permit only*. You must obtain a parking permit at the registration desk and park ONLY at the Bar Neck parking lot. Directions will be given to you at the registration desk. Do NOT park in any of the other MBL lots or on the street—ticketing and towing are considered a local recreation.

**Luggage Storage:** There is a carriage house outside Swope where luggage can be stored if you have arrived early or need to store your items beyond check out. The front desk staff will assist you in locating the building should you need help.

**Library Usage:** The MBL Library has a world-renowned collection, and visitors are welcome at all times. The Library is staffed Monday through Friday, 8:00 a.m. to 5:00 p.m.

**Mail/Messages:** Private phones are available in nearly all guest rooms. Messages can be left at 508-548-3705 24 hours a day and are available for pickup at the front desk of Swope Center (phones in rooms do not have voice mail). Packages and mail should be addressed to your name, *Society of General Physiologists, Marine Biological Laboratory, 7 MBL Street, Woods Hole, MA 02543-1015*.

*A map of the MBL Campus is featured on page iii of this program.*
70th Annual Meeting and Symposium
of the Society of General Physiologists
Marine Biological Laboratory,
Woods Hole, Massachusetts
September 7–11, 2016

Genetic & Animal Models for Ion Channel Function in Physiology and Disease
Organized by Andrea Meredith (University of Maryland School of Medicine) and Mark Nelson (University of Vermont)

**Schedule of Events**

**Wednesday, September 7**

2:00 – 9:00 Registration and Room Assignments (Swope Center, 1st Floor)

5:30 – 7:00 Dinner (Swope Center, Dining Room, 2nd Floor)

7:15 – 7:25 Opening Remarks (Lillie Auditorium, Lillie Building)
Conference Organizers, Andrea Meredith and Mark Nelson
Mark Shapiro, President, Society of General Physiologists

7:25 – 7:30 Introduction of Friends of Physiology Keynote Speaker
Andrea Meredith and Mark Nelson

7:30 – 8:30 Friends of Physiology Keynote Lecture
Jeff Noebels (Baylor College of Medicine)
‘Precision Physiology and Rescue of Ion Channel Disorders: The Gate is Now Wide Open’ [Abstract 62]

8:30 – 9:00 Rob Taft (The Jackson Laboratory)
‘CRISPR/Cas-Mediated Genome Engineering’ [Abstract 78]

9:00 – 11:00 Mixer (MBL Club)
Travel Awards Presented
THURSDAY, SEPTEMBER 8

7:00 – 8:30  Breakfast (Swope Center, Dining Room)

8:45– 9:55  Session I:  Channelopathies I (Lillie Auditorium, Lillie Building)
Session chair:  Al George

(8:45-9:20) Geoff Pitt (Weill Cornell Medicine)
‘The role of Ca_{V}1.2 L-type Ca^{2+} channels in physiology and disease’ [Abstract 69]

(9:20-9:55) Amy Lee (University of Iowa)
‘Pharmacological and genetic modifiers of a Ca_{V}2.1 channelopathy’ [Abstract 46]

9:55 – 10:15  Break

10:15 – 12:35  Session II:  New Transgenic Models
Session chair:  Lori Isom

(10:15 –10:50) Steven Marx (Columbia University)
‘Elucidating the mechanisms underlying the regulation of voltage-gated Ca^{2+} channels using transgenic mice’ [Abstract 42]

‘Using knockout approaches to tease apart physiological roles of Slo family ion channels and the contributions of regulatory subunits’ [Abstract 47]

(11:25-12:00) Murali Prakriya (Northwestern University)
‘Regulation of neurogenesis and astrocyte function by store-operated calcium channels’ [Abstract 72]

(12:00-12:25) Flash Poster Preview A (20mins)
Abstract 6:  Selective Activation of Muscarinic Receptors in Dentate Gyrus-CA3 Promotes Hyperexcitability and Seizure Susceptibility.  Chase Carver.
Abstract 16:  Trafficking of the cardiac sodium channel Na V 1.5 is regulated by the lateral membrane-specific protein CASK through its GUK and L27B domains.  Catherine Eichel.
Abstract 31:  Endothelial PIP2 bidirectionally modulates TRPV4 and Kir2.1 signaling in the brain capillary endothelium.  Osama Harraz.
Abstract 33: Discovery of a persistent inward current in skeletal muscle with characteristics suggesting it plays a central role in triggering myotonia in myotonia congenita. Ahmed Hawash.


Abstract 71: Role of Chloride Intracellular Channel (CLICs) Proteins in Maintaining Cardiac Mitochondrial Physiology. Devasena Ponnalagu.

12:35– 1:45 Lunch (Swope Center, Dining Room)

1:45 – 3:00 Roundtable with Journal of General Physiology and Journal of Clinical Investigation Editors (Meigs Room)

3:00 – 5:00 Poster Session A (Swope Center, 2nd Floor)
Session A Authors at Posters for discussion with attendees

5:00 – 6:30 Dinner (Swope Center, Dining Room)

6:30 – 9:00 Session III: Ion Channels in Disease Models
Session chair: Geoffrey Pitt

(6:30-7:05) Hugues Abriel (University of Bern)
‘Role of TRPM4 in human cardiac disorders’ [Abstract 1]

(7:05-7:40) Cathy Proenza (University of Colorado)
‘Electrical remodeling in the sinoatrial node associated with diabetes and aging’ [Abstract 74]

(7:40-8:15) William Catterall (University of Washington)
‘Dissecting Phenotypes by Gene Deletion in a Mouse Model of Dravet Syndrome’ [Abstract 7]

(8:15-8:50) Colin Nichols (Washington University)
‘Cantu Syndrome: Multi-organ complexities from KATP gain-of-function’ [Abstract 61]

9:00 – 11:00 Mixer (MBL Club)
FRIDAY, SEPTEMBER 9

7:00 – 8:30 Breakfast (Swope Center, Dining Room)

8:45 – 9:55 Session IV: Ion Channels in Disease Models II
(Lillie Auditorium, Lillie Building)
Session chair: Stephen Cannon

(8:45-9:20) Jorge Contreras (Rutgers University)
‘Cardiac dysfunction in Duchenne Muscular Dystrophy: Role of Connexin 43’ [Abstract 22]

(9:20–9:55) Erika Piedras-Renteria (Loyola University Chicago)
‘The KLHL1 KO as a model of altered T-type Ca channel function’ [Abstract 68]

(9:55-10:10) Short talk 1: Mike Bennett (Albert Einstein College of Medicine)
‘HIV-associated cardiovascular disease: Role of Connexin 43’ [Abstract 73]

10:10 – 10:30 Break

10:30 – 12:35 Session V: iPSCs and Patient-Derived Stem Cell Models
Session chair: Gail Robertson

(10:30-11:05) Jack Parent (University of Michigan)
‘Modeling epileptic encephalopathies with patient-derived neurons’ [Abstract 50]

(11:05-11:40) Rocky Kass (Columbia University)
‘Patient-specific induced pluripotent stems as a platform for precision therapeutics in the treatment of an inherited cardiac arrhythmia’ [Abstract 41]

(11:40-12:15) Peter Mohler (Ohio State University)
‘Defining new mechanisms underlying cardiovascular disease’ [Abstract 58]

(12:15-12:30) Short talk 2: Ivy Dick (University of Maryland)
‘Developing a New Approach for the Treatment of Timothy Syndrome’ [Abstract 13]

12:35 – 1:45 Lunch (Swope Center, Dining Room)

2:00 – 5:00 FREE AFTERNOON: Harbor Cruise Option 2:30-4:00
No charge. Sign-up sheet at registration table. Limited space.
FRIDAY, SEPTEMBER 9—Continued

5:00 – 6:30  Dinner (Swope Center, Dining Room)

6:30 – 9:00  Session VI: Channelopathies II
Session chair: Gary Yellen

(6:30–7:05) Lori Isom (University of Michigan)
‘Discovering Epilepsy Mechanisms in Dravet Syndrome’
[Abstract 39]

(7:05–7:40) Robert Brenner (University of Texas)
‘Gain-of-function mutations of BK potassium channels and pro-excitatory effects’ [Abstract 81]

(7:40–8:15) Stephen Cannon (UCLA)
‘Physiological foundations of susceptibility to periodic paralysis revealed by knock-in mouse models’ [Abstract 5]

(8:15–8:45) Flash Poster Preview B (30mins) (<2 mins, 1 slide):
Abstract 2: The Late L-type Ca Current as a Target for a New Class of Antiarrhythmics. Marina Angelini.
Abstract 29: Munc13 control of presynaptic voltage-gated Ca2+ channels. Geraldine Gouzer.
Abstract 38: Pyramidal neuron subpopulation excitability increases with reduced function and expression of voltage gated Na+ channels in the Scn1b null mouse model of Dravet syndrome. Jacob Hull.
Abstract 59: Potentiation of excitatory synaptic transmission in the superficial dorsal horn by low concentrations of Kv channel inhibitors. Tanziyah Muqeeem.
Abstract 70: Single nucleotide polymorphisms (SNPs) alter current from human BK channels. Amber Plante.
Abstract 76: Fast Voltage-Gated Sodium Channel Activity in Huntington’s Disease. Eric Reed.
Abstract 83: BK Channels Are Activated by Distinct Calcium Sources During Day and Night in SCN Neurons. Joshua Whitt.
Abstract 87: SWELL1 is a regulator of adipocyte insulin signaling and glucose homeostasis. Yanhui Zhang

9:00 – 11:00  Mixer (MBL Club)
SATURDAY, SEPTEMBER 10

7:00 – 8:30  Breakfast (Swope Center, Dining Room)

8:45 – 9:55  Session VII: Channelopathies III (Lillie Auditorium, Lillie Building)
Session chair: Murali Prakriya

(8:45 - 9:20) **Alfred George** (Northwestern University)
‘Decrypting Variants of Unknown Significance in Cardiac and Brain Channelopathies’ [Abstract 26]

(9:20 – 9:55) **Mark Nelson** (University of Vermont)
‘Potassium channelopathy-like defect underlies early-stage cerebrovascular dysfunction in a genetic model of small vessel disease’ [Abstract 60]

(9:55 - 10:10) **Short Talk 3: Scott Earley** (University of Nevada)
‘Redox Modulation of TRPA1 Channels in Brain Capillary Endothelial Cells Contributes to Neurovascular Coupling’

10:10 – 10:30  Break

10:30 – 12:35  Session VIII: Imaging Physiological Function & Optical Sensors
Session chair: Mark Nelson

(10:30-11:05) **Mark Rizzo** (University of Maryland)
‘FLARE biosensors for multiparametric imaging’ [Abstract 53]

(11:05-11:40) **Adam Cohen** (Harvard University, HHMI)
‘Bringing Bioelectricity to Light’ [Abstract 10]

(11:40-12:15) **Gary Yellen** (Harvard University)
‘Quantitative imaging of genetically-encoded metabolic sensors in mouse brain’ [Abstract 85]

(12:15 - 12:30) **Short Talk 4: Anna Greka** (Harvard University)
‘A small molecule inhibitor of TRPC5 suppresses proteinuric kidney disease progression’ [Abstract 103]

12:35 – 1:45  Lunch (Swope Center, Dining Room)

2:00 - 2:30  SGP Annual Meeting - All members welcome! (Meigs Room)

3:00 – 5:00  **Poster Session B** (Swope Center, 2nd Floor)
Session B Authors at Posters for discussion with attendees
SATURDAY, SEPTEMBER 10—Continued

5:00 – 6:30 Lobster Dinner/Clambake (Swope Center, Dining Room)

6:30 – 9:00 Session IX: Genetic Regulation
Session chair: Georgi Petkov

(6:30-7:05) Gail Robertson (University of Wisconsin)
‘Translational control of ion channel composition’ [Abstract 48]

(7:05-7:40) Andrea Meredith (University of Maryland)
‘Rhythmic regulation of BK splice variants in the circadian clock’ [Abstract 57]

(7:40- 8:55) Short Talk 5: Tracey Hermanstyne (Washington University)
‘Kv12-encoded channels selectively regulate nighttime firing rates in the suprachiasmatic nucleus’ [Abstract 34]

9:00 – 11:00 Mixer & Awards Ceremony (MBL Club)
Poster Awards Presented by Jeff Noebels

SUNDAY, SEPTEMBER 11

7:00 – 8:30 Breakfast (Swope Center, Dining Room)

10:00 Checkout deadline

12:35 – 1:45 Lunch (Swope Center, Dining Room)
Pick up box lunches after checkout
ABSTRACTS OF PAPERS AT THE SEVENTIETH ANNUAL MEETING OF THE SOCIETY OF GENERAL PHYSIOLOGISTS

Genetic and Animal Models for Ion Channel Function in Physiology and Disease

Marine Biological Laboratory
Woods Hole, Massachusetts
7–11 September 2016

Organized by
ANDREA MEREDITH and MARK NELSON
1. Role on TRPM4 in Human Cardiac Disorders. 
HUGUES ABRIEL, Department of Clinical Research, University of Bern, and NCCR TransCure, Switzerland

The TRPM4 channel is one of the 28 transient receptor potential channels expressed in the human body. It is expressed in a wide range of cells and tissues, but its function in physiology and pathophysiology is scarcely understood. Recently, its roles in human disorders such as cardiac electrical disturbances and multiple sclerosis have been demonstrated. In this talk, new data on genetic variants found in human TRPM4 linked to cardiac conduction defect, congenital atrioventricular block and Brugada syndrome will be presented. Our group is also currently developing more specific and potent TRPM4 small molecule inhibitors. Recent data on the characterization of these inhibitors used as either chemical chaperones or tools to understand the physiology of TRPM4 in vivo will be shown. Finally, we will present our new findings on the characterization of the cardiac electrical properties of a cardiac-specific TRPM4 knockout mouse line that we generated.

2. The Late L-type Ca Current as a Target for a New Class of Antiarrhythmics. M. ANGELINI,1 A. PEZHOUMAN,2 N. SAVALI,1 A. PANTAZIS,1 H.S. KARAGUEUZIAN,2,3 J.N. WEISS,2,3,4 and R. OLCESE,1,2,4 1Division of Molecular Medicine, Department of Anesthesiology, 2Cardiovascular Research Laboratory, 3Department of Medicine (Cardiology), David Geffen School of Medicine, and 4Department of Physiology, University of California, Los Angeles, Los, Angeles, CA 90095

There is an unmet need for an effective pharmacological therapy for cardiac arrhythmias to circumvent the limitations of class IV antiarrhythmics (e.g., diltiazem and verapamil) that block the peak L-type Ca current (I_{Ca,L}), compromising EC coupling. We recently discovered that early afterdepolarizations (EADs) of the cardiac action potential (AP), well-recognized triggers of fatal arrhythmias, can be potently suppressed by reduction of the late I_{Ca,L} component, without perturbing peak I_{Ca,L}. (Madhvani et al. 2015. J. Gen. Physiol. 145:395–400). Motivated by these findings, we are studying pilot compounds that could constitute a new, safer class of antiarrhythmics that do not compromise inotropy. One of the promising drugs is roscovitine, a purine analogue. We observed a selective reduction of late versus peak I_{Ca,L} by roscovitine in both native I_{Ca,L} from isolated rabbit ventricular myocytes under AP clamp and human Ca_{L,2} channels expressed in Xenopus oocytes. The EAD-suppressing ability of this drug was tested in isolated rabbit ventricular myocytes after induction of a stable EAD regimen with 600 μM H_2O_2. The oxidative stress caused EADs in 81% (95% CI: 66–95%) of the APs, prolonging AP duration (APD_90) from 276 ms (CI: 97–433 ms) to 840 ms (CI: 285–1,967 ms; n = 4). Addition of roscovitine (20 μM) completely abolished EADs and restored the APD_90 to 189 ms (CI: 155–222 ms). Moreover, roscovitine did not significantly perturb the Ca transient, implying the preservation of normal excitation–contraction coupling. We also tested roscovitine on isolated perfused aged rat hearts exposed to 0.1 mM H_2O_2 to induce EAD-mediated ventricular tachycardia/ fibrillation (VT/VF). Roscovitine addition suppressed these VT/VF in five out of six hearts within 10 min of perfusion. These findings set the basis for the development of a conceptually new class of antiarrhythmics (L-type Ca channel gating modifiers) that selectively reduce late I_{Ca,L}, with minimal or no effect on contractility.

3. HIV-tat Increases Connexin43 Expression and Alters Trafficking in Human Astrocytes: Role in NeuroAIDS. MICHAEL V. BENNETT1 and ELISEO A. EUGENIN,2,3 1Department of Neuroscience, Albert Einstein College of Medicine, Bronx, NY 10461; 2Public Health Research Institute (PHRI), Newark, NJ 07103; 3Departments of Microbiology, Biochemistry, and Molecular Genetics, Rutgers New Jersey Medical School, Rutgers the State University of NJ, Newark, NJ 07103

HIV-associated neurocognitive disorders (HAND) are a major complication in at least half of the infected population despite effective antiretroviral treatment and immune reconstitution. HIV-associated CNS damage is not correlated with viral replication, but instead is associated with mechanisms that regulate inflammation and neuronal compromise. Our data indicate that one of these mechanisms is mediated by gap junction channels. Normally, gap junction channels shut down under inflammatory conditions. However, HIV infection up-regulates Cx43 expression and maintains gap junctional communication. Here, we demonstrate that HIV-tat, the transactivator of the virus, and no other HIV proteins tested, increases Cx43 expression and maintains functional gap junctional communication in human astrocytes. Cx43 up-regulation is mediated by binding of the HIV-tat protein to the Cx43 promoter, resulting in increased Cx43 mRNA and protein, as well as increased gap junctional communication. Therefore, we propose that HIV-tat DNA persistent in the host genome is responsible for continued expression of the HIV-tat protein and that the resulting increased Cx43 expression allows spread of intracellular toxic signals generated in a few HIV-infected cells into surrounding uninfected cells. HIV-tat, which is membrane permeable, gets out of the infected cells and into adjacent cells where it increases Cx43 expression. Formation of gap junctions requires Cx43 in both apposed cells, and apoptosis is observed across astrocyte–astrocyte, astrocyte–neuron, and astrocyte–endothelial cell contacts. In the current antiretroviral era, where HIV replication is often completely suppressed, viral proteins such as HIV-tat are still produced and released from infected cells. Thus, blocking the effects of HIV-tat could be a new strategy to


SAMUEL J. BOSE,1 JIA LIU,1 ZHIWEI CAI,1 ALICE G.M. BOT,2 MARCEL J.C. BJVELDS,2 TIMEA PALMAI-PALLAG,3 MICHAEL J. MUTULO,1 ANN HARRIS,3 HUGO R. DE JONGE,2 DAVID N. SHEPPARD,1 1School of Physiology, Pharmacology and Neuroscience, University of Bristol, Bristol, England, UK; 2Department of Gastroenterology and Hepatology, Erasmus University Medical Center, Rotterdam, Netherlands; 3Human Molecular Genetics Program, Lurie Children’s Research Center, and Department of Pediatrics, Northwestern University Feinberg School of Medicine, Chicago, IL 60611

Cystic fibrosis (CF) is caused by mutations in the cystic fibrosis transmembrane conductance regulator (CFTR). Animal models of CF exhibit phenotypic variation, for example mice with the most common CF-causing mutation F508del do not develop the lung pathology observed in CF patients. Models with closer physiology to humans, including pig and ferret, have therefore been developed, and a sheep model has been proposed. To understand better the function of CFTR orthologues from model species, we used single-channel recordings to compare human, mouse, and sheep wild-type (WT) and F508del-CFTR and determine their response to the potentiator ivacaftor (VX-770; Vertex Pharmaceuticals).

The gating behavior of sheep CFTR closely resembled that of human CFTR, whereas mouse CFTR showed prolonged openings to a subconductance state (O2) with brief transitions to the fully open state (O3). Sheep CFTR single-channel current (i) and open probability (Po) were increased compared with those of human CFTR. However, the i and Po of mouse O2 were reduced compared with human CFTR. The F508del mutation reduced the Po of sheep CFTR less severely than its effect on human CFTR, whereas the mutation was without effect on mouse O2. For human and sheep CFTR, ivacaftor potentiated WT and F508del-CFTR; the drug was without effect on mouse WT and F508del-CFTR. Both human and sheep F508del-CFTR showed thermal instability at 37°C, whereas mouse F508del-CFTR remained stable at this temperature.

Our data indicate that CFTR from human, sheep, and mice show variation in function and pharmacology with relevance for animal models of CF. They also demonstrate that the F508del mutation does not have the same impact in CFTR orthologues from diverse species. These observations have implications for understanding the structure, function, and pharmacology of CFTR.

Supported by Cystic Fibrosis Trust; S.J. Bose is the recipient of a studentship from the UK Medical Research Council.

5. Physiological Foundations of Susceptibility to Periodic Paralysis Revealed by Knock-In Mouse Models.

STEPHEN C. CANNON, Department of Physiology, David Geffen School of Medicine, University of California, Los Angeles, Los Angeles, CA 90095

Hypokalemic periodic paralysis (HypoKPP) is a dominantly inherited disorder of skeletal muscle with recurrent episodes of severe weakness lasting hours to days, associated with low serum K+ (<3.5 mM). Weakness is caused by reduced fiber excitability resulting from sustained depolarization of the Vrest. Missense mutations, almost always at arginine residues of S4 voltage-sensor domains (VSDs), in Ca9.1 (60% of families) or Na9.1,4 (20%) are established causes of HypoKPP. Expression studies of HypoKPP mutant Na9.1,4 channels in oocytes revealed a common anomaly, the gating pore current conducted by a leaky VSD that is permissive to cation flux at hyperpolarized potentials and nonconducting when depolarized (V1/2 = −52 mV). Based on oocyte data, the gating pore conductance in HypoKPP muscle at a resting potential of ~90 mV is estimated to be 10 μS/cm² or 1% of the total resting conductance.

Muscle from knock-in mutant mice (Ca9.1,1-R528H or Na9.1,4-R669H) has a HypoKPP phenotype with severe loss of force in low K+ challenge, and voltage-clamp studies confirmed gating pore currents for mutant Ca9.1,1 as well as Na9.1,4 channels. Simulations of fiber excitability reveal a mechanism by which a small gating pore conductance may produce susceptibility to anomalous depolarization in low K+. When the K+ current that sets Vrest is conducted by Kir channels, then this outward current amplitude is compromised and Vrest depolarizes if external K+ is low enough (even though Eg is more negative). In normal muscle, this paradoxical depolarization occurs for K+ <2 mM. The HypoKPP gating pore current shifts this critical K+ to the 3–4 mM range, creating Vrest bistability of ~90 mV and ~60 mV. The pathogenic depolarized Vrest is favored by raised internal Cl−, and reducing Cl− influx via inhibition of the NKCC transporter with bumetanide prevents low-K−–induced loss of force in HypoKPP mice.

Supported by NIAMS of the NIH (AR42703, AR06182) and the MDA (381149).

6. Selective Activation of Muscarinic Receptors in Dentate Gyrus-CAL3 Promotes Hypercactivity and Seizure Susceptibility. CHASE M. CARVER and MARK S. SHAPIRO, Department of Physiology, University of Texas Health Science Center, San Antonio, TX 78229

Epileptogenesis describes the complex, plastic changes that alter a normal brain into a hyperexcitable network debilitated by recurrent seizures. The dentate gyrus (DG) and granule cell neurons control excitatory inputs into the hippocampus. Voltage-gated KCNQ2/3 K+ channel (M-channel) current maintains homeostatic control over neuronal resting membrane potential and
firing frequency. M-current is sensitive to suppression by muscarinic acetylcholine receptor activation. Cholinergic input to M1 muscarinic receptors and the dominant function of M-current in neuronal excitability suggest involvement in epileptogenesis pathways. We hypothesized that in DG granule cells, muscarinic receptor activity induces maladaptive changes in which M-current is more susceptible to muscarinic depression, conferring hyperexcitability and recurrent seizures.

We induced epileptogenesis by transgenic expression of muscarinic “Designer Receptors Exclusively Activated By Designer Drug” (DREADD) exclusive to the DG-CA3 circuit using Cre-loxP mice. Mice were administered the designer drug clozapine-N-oxide (CNO) to elicit remote activation of the muscarinic DREADDs. EEG electrodes were implanted in mice to record electrographic seizure activity. Stimulation of the muscarinic DREADD by CNO delivery was repeated once daily, and seizure behavior was scored. Active and passive discharge properties and M-channel currents of DG granule cells were characterized with patch-clamp electrophysiology in the brain slice.

Muscarinic stimulation of DG granule cells was sufficient to induce epileptogenic activity in vivo, as mice displayed focal seizure activity after administration of CNO. Repeated kindling of seizures resulted in generalized clonic seizures that persisted for 30–60 s. DG granule cells exhibited robust M-current suppression and greater hyperexcitability upon DREADD receptor stimulation.

Muscarinic receptor stimulation in the DG-CA3 circuit is sufficient to induce epileptiform excitation that progressed into network seizures. Chemogenetic spatial and temporal control represents a promising advancement in understanding epileptogenesis over traditional chemogenetic models. The interaction of muscarinic receptors and M-channels may strongly contribute to maladaptive, epileptogenic changes in the DG to promote seizure.

7. Dissecting Phenotypes by Gene Deletion in a Mouse Model of Dravet Syndrome. WILLIAM CATTERALL, Department of Pharmacology, University of Washington, Seattle, WA 98195

Dravet syndrome is a complex genetically dominant neuropsychiatric disease involving intractable epilepsy, profound cognitive deficit, autistic-like phenotypes, and a high frequency of sudden unexplained death in epilepsy (SUDEP). It is caused by loss-of-function mutations in the gene SCN1A encoding brain sodium channel Nav1.1, which specifically impair action potential firing by GABAergic inhibitory interneurons and disinhibit neural circuits in the brain. Deletion of Nav1.1 channels specifically in forebrain GABAergic interneurons using the Cre-Lox method recapitulates all of the manifestations of this disease, including epilepsy, SUDEP, cognitive deficit, and autistic-like behaviors. These results demonstrate that disinhibition of neural circuits by mutation of Nav1.1 channels causes both epilepsy and comorbidities in Dravet syndrome. Dissection of the cellular basis for Dravet syndrome further with the Cre-Lox method shows that mutation in parvalbumin-expressing fast-spiking interneurons causes pro-epileptic effects and autistic-like behaviors, deletion in somatostatin-expressing interneurons causes pro-epileptic effects and hyperactivity, and deletion in both types of interneurons gives synergistic effects on epilepsy and SUDEP and a partial phenotype for cognitive deficit. Sudden death is caused by excess parasympathetic outflow from the central nervous system during and following seizures, which causes intense bradycardia and frequently leads to sudden cardiac death by ventricular arrhythmia within 1–2 min. Further genetic dissection of pathophysiology and novel approaches to therapeutics for Dravet syndrome will be discussed.

8. The MAGUK CASK Regulates the Cardiac Sodium Channel Nav1.5, Which is Mediated by Calcineurin. MORGAN CHEVALIER, SARAH VERMIL, SABINE NAFLZGER, JEAN-SEBASTIEN ROUGIER, and HUGUES ABRIEL, Department of Clinical Research, University of Bern, Bern, Switzerland

Background: The voltage-gated Na\(^+\) channel Na\(_{1.5}\) is responsible for the rapid depolarization phase of the cardiac action potential (AP). Na\(_{1.5}\) is present in different membrane domains in cardiomyocytes and interacts with specific partners such as MAGUKs (membrane-associated guanylate kinase), which are key regulators of ion channels. The MAGUK family member CASK (calcium/calmodulin-dependent serine protein kinase) regulates ion channels in the brain. Here, we investigate the role of CASK in the regulation of Na\(_{1.5}\) in cardiomyocytes.

Methods and results: First, immunostainings in isolated mouse cardiomyocytes show that CASK is exclusively expressed at the lateral membrane. We also showed the SAP97-independent interaction of CASK with Nav1.5 by pulldown experiments with the Nav1.5 C-terminal domain in SAP97 KO and WT cardiomyocytes. To assess the functional consequences of the interaction between CASK and Na\(_{1.5}\) channels, patch-clamp experiments in cardiac-specific CASK KO cardiomyocytes revealed that \(I_{\text{Nav}}\) is increased by 40% without any significant modifications of the steady-state activation and inactivation. Additionally, AP recordings revealed that the AP threshold is significantly lower in CASK KO cardiomyocytes, while d\(V/dt\) and resting membrane potentials are not modified. Concomitantly, Western blots in CASK KO or WT cardiomyocytes revealed an increase in the expression of Na\(_{1.5}\). All together, these data show that CASK is a negative regulator of Na\(_{1.5}\) channels.
CASK also interacts with the calcium-dependent serine-threonine phosphatase calcineurin in cardiomyocytes, which inhibits its activity. To assess the role of calcineurin in the CASK-dependent regulation of Na⁺, I₅ᵥ was recorded in CASK-silenced TSA-201 cells. Interestingly, treatment with the calcineurin inhibitor cyclosporin A (10 μM) abolished the effect of CASK silencing on I₅ᵥ.

**Conclusion:** According to these results, CASK appears to be as negative regulator of Nav1.5, which activity is mediated by calcineurin.

9. Optogenetic Induction of Cortical Spreading Depressions. DAVID Y. CHUNG,¹,²,³ HOMA SADEGHIAN,¹ FUMIAKI OKA,¹,³ TAO QIN,¹ CENK AYATA,¹,² JGEMOSSOVA,¹,² Neurovascular Research Unit, Department of Radiology, Massachusetts General Hospital, Harvard Medical School, Boston, MA 02114; Stroke Service and Neuroscience Intensive Care Unit, Department of Neurology, Massachusetts General Hospital, Harvard Medical School, Boston, MA 02114; Department of Neurosurgery, Yamaguchi University School of Medicine, Ube, Japan; *presenting author

**Background:** Cortical spreading depression (CSD) has a critical role in the pathogenesis of secondary damage after traumatic brain injury, ischemic stroke, intracerebral hemorrhage, and aneurysmal subarachnoid hemorrhage. Experimental studies targeting CSDs continue to be important in the development of new therapeutic approaches; however, current CSD induction methods are cumbersome, cause primary brain injury, and obscure events at the nidus of the CSD.

**Methods:** We developed a minimally invasive method for the study of CSDs. Transgenic mouse lines expressing channelrhodopsin (ChR2) in cortical neurons were used to determine regional thresholds for optogenetic precipitation of CSDs through intact skull. In vivo laser speckle and Doppler flowmetry, bright field imaging, and electrocorticography were used to detect CSDs. Additionally, we measured local field potential shifts (LFP) and extracellular potassium using an ion-selective glass microelectrode inserted within the area of light stimulation.

**Results:** We were able to induce CSDs in multiple regions without causing primary brain injury. We observed regional differences in thresholds for optogenetically-induced CSDs (from lowest to highest threshold): (1) whisker barrel, (2) motor, (3) sensory, and (4) visual cortex. CSDs were reliably induced in whisker and motor cortices for all tested stimulation paradigms; however, it was not always possible to predict whether a CSD could be induced in sensory or visual cortex. Dynamic changes in LFP and increased extracellular potassium concentrations at the site of light stimulation preceded precipitation of a CSD.

**Conclusions:** Minimally invasive optogenetic stimulation through intact skull can reliably induce CSDs and enables observation of changes in extracellular potassium at the origin of a CSD. Optogenetic induction of CSDs in ChR2 transgenic mice is a potentially useful method for the study of secondary brain injury and can serve as an important tool for the future study of fundamental mechanisms of the phenomenon.

10. Bringing Bioelectricity to Light. ADAM COHEN, HHMI, Harvard University, Cambridge, MA 02138

We are developing tools for all-optical electrophysiology: simultaneous optical perturbation and optical measurement of membrane voltage in electrically active cells. We developed a protein-based fluorescent voltage indicator, QuasAr3, derived from a gene found in a Dead Sea microorganism. When expressed in neurons or cardiac myocytes, this gene reports firing via flashes of near-infrared fluorescence. We paired the reporter with a blue light–activated ion channel, called CheRiff. Using patterned optical stimulation, we have probed excitability, synaptic transmission, and network dynamics in primary neurons in culture, in human iPSC-derived neurons, in acute brain slice, and in vivo. We have also explored electrical activity in cardiomyocytes, vascular endothelial cells, and even bacteria. I will also describe applications to human iPSC-derived models of ALS and epilepsy and to modeling arrhythmia in human iPSC-derived cardiomyocytes.

11. Physiological Levels of Extracellular Histones Increase TRPV4 Activity While Trauma Levels Increase P2X-Mediated Ca²⁺ Influx. DANIEL COLLIÉ,¹ NURIA VILLALBA,¹ ADRIAN SACKHEIM,² KALEV FREEMAN,¹,² and MARK NELSON,¹,² Department of Pharmacology and ²Department of Surgery, University of Vermont, Burlington, VT 05405

Elevated levels of histone proteins are found in the circulation of patients following traumatic injury (healthy: 2.3 μg/ml, 4 h post trauma: 10–230 μg/ml; Abrams et al. 2013. *Am. J. Respir. Crit. Care. Med.* 187:160–169) and are associated with vascular dysfunction, coagulopathy, sepsis, and poor patient outcome. However, the mechanism and specific ion channels involved have not been identified. We used resistance-sized mouse mesenteric arteries (MAs) to study the effects of extracellular histones on the vasculature. To study the spatial and temporal characteristics of histone-induced endothelial cell (EC) Ca²⁺ signals, we used spinning disk confocal microscopy of en face MAs from EC-specific genetically encoded Ca²⁺ indicator mice (GECI, cx40-GCaMP5-mCherry). Physiological histone levels (1 μg/ml, unfractionated histone protein mixture of H1, H2a, H2b, H3, and H4) increased local Ca²⁺ events that were inhibited by 10 nM GSK219, a TRPV4-specific antagonist. Trauma levels of histones (10 μg/ml), within seconds of application, triggered large Ca²⁺ events that propagate within and between
ECs. 10 μg/ml histones increased intracellular Ca\(^{2+}\)
within minutes, and exposure for 30 min induced EC
death and decreased EC-dependent dilation in ex vivo
pressurized MA s. Propagating events were due to Ca\(^{2+}\)
influx rather than IP\(_{3}\)-R-mediated release from intra-
cellular stores, yet events were not sensitive to inhibi-
tion or genetic ablation of TRPV4. Propagating events
were suppressed by selective and nonselective purinu-
ergic antagonists (100 μM amiloride, 1 μM TNPATP, and
50 μM Suramin), suggesting a role for ionotropic puri-
ergic receptor (P2X)–mediated Ca\(^{2+}\) influx in ECs.
The data demonstrate that circulating histones are robust
activators of Ca\(^{2+}\) signaling in the vascular endo-
thelium from physiology to disease.

12. The Wistar Kyoto Rat Strain as a Model for Irrit-
able Bowel Syndrome. JULIE E. DALZIEL, WAYNE
YOUNG, KARL FRASER, SHALOME BASSETT, and
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The Wistar Kyoto (WKY) rat strain is hyper-responsive
to stress and considered a model for stress-associated
functional gastrointestinal (GI) disorders such as irrita-
ble bowel syndrome (IBS). Although GI transit and mi-
crobiota composition are altered in IBS, these aspects
have not been thoroughly investigated in WKY rats. We
therefore compared GI transit in WKY with Sprague
Dawley (SD) rats and characterized differences in cecal
microbiome composition and plasma metabolites asso-
ciated with stress. Transit of metallic beads was tracked
along different regions of the GI tract over 12 h by high
resolution x-ray imaging. Stomach emptying (4 h) and
transit to the small (9 h) and large intestine (12 h) were
assessed using a rating scale to classify bead location
Plasma extracts were analyzed with both positive and
negative ionization modes of lipid and HILIC (polar
compounds) liquid chromatometry mass spectrometry
(LC-MS) streams. Cecal microbial composition was de-
termined by Illumina MiSeq 16S rRNA amplicon se-
quencing and analysis using the QIIME pipeline. In
WKYs, 77% of beads were retained in the stomach com-
pared with 35% retention for SDs. Subsequent transit
was decreased by 36% at 9 h and by 21% at 12 h in WKY
compared with SDs. Interestingly, excluding those re-
tained in the stomach at 9 h, transiting beads had moved
39% further through the small intestine over 4–9 h for
WKY compared with SDs. Phylogenetic differences be-
tween rat strain cecal microbiota communities were
clearly distinguished and plasma metabolite differences
detected. The results demonstrate impaired stomach
emptying, yet rapid small intestine transit, in WKY rats
compared with SD animals. This was unexpected and
reveals that the slower GI transit in WKY can be largely
attributed to delayed stomach emptying. These observa-
tions suggest that WKY rats may also be considered an
appropriate model for functional gastric disorders.

13. Developing a New Approach for the Treatment of
Timothy Syndrome. IVY E. DICK, WORAWAN B.
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Timothy syndrome (TS) is a multisystem disorder,
including neurological impairment, autism, and car-
diac action potential prolongation (long QT) with life-
threatening arrhythmias. The underlying basis of TS is
typically a single point mutation in either exon 8a (TS
channel) or exon 8 (TS channel) such that the expression pattern of these mutually
exclusive exons significantly contributes to the overall
severity of the disease. The effect of these mutations on
channel function, blockers of these channels such as verapamil have only partially counter-
acted the severe symptoms of TS patients. Here, we
show that this is likely due to a differential effect of vera-
pamil on TS versus wild-type channels. In particular, the
use-dependent block of verapamil is significantly attenu-
ated in the context of the TS channel, thus decreasing
the efficacy of verapamil in TS patients. We therefore
propose an alternate therapeutic strategy in which in-
clusion of the TS-containing exon is decreased through
manipulation of the splice expression pattern via an ant-
sense oligonucleotide. Remarkably, treatment of in-
duced pluripotent stem cells (iPSCs) derived from a TS
channel patient resulted in not only a decrease of the del-
etious exon, but a corresponding increase in the non-
affected exon. Further, treatment of the cells normalized
the action potential duration, validating the functional
efficacy of the treatment strategy. Overall, the potential
benefits for TS patients are significant, and the appro-
ach may serve as a model for developing new ther-
peutic strategies for any channelopathy in which the
mutation occurs within a mutually exclusive exon.

14. Structural Elements of Peripheral Coupling Sites
Regulating Cerebral Artery Smooth Muscle Contractil-
ity. SCOTT EARLEY, Department of Pharmacology, Center
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Medicine, Reno, NV 89557

Junctional membrane complexes (JMCs) are static
structures that regulate close contacts (~20 nm) be-
tween the plasma membrane (PM) and sarcoplasmic
reticulum (SR) and are critically important for the ex-
citability and contractility of cardiac and skeletal muscle
cells. In smooth muscle cells (SMCs), JMCs occur as sites of peripheral coupling between the PM and the SR. These sites may be important for the activity of Ca\(^{2+}\)-sensitive ion channels on the PM by Ca\(^{2+}\) released from the SR, such as the activation of large-conductance Ca\(^{2+}\)-activated K\(^+\) channels (BK\(_{Ca}\)) by Ca\(^{2+}\) sparks generated by release of Ca\(^{2+}\) from the SR through type 2 ryanodine receptors (RyR2). Here we investigate the structural basis and functional significance of peripheral coupling in native contractile cerebral artery SMC. High-resolution live-cell confocal imaging experiments using PM-, SR-, and tubulin-specific fluorescent dyes revealed distinct arching microtubule structures beneath the peripheral SR proximal to the PM. These structures were also apparent when ground state depletion with individual molecule return (GSDIM) super-resolution nanoscopy was used to image single microtubule filaments. Surprisingly, GSDIM images also showed that immunolabeled RyR2 clusters selectively colocalized with microtubules in freshly isolated SMCs. To investigate the influence of microtubule arches on the formation of peripheral coupling sites at the molecular level, a combination of high-resolution live cell confocal microscopy and GSDIM super-resolution nanoscopy was used. SMCs labeled with PM- and SR-specific dye showed areas of peripheral coupling that were disrupted when nocodazole (10 μM) was used to depolymerize microtubules, but not when the actin cytoskeleton was disrupted with a combination of latrunculin B (1 μM) and swinholide A (0.1 μM). Peripheral coupling was also maintained during maximal agonist-induced contraction (uridine trisphosphate, 30 μM). Freshly isolated SMCs were immunolabeled with anti–BK\(_{Ca}\)-α subunit (BKα) and anti-RyR2 and imaged using GSDIM super-resolution nanoscopy. Using object-based analysis, we found that 1.9 ± 0.4% of BK\(_{Ca}\) puncta colocalized with RyR2, far greater than the colocalization of these two proteins in randomized control images (0.2 ± 0.1%, n = 12, p < 0.001). In addition, nocodazole treatment (10 μM, 1 h), significantly reduced colocalization of BKα and RyR2 (n = 12, P < 0.05), suggesting that intact microtubule networks are required for the formation of this signaling complex. Additional experiments were performed to establish functional consequences of disrupting peripheral coupling sites. Ca\(^{2+}\) sparks were recorded using high-speed, high-resolution confocal Ca\(^{2+}\) imaging in smooth muscle cells loaded the Ca\(^{2+}\) indicator Fluo-4AM. Cells treated with nocodazole showed no change in Ca\(^{2+}\) spark amplitude and rise time and a slight decrease in frequency. Ca\(^{2+}\) spark duration, decay time, and spatial spread were significantly increased. In perforated-patch clamp electrophysiology experiments (V\(_{H} = -30\) mV), nocodazole treatment abolished spontaneous Ca\(^{2+}\)-dependent BK\(_{Ca}\) activity, recorded as spontaneous transient outward currents (STOCs), suggesting that microtubule depolymerization uncouples Ca\(^{2+}\) spark-dependent activation of BK\(_{Ca}\) channels. Pressure myography was used to determine the effects of this response on arterial contractility. Myogenic vasoconstriction of cerebral resistance arteries (∼100–200 μm) treated with nocodazole was significantly greater than controls, likely due to loss of BK\(_{Ca}\) activity, which has a negative feedback on the membrane potential during pressure-induced vasoconstriction. Taken together, these findings demonstrate that microtubule-dependent formation of peripheral coupling sites is functionally significantly for the regulation of membrane excitability and contractility of arterial SMC.

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15. Recombinant ApoL1 Confers pH-Dependent Anion Permeability to Phospholipid Vesicles.

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**Background:** Variants in ApoL1 confer increased risk of certain chronic kidney diseases in people of African ancestry. ApoL1 has been reported to function as an ion channel, but reports vary on the nature of this activity. We sought to characterize ApoL1 ion transport activity more fully.

**Methods:** Recombinant N-terminal His-tagged ApoL1 was expressed in bacteria and purified using methods of Thompson and Finkelstein (2015. Proc. Natl. Acad. Sci. 112:2894–2899). Ion permeability was assessed using vesicle-based, voltage-dependent Cl and K efflux assays using ion-selective electrodes. Single channel properties were investigated using the tip-dip lipid bilayer approach. Protein structure was probed using intrinsic tryptophan fluorescence.

**Results:** The preparation yields large amounts of very highly purified His-tagged ApoL1, which is active in a trypanosome killing assay. Direct addition of ApoL1 to phospholipid vesicles yields robust Cl-selective permeability that supports voltage-driven Cl transport. The activity shows a strong dependence on pH, with a sharp drop in activity as pH is raised above 6.5. Activity is linearly dependent on mass of protein and shows strong dependence on lipid composition of the vesicles, requiring the presence of negatively charged phospholipids. We do not find cation-selective permeability when assayed at either pH 5.0 or 7.5. Biophysical studies of ApoL1 in solution reveal a significant structural transition in the same pH range in which the channel is activated (pH 7–6.5). In tip-dip bilayers, ApoL1 inserts at low pH, generating transitions with a range of conductances between 2 and 10 pS and with a nonrectifying current-voltage relationship. We do not find enhanced channel activity if the bath solution is changed to pH 7.5.

**Conclusions:** Purified recombinant ApoL1 can insert directly into phospholipid membranes at low pH and function as an anion-selective channel. Whether the disease-associated variants show altered channel properties remains to be determined.
16. Trafficking of the Cardiac Sodium Channel Na\textsubscript{v1.5} is Regulated by the Lateral Membrane-Specific Protein CASK Through Its GUK and L27B Domains. C. EICHEL, A. BEURIOT, F. LOUAULT, G. DILANIAN, A. COULOMBE, S. HATEM and E. BALSE, Paris 6 University, INSERM UMRS1166, Faculté de Medicine Pitie-Salpetriere, 75013 Paris

**Introduction:** Targeting and clustering of transmembrane proteins to specialized membrane domains of cardiac myocytes (CMs) are critical for proper cardiac function. MAGUK proteins have emerged as key partners in this organization. We have previously reported that the MAGUK protein CASK localizes at lateral membranes (LMs) of CMs, in association with the dystrophin–glycoprotein complex (DGC). We have also shown that CASK interacts with the main cardiac sodium channel, Na\textsubscript{v1.5}, and negatively regulates the sodium current (I\textsubscript{Na}).

**Methods:** Using adenoviral transfer technology to manipulate adult CMs, high resolution 3-D deconvolution and total internal reflection fluorescence (TIRF) microscopy, GST pull-down, biotinylation, RT-qPCR, and patch clamp electrophysiology, we investigated the mechanism of CASK-dependent regulation of I\textsubscript{Na} in CMs and examined the role of CASK in the organization of the Na\textsubscript{v1.5} macromolecular complex at LMs.

**Results:** Using a purified protein CASK, we showed that CASK directly binds to Na\textsubscript{v1.5} C terminus via the PDZ-binding motif SIV, regardless of syntrophin presence. Overexpression of WT CASK decreased I\textsubscript{Na} in CMs, whereas overexpression of CASK constructs deleted for either the L27B or GUK domain restored I\textsubscript{Na}. Neither the expression level of scn5a nor the total amount of Na\textsubscript{v1.5} proteins was modified upon CASK silencing or overexpression. Furthermore, the protein transport inhibitor brefeldin-A prevented I\textsubscript{Na} and Na\textsubscript{v1.5} surface expression increase in CASK-silenced CMs. Finally, quantification of Na\textsubscript{v1.5} expression at the LM and ID revealed that the LM pool was only increased upon CASK silencing.

**Conclusion:** CASK directly interacts with Na\textsubscript{v1.5} and controls its surface expression by regulating anterograde trafficking and/or stabilization at the LM. Both L27B and GUK domains of CASK are likely involved in this regulation. These results strengthen the concept of differentially regulated pools of Na\textsubscript{v1.5} and suggest that CASK could participate in maintaining low level of Na\textsubscript{v1.5} at the LM.

17. Excitation-Secretion Coupling in Zebrafish Pancreatic Islets. CHRISTOPHER EMFINGER,\textsuperscript{1,2,3} YIXI WANG,\textsuperscript{1} CHRIS REISSAUS,\textsuperscript{1} ALECIA WELSCHER,\textsuperscript{2} ZIHAN YAN,\textsuperscript{2} KRYS HYRC,\textsuperscript{3} MARIA S. REMEDI,\textsuperscript{1,2,3} and COLIN NICHOLS,\textsuperscript{1,3} \textsuperscript{1}Department of Medicine, Division of Endocrinology, Metabolism and Lipid Research, \textsuperscript{2}Department of Cell Biology and Physiology, \textsuperscript{3}Department of Medicine, and \textsuperscript{3}Center for the Investigation of Membrane Excitability Diseases (CIMED), Washington University in St. Louis, St. Louis, MO 63130

Zebrafish are becoming more frequently used to model development and metabolic diseases as well as to screen for genes modulating many pathways of interest. However, many questions regarding basic physiology in the zebrafish remain unanswered. In mammalian pancreata, ATP-sensitive potassium channels (K\textsubscript{ATP} channels) and voltage-dependent calcium channels (VDCCs) link metabolism and membrane excitability to insulin secretion and are critical in controlling blood glucose. Whether this process occurs in zebrafish and to what extent it is essential for overall glucose homeostasis are not well understood. While zebrafish possess orthologous genes for many mammalian proteins, whether key proteins are expressed or functional in the pancreas has not previously been determined. We characterized K\textsubscript{ATP} channels and VDCCs in islets and individual β cells isolated from zebrafish and show that the fish pancreatic cells express functional K\textsubscript{ATP} channels with similar subunit composition, pharmacology, and function as their mammalian counterparts. We also show that these channels are essential for metabolic control of insulin secretion ex vivo using isolated islets and in vivo via intraperitoneal glucose tolerance tests. Further, we demonstrate that fish β cells possess functional calcium channels that are active when islets are exposed to high glucose, sulfonylureas, or membrane depolarization, indicating that calcium fluxes in these cells are downstream of K\textsubscript{ATP}-dependent membrane depolarization. Strikingly, in contrast to mammals, zebrafish β cells are not electrically coupled, perhaps explaining differences in glucose tolerance between fish and mammals. Further, we have developed essential components for screening zebrafish for modulators of whole body responses to β-cell membrane inexcitability. Given that some but not all of the mammalian components for metabolism-secretion coupling are present in zebrafish, the fish provide an opportunity to understand the evolutionary conservation of many essential components of insulin secretion as well as to identify the pathways modulating whole-body responses to islet electrical inexcitability.

18. Underlying Mechanisms of Remission in a Mouse Model of Neonatal Diabetes. CHRISTOPHER EMFINGER,\textsuperscript{1,2} ALECIA WELSCHER,\textsuperscript{1} ZIHAN YAN,\textsuperscript{1} COLIN NICHOLS,\textsuperscript{1,2,3} and MARIA S. REMEDI,\textsuperscript{1,2,3} \textsuperscript{1}Department of Medicine, Division of Endocrinology, Metabolism and Lipid Research, \textsuperscript{2}Department of Cell Biology and Physiology, and \textsuperscript{3}Center for the Investigation of Membrane Excitability Diseases (CIMED), Washington University in St. Louis, St. Louis, MO 63130

Gain-of-function (GOF) mutations in the K\textsubscript{ATP} channel cause the channels to remain open despite rising glucose levels preventing insulin secretion, triggering neonatal diabetes mellitus (NDM). Individuals with identical mutations can have different levels of symptom severity and treatment efficacy, ranging from permanent...
to transient NDM, suggesting other factors modulate responses to β-cell inexcitability. Mice expressing tamoxifen-inducible K\textsubscript{ATP}-GOF mutations in pancreatic β-cells develop severe diabetes following transgene induction, which persists if untreated. However, if mice are treated with glibenclamide (GB) for two weeks (starting at disease induction) they show two distinctive outcomes at the end of treatment: (a) one group of mice enters into remission (TND), with normalization of glucose which persists long after treatment ended, and (b) another group develops severe and permanent diabetes (PND). Notably, remission in TND mice is not an off-target effect of GB, as some mice also remit with insulin treatment. To explore causes of these outcomes, we subjected K\textsubscript{ATP}-GOF mice to hyperinsulinemic-euglycemic clamps. These reveal that insulin sensitivity is greatly reduced in PND mice but improved in TND mice. Initial tracer studies show no differences between groups in glucose uptake rates to explain the divergent baseline blood glucose and clamp glucose infusion rates. To explore the temporal link between changes in insulin sensitivity and treatment response, insulin tolerance tests before disease induction and at the end of GB treatment were done. These reveal that blood glucose diverges before changes in insulin sensitivity. In addition, plasma lipids are elevated in PND mice and not in TND mice, but only after differences in glucose appear. Together, the results suggest that insulin sensitivity changes are a result of rather than a cause of diverging blood glucose in TND and PND and suggest that other signaling pathways may determine which individuals enter remission in NDM.

19. Ionic Bases of Conditioned Inhibition-Produced Changes in Hermisenda Type B Photoreceptors.
JOSEPH FARLEY, Neuroscience, Indiana University, Bloomington, IN 47405

While the neural mechanisms involved in simple forms of associative learning have been extensively studied, most research has focused upon paradigms in which stimuli are repeatedly paired. In contrast, little is known about the mechanisms underlying conditioned inhibition (CI), where an organism learns that one stimulus signals the absence of a second. In aversive conditioning paradigms, CI learning is often referred to as safety-signal learning. Our previous research has shown that CI in Hermisenda, established by repeated explicitly-unpaired (EU) presentations of light and rotation, separated by a fixed, lengthy temporal gap, results in increased phototactic behavior and decreased photoresponses and spike activity of ocular type B photoreceptors. Here I describe the ionic bases of these changes. Intracellular recordings confirmed our prior reports that B cells from EU-animals showed selective decreases in light-evoked generator potentials and spike frequencies. Voltage clamp analyses of type B cells revealed that two somatic K\textsuperscript{+} currents (I\textsubscript{A} and I\textsubscript{K\textsubscript{Ca}}) were increased for days following CI training. Elimination of I\textsubscript{K\textsubscript{Ca}} through chelation of intracellular Ca\textsuperscript{2+} abolished the CI-produced difference in light response, while 4-aminopyridine block of I\textsubscript{A} abolished CI-produced difference in spiking. Peristimulus time histogram analyses of spiking in B cells from untrained and random control animals revealed bimodal distributions, with the second longer mode being blocked by 4-AP and reflective of I\textsubscript{A}. In B cells from EU-trained animals, the second mode was more prominent than in control cells, indicative of greater I\textsubscript{A}. Hodgkin-Huxley style computational models constructed for B cells that incorporated the changes in I\textsubscript{A} and I\textsubscript{K\textsubscript{Ca}} produced by paired- or EU-conditioning produced changes in simulated light responses and spiking that mimicked those observed in physiological studies. These data are the first to correlate biophysical changes in ionic conductance systems with CI-produced alterations in excitability of identified neurons that are mediators of CI memory.

20. Mechanisms of Suppression of Kv1.1 Channel Activity by Aβ(1–42), JOSEPH FARLEY, KRISTI DEBOEUF, and MOHAMMAD FARIDUL ISLAM, Neuroscience, Indiana University, Bloomington, IN 47405

Many studies have found that Aβ-peptides participate in the pathogenesis of Alzheimer’s disease (AD), leading to disruption of calcium (Ca\textsuperscript{2+}) homeostasis and eventual cell death. The mechanisms underlying these effects remain unclear. Our work suggests that Aβ inhibition of voltage-dependent K\textsuperscript{+} channel activity (e.g., Kv1.1) is among the earliest steps. Our previous work, using murine Kv1.1 expressed in Xenopus oocytes, elucidated a pathway in which Ca\textsuperscript{2+}-dependent activation of protein phosphatase 2B (PP2B/calcineurin), protein kinase C (PKC), protein tyrosine kinases (PTKs), and RhoA all participate to produce rapid and strong suppression of Kv1.1 activity. This pathway is recruited by a variety of stimuli that increase [Ca\textsuperscript{2+}], including GPCRs that couple to G\textsubscript{q/11}-PLC, calcium ionophore (A23187), and LGICs that flux Ca\textsuperscript{2+}. Because Kv1.1 and related channels are activated during an action potential and regulate Ca\textsuperscript{2+} influx during depolarization and inhibition of Kv1 channels is often neurotoxic, we speculate that Aβ suppression of Kv1 channels could lead to hyperexcitability, altered synaptic transmission, disruption of Ca\textsuperscript{2+} homeostasis, and neurotoxicity. We assessed the effects of Aβ(1–42) peptide (AnaSpec, monomers and low-n oligomers) on Kv1.1 channels in oocytes. Aβ(1–42) (10 nM to 1 μM) produced a dose-dependent inhibition of macroscopic Kv1.1 current: ~50% reductions within 30 m for 1 μM. Reverse sequence (40–1) peptide and other controls failed to suppress Kv1.1. Aβ suppression of Kv1.1 was partially Ca\textsuperscript{2+} and PP2B dependent, being reduced by ~50% when cells were either loaded with BAPTA-AM or exposed to the PP2B-inhibitor, cyclosporine A. The Aβ-stimulated
Ca$^{2+}$ increase did not occur via Ca$^{2+}$ influx. Complete removal of extracellular Ca$^{2+}$ failed to attenuate Aβ suppression of Kv1.1. Instead, Aβ appears to release Ca$^{2+}$ from internal stores. Single channel recording results suggest the possibility that Aβ suppression of Kv1.1 also involves direct protein–protein interaction of Aβ with Kv1.1 channel subunits.

21. Looking for the Line Between Agonists and Inhibitors. Prokaryotic Sodium Channels and Allosteric Modulation of Voltage-Sensitive Gating. ROCIO K. FINOL-URDANETA,1,2 JEFF R. MCARTHUR,2 RACHIELLE GAUDET,2 DENIS TIKHONOV,3 BORIS ZHOROV,3,4 and ROBERT J. FRENCH,1 1Department of Physiology and Pharmacology, University of Calgary, Calgary, Alberta, Canada T2N 4N1; 2Harvard University, Department of Molecular and Cellular Biology, Cambridge, MA 02138; 3Sechenov Institute of Evolutionary Physiology and Biochemistry, Russian Academy of Sciences, St. Petersburg 194223, Russia; 4Department of Biochemistry and Biomedical Sciences, McMaster University, Hamilton, Ontario, Canada L8S 4L8

Both agonistic biological toxins, such as batrachotoxin (BTX), and peptide inhibitors, such as µ-conotoxins (µCTXs), show use-dependent, apparently allosteric actions on voltage-gated sodium (Nav) channels. At this time, available high-resolution structural data, and the relatively simple, homo-tetrameric structures, of prokaryotic Nav channels offer an excellent opportunity to better understand these complex ligand–channel interactions. We have recently found that both BTX (Finol-Urdaneta et al. 2016. Biophys. J. 110:109a.) and µCTXs show potent interactions with NavBac channels. For BTX concentrations in the range of 1–10 µM, NaChBac channel modulation was facilitated, under voltage clamp, by trains of conditioning depolarizing pulses. Tail currents, on repolarization, increased in amplitude and slowed throughout the train, coincident with a negative shift in activation voltage. µCTX PIIIA inhibited NaChBac in the picomolar range (IC50 ~0.5 pM; Finol-Urdaneta et al. 2015. Biophys. J. 108:583a.), even more potent than striking earlier predictions (Chen and Chung. 2012. Biophys. J. 102:483–488.). Inhibition/block approached a quasi-steady-state, while being monitored with repeated depolarizing test pulses to −10 mV (Vh = −120 mV). Furthermore, at concentrations well below the IC50, single-pulse inactivation decay rates increased. Thus, actions of both of these structurally disparate toxins were facilitated by voltage protocols that normally drive transitions of the channels from resting to activated states. Given that the binding sites for BTX and the µCTXs appear to be on opposite sides of the selectivity filter, these data seem to be consistent with considerable flexibility and coupling along the pore lining.

We explored BTX interactions with a NaChBac model, based on a cryo-EM structure of NavCt (Tsai et al. 2013. J. Mol. Biol. 425:4074–4088.). Modeling and mutagenesis studies suggest that the BTX-binding site is separate from the µCTX receptor but overlaps residues homologous to those important for both local anesthetic and BTX binding in eukaryotic channels.

22. Cardiac Dysfunction in Duchenne Muscular Dystrophy: Role of Connexin 43. DIEGO FRAIDENRACH,1 PATRICIO GONZALEZ,1 NATALIA SHIROKOVA,2 JAYALAKSHMI RAMACHANDRAN,2 and JORGE E. CONTRERAS,2 1Department of Cell Biology and Molecular Medicine and 2Department of Pharmacology, Physiology, and Neuroscience, Rutgers University, New Jersey Medical School, Newark, NJ 07103

Duchenne muscular dystrophy (DMD) is characterized by the lack of dystrophin in cardiac and skeletal muscle cells. This leads to progressive loss of functional myocardium, to heart failure, and to damage of skeletal muscle. In the working myocardium, gap junctions located at intercalated disks mediate proper intercellular propagation of electrical signals and contractility. The gap junction channels are composed of connexin protein. Recently, we found that connexin 43 (Cx43), the dominant connexin in the ventricular myocardium, is up-regulated and mislocalized from the intercalated disks to lateral regions in cardiomyocytes of DMD mice and humans. These lateralized Cx43 proteins seem to be in the form of active unopposed plasma membrane channels ("hemichannels") rather than junctional channels. Strikingly, we found that specific pharmacological inhibition of Cx43 hemichannels prevents isoproterenol-induced arrhythmias and death in DMD mice. Furthermore, we found that the lateralized Cx43 hemichannels in DMD cardiomyocytes cause increased membrane permeability, which may cause damage to cardiac myocytes via enhanced free radical production and cytosolic Ca$^{2+}$ increase. Together, these results suggest that the DMD-induced mislocalization of Cx43 has pathological consequences regarding both cardiac function and myocyte health/survival. This is a novel and potent mechanism by which DMD causes cardiac pathology. Because Cx43 lateralization occurs in many cardiac pathologies (e.g., hypertension, ischemic reperfusion, myocardial infarction, and hypertrophy), we propose that opening of these hemichannels can have a key role in many cardiovascular diseases.

23. Genomic Insights from Comparative Analyses of Teleost Fish: Implications for Myopathies Related to Excitation-Contraction Coupling. JENS FRANCK, Department of Biology, University of Winnipeg, Winnipeg, MB, Canada R3B 2E9

Excitation-contraction (EC) coupling describes the relationship between the depolarization of the muscle membrane and the subsequent contraction of the muscle cell. In skeletal muscle cells, the depolarization of the muscle membrane triggers a conformational change
in the L-type calcium channel (Cav1.1). In skeletal muscle, the Cav1.1 directly interacts and mechanically gates the intracellular ryanodine receptor (RyR) channel to release calcium from the intracellular stores of the sarcoplasmic reticulum, a mechanism termed depolarization-induced calcium release (DICR). In contrast, the L-type channel in cardiac muscle (Cav1.2) opens in response to the depolarization signal, and the extracellular calcium subsequently acts as a ligand to gate open the intracellular RyR channels, a mechanism termed calcium-induced calcium release (CICR). The DICR mode of calcium release is believed to be a vertebrate innovation. We previously described fiber type-specific expression of RyR1 paralogues in slow twitch (yr1α) and fast twitch (yr1β) muscle fibers in fish (Franck et al. 1998. Amer. J. Physiol. 275: C401–C415; Darbandi and Franck. 2009. Comp. Biochem. Physiol. B. 154:443–448). More recently, it has been reported that the α1S subunits (α1S) of the multimeric Cav1.1 channel are also duplicated and expressed discretely in the slow twitch (α1Sα) and fast twitch (α1Sβ) muscles of zebrafish (Schredelseker et al. 2010. Proc. Natl. Acad. Sci. 107:5658–5663). Interestingly, the Cav1.1 channels in teleosts do not conduct any extracellular calcium and rely solely on the DICR mode of intracellular calcium release (Schredelseker et al. 2010. Proc. Natl. Acad. Sci. 107:5658–5663). Diseases related to abnormalities in the EC coupling process include malignant hyperthermia (MH), hypokalemic periodic paralysis (HypoKPP), and central core disease (CCD). Mutations in both the Cav1.1 channels and α1S genes have been linked to MH. In the α1S gene, five mutations, R174W, R1086H, R1086C, R1086S, and T1354S have been identified as causative (Bannister and Beam. 2013. Biochim. Biophys.Acta. 1828:1587–1597). The T1354S mutation is proposed to act by increasing the rate of inward calcium current through the Cav1.1 channel and an increase in caffeine sensitivity of the intracellular RyR1 channel (Pirone et al. Am. J. Physiol. Cell Physiol. 299:C1345–C1354). In light of the fact that teleost fish do not rely on extracellular calcium, it is interesting to observe that the α1S genes in teleosts all encode S at position 1354. This observation therefore provides potential insight into the relationship between Cav1.1 and RyR1 coupling in disease states such as MH.

25. Regulation of Cerebral Artery Endothelial TRPV4 Channel Function by cGMP-Dependent Protein Kinase. KALEV FREEMAN, ADRIAN BONEV, DANIEL COLLIER, NURIA VILLALBA, OSAMA HARRAZ, SWAPNIL SONKUSARE, and MARK NELSON. 1Department Surgery, University of Vermont, Burlington, VT 05405; 2University of Virginia, Charlottesville, VA 22908

Cerebral arteries (CAs) have uniquely different vaso-dilatory mechanisms than mesenteric arteries (MAs); yet, both arteries have functional endothelial TRPV4 channels. In MAs, TRPV4 Ca<sup>2+</sup> events in myoendothelial projections (MEPs) control vasodilation. We measured vasodilation in response to the TRPV4 agonist GSK1016790A (GSK101) in similarly sized mouse CA.
and MAs. The concentration of GSK101 necessary to induce CA vasodilation was 10× higher than for MAs (EC50, 322 nM vs. 32 nM). We next measured elementary TRPV4 Ca²⁺ influx events (sparklets) in CAs from mice expressing a genetically encoded endothelial Ca²⁺ biosensor (GCaMP2). The amplitude and quantal levels of TRPV4 sparklets were the same in CAs as MAs. Because the coupling of elementary Ca²⁺ influx events depends on A-kinase anchoring protein (AKAP150), we analyzed the relationship of sparklets to AKAP and MEPs in CAs. CAs had the same TRPV4 coupling strength between sparklet events and similar proximity to AKAP150 at MEPs, as MAs. However, the baseline TRPV4 open channel probability in CAs was substantially lower. In conventional whole cell patch clamp of CA endothelial cells, introduction in CAs was substantially lower. In conventional whole cell patch clamp of CA endothelial cells, introduction of nonhydrolyzable ATP increased TRPV4 currents. In perforated patch clamp experiments, inhibition of cGMP-dependent protein kinase (PKG) with Rp-8-Br-PET-cGMP caused a 227 (±15) % increase in TRPV4 currents in response to 300 nM GSK101; yet, this did not completely replicate the 10× difference between tissue beds. Ca²⁺ imaging experiments in CAs in the presence of Rp-8-Br-PET-cGMP or NO inhibitor L-NNA showed that both interventions increase Ca²⁺ sparklets in CAs. These data suggest phosphorylation of the TRPV4 channel by PKG, accounts for approximately twofold regulation in TRPV4 function in CAs. However, our data also suggest that other endogenous factors, possibly other kinases, must also be involved in strong negative regulation of TRPV4 channels in the brain, possibly to protect CA endothelial cells from excessive Ca²⁺ influx.

26. Decrypting Variants of Unknown Significance in Cardiac and Brain Channelopathies. ALFRED GEORGE, Northwestern University Feinberg School of Medicine, Chicago, IL 60611

Genetic testing has become standard-of-care for many diseases including channelopathies affecting cardiac rhythm such as the congenital long-QT syndrome (LQTS) and those associated with severe childhood epilepsy. However, interpreting genetic test results is often confounded by the discovery of “variants of unknown significance” for which there is insufficient data to establish whether or not a particular variant predisposes to a disease. This problem is particularly vexing for genetic disorders with strong allelic heterogeneity and a preponderance of “private” mutations. For channelopathies, in vitro functional assessments have been valuable for determining the potential pathogenicity of variants discovered in the research setting, but the value of this approach for variant classification in the clinical setting has not been evaluated. Functional annotation experiments (e.g., patch-clamp recording) have become the gold standard in assessing the likely pathogenicity of ion channel variants, but the extreme time and labor intensity of this approach are insufficient to tackle the thousands of known variants.

We have implemented a high-throughput (dual 384-well), automated patch-clamp system to enable functional annotation and pharmacological profiling of human ion channel variants at an unprecedented scale. Data from studies of cardiac potassium channels in LQTS as well as brain sodium and potassium channels associated with early onset epileptic encephalopathy will be presented to illustrate the power of this approach.

27. Pericyte-Mediated Alterations of Blood Flow Distribution at Capillary Bifurcations in a Genetic Model of Cerebral Ischemic Small Vessel Disease. ALBERT L. GONZALES, THOMAS A. LONGDEN, FABRICE DABERTRAND, BO SHUI, MICHAEL I. KOTLIKOFF, and MARK T. NELSON. 1Department of Pharmacology, University of Vermont, Burlington VT 05404; 2Department of Biomedical Sciences, College of Veterinary Medicine, Cornell University, Ithaca, NY 14853

Cerebral autosomal-dominant arteriopathy with subcortical infarcts and leukoencephalopathy (CADASIL) is the most common heritable cause of stroke and vascular dementia and is caused by dominant mutations of NOTCH3, which is expressed only in smooth muscle cells and pericytes in the brain. Pericytes cover a significant portion of the vast capillary network within the brain. However, only pericytes closest to the terminal arterioles are contractile. The goal of the current study is to elucidate the functional role of contractile pericytes at capillary bifurcations and to examine the consequence of the NOTCH3 mutation on pericycle-controlled capillary blood flow. We tested the hypothesis that functional pericytes asymmetrically control the luminal diameter of branches at capillary bifurcations independently of one another and, in doing so, are capable of controlling the distribution of red blood cells (RBCs) within the capillary system. Using a novel transgenic mouse expressing a genetically encoded Ca²⁺ indicator in contractile pericytes (acta2-GCaMP5-mCherry), we characterize the ion channels contributing to local Ca²⁺ signals regulating the asymmetry constriction by the pericycle processes of capillary branches. Using in vivo multiphoton microscopy, we observed distribution of RBCs at capillary junctions in correlation with the angle of bifurcation and diameter of daughter branches. Using a transgenic mouse model of CADASIL, we detected a disruption in RBC flow at capillary bifurcations, suggesting that pericytes play an integral role in controlling the directional flow of blood within the capillary bed.

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28. BK Channel α and γ1 Subunits Assemble with a Stoichiometry of Up To 4:4, But One γ1 Is Sufficient To Produce the Full γ1-Induced Gating Shift. VIVIAN GONZALEZ-PEREZ,1 MANU BEN-JOHNY,2 and CHRISTOPHER J. LINGLE,1 1Department of Anesthesiology, Washington University School of Medicine, St Louis, MO, 63110; 2Department of Biomedical Engineering, The Johns Hopkins University School of Medicine, Baltimore, MD, 21205

Ca2+ and voltage-gated K+ channels of large conductance (BK) are symmetrical complexes whose minimal functional unit is a tetramer composed of four identical α-subunits. BK α-subunits can coassemble with auxiliary subunits belonging to two different families, β and γ, which drastically influence channel gating. The presence of β2 subunit produces a complete inactivation of BK currents, and the inactivation rate at single channel level is a functional reporter of the number of β2 subunits/channel. This methodology has established that up to four β-subunits can assemble with the α-subunits in a BK channel complex, where each one contributes an identical additive increment to the total β-induced gating shift. However, the stoichiometry of the assembly of α and γ subunits is still unknown. To solve it, we designed an “inactivating” γ1 subunit (chimera β2/γ1) and measured the inactivation rates at the single channel level. Our results show that, similar to β subunits, up to four β2/γ1 subunits can assemble in BK channels expressed in Xenopus oocytes. Unlike β subunits, the presence of one β2/γ1 subunit in a single BK channel is sufficient to produce the full γ1-induced gating shift. Our results confirm the “all-or-none” type of functional regulation produced by γ1 on BK channels.

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29. Munc13 Control of Presynaptic Voltage-Gated Ca2+ Channels. GERALDINE GOUZER, MINGYU XUE, and TIMOTHY RYAN, Department of Biochemistry, Weill Cornell Medicine, New York, NY 10065

Synaptic transmission relies on precise control of presynaptic voltage-gated Ca2+ channels to drive exocytosis of neurotransmitters. Given the steep relationship between Ca2+ influx and release probability, it is critical to understand the machinery responsible for tuning Ca2+ influx at nerve terminals and how this is coordinated with tuning of the release machinery itself. We previously showed that the critical active zone SNARE assembly molecule Munc13-1, which is essential for all known forms of fast neurotransmitter release, also controls the gating of voltage-gated Ca2+ channels in nerve terminals (Calloway et al. 2015, eLife. 4:e07728). Munc13-1 contains a SNARE-binding Mun domain and an adjacent tandem CI-C2B domain that mediates both plasma membrane and Ca2+ channel interactions. The CI domain of Munc13 was previously shown to be critical for mediating the strong modulatory impact of the diacylglycerol (DAG) mimetic phorbol-ester on synaptic transmission, presumably by tuning SNARE assembly such as to lower the energy barrier for fusion. We show here that activation of the CI domain in Munc13 likely disrupts the C2 domain interaction with Ca2+ channels since application of phorbol ester rapidly decreases action potential–driven presynaptic Ca2+ influx by ~25%. Genetic ablation of Munc13 reduces Ca2+ influx by a similar amount, and the remaining AP-driven Ca2+ influx is no longer sensitive to phorbol ester. Furthermore phorbol-ester modulation of AP-driven presynaptic Ca2+ influx is eliminated when Munc13-1 is replaced with a variant harboring a point mutation in the CI domain that renders it insensitive to DAG. Our data are thus consistent with a model by which interactions of the CI domain with the plasma membrane changes the positioning of the C2B domain with respect to the Ca2+ channel, breaking the interaction, while in turn positioning the Mun domain to drive more favorable SNARE assembly and potentiation of neurotransmitter release at the expense of lowered Ca2+ influx.

30. Physiological and Behavioral Comorbidities in a Zebrafish Model for Dravet Syndrome. BRIAN P. GRONE and SCOTT C. BARABAN, Department of Neurological Surgery, University of California, San Francisco, San Francisco, CA 94143

Mutations in SCN1A, which encodes Nav1.1, cause Dravet syndrome (DS), a catastrophic childhood epilepsy. In addition to early-life seizures, DS patients experience comorbid conditions such as movement disorders, sleep disturbances, anxiety, early fatality, and cognitive decline. To study the functional consequences of SCN1A mutations, we use zebrafish with a loss-of-function mutation in scn1lab, one of two zebrafish homologues of SCN1A. Homozygous scn1lab+/+ mutant larvae exhibit early-life seizures, metabolic deficits, and early death. To establish zebrafish as a model for quantifying comorbid conditions in DS, we performed a battery of in vivo assays with scn1lab+/+ mutants between 3 and 6 d postfertilization (dpf). To study motor activity during a seizure, we used high-speed video imaging with simultaneous EEG recording in head-fixed zebrafish. In mutants (n = 8), long-duration ictal events were associated with high-velocity, complex sinusoidal tail deflections >50 degrees and lasting 600–1200 ms; interictal events were associated with briefer tail movement and smaller angle deflections. To study anxiety, we used locomotion tracking to monitor exploratory behavior in an open field arena. Mutants exhibited impaired exploratory behavior, with significantly increased time spent freezing and more time spent in the periphery. To evaluate nighttime arousal disturbances, we tracked larvae in 96-well plate format for 24 h. Locomotor activity during night (sleep phase) was higher in mutants compared with controls. To assess cardiac function, we measured
heart rate using video recordings. No significant differences in heart rate compared with controls were observed. We also quantified the distribution of onset of fatality, which occurs in the first 14 dpf. Our results demonstrate conserved features of movement disorders, anxiety, sleep disturbances, and early fatality in *sonlab* mutant zebrafish. These studies establish the zebrafish as a model for studying comorbid disease states, in addition to seizures, in a zebrafish DS model.

31. Endothelial PIP2 Bidirectionally Modulates TRPV4 and Kir2.1 Signaling in the Brain Capillary Endothelium. OSAMA F. HARRAZ and MARK T. NELSON,1,2 1Department of Pharmacology, College of Medicine, University of Vermont, Burlington, VT 05405; 2Institute of Cardiovascular Sciences, University of Manchester, Manchester, England, UK

Brain capillaries form a sensory web that can detect neuronal activity. We recently showed that brain capillary endothelial cells (cECs) express ideal candidates for the transduction of Ca2+ and electrical signals, namely TRPV4 and Kir2.1 channels, respectively. Here, we tested the hypothesis that the same regulatory pathway can simultaneously exert opposite effects on TRPV4 and Kir2.1 channels in freshly isolated cECs from C57BL/6 mouse brain. TRPV4 currents were inhibited by the inclusion of hydrolysable ATP in the patch pipette. Scavenging phosphatidylinositol 4,5-bisphosphate (PIP2) or inhibition of its synthesis reversed ATP effects. In contrast, Kir2.1 currents were unaffected by exogenous PIP2 or ATP. Kir2.1 current, however, declined over time, and this rundown did not occur when PIP2 was included in the pipette. Activation of G protein–coupled receptors (GqPCRs) hydrolyzes PIP2 to produce IP3 and diacylglycerol (DAG). We tested the hypothesis that GqPCR activation through hydrolysis of PIP2 has divergent consequences on TRPV4 and Kir2.1 currents. Prostaglandin E2 (PGE2; an agonist of the GqPCR EP1), which has been implicated in neurovascular coupling, relieved the inhibition of TRPV4 channels by intracellular ATP. The latter effect was attributed to PIP2 hydrolysis rather than enhanced IP3/Ca2+ or DAG/PKC signaling. On the other hand, Kir2.1 current rundown was significantly augmented by PGE2. In conclusion, our studies support the concept that PIP2 tonically inhibits TRPV4 channels and activates Kir2.1 channels in brain capillary endothelial cells. GqPCR activation acts as a molecular switch to alter the balance between electrical (Kir2.1) and Ca2+ (TRPV4) signaling, which would change the signaling modality to upstream penetrating arterioles and likely would have profound effects on the control of blood flow into the brain.

Through collaboration with the Regeneron Genetics Center, we have performed whole exome sequencing on more than 50,000 individuals for whom we have comprehensive clinical data from the electronic health records including ECG records. We have discovered a large number of variants in ion channels that we are assessing for association with various disease states. As a first example, we identified a patient who carried a novel truncation mutation in KCNJ3 gene (Kir3.1 or GIRK1 channel). The patient had a strong personal and family history of heart disease and has since died of a suspected arrhythmia. Kir3.1 and Kir3.4 (GIRK4) form a hetero-tetramer, traffic to the cell surface, and comprise IK(A), in atria. The stop-gain mutation, R286stop identified in this patient, removes the vast majority of the channel C terminus. To test the functional relevance of this novel mutation to the patient’s arrhythmia, we Flag tagged Kir3.1 on the N terminus and mutated Arginine 286 to a stop codon (R286stop). In HEK293 cells, R286stop degraded more rapidly compared with wild-type Kir3.1. That degradation was prevented by proteasome inhibitors. Kir3.4 and WT Kir3.1 interact with R286stop but also appear to degrade when coexpressed with R286stop, suggesting a dominant negative effect on the interacting proteins caused by the truncated channel. Kir3.1 alone resides in the ER and is targeted to the plasma membrane when coexpressed with Kir3.4. R286stop, on the other hand, localizes to the ER and vesicle type puncta in the cell and does not traffic to the plasma membrane when Kir3.4 is coexpressed. Interestingly, R286stop traps some of the Kir3.4 inside the cell, reducing its cell surface expression. Our data suggest the R286stop mutation could have resulted in diminished IK(A), leading to the arrhythmia in the patient.

32. Whole Exome Sequencing in 50,000 Clinic Patients Leads to Discovery of a Novel Kir3 Mutation Implicated in Arrhythmia. CASSANDRA M. HARTLE, BRYN S. MOORE, and TOORAJ MIRSHAHI, Weiss Center for Research, Geisinger Clinic, Danville, PA 17832

33. Discovery of a Persistent Inward Current in Skeletal Muscle with Characteristics Suggesting It Plays a Central Role in Triggering Myotonia in Myotonia Congenita. AHMED HAWASH, ANDREW VOSS, and MARK RICH, Wright State University, Dayton, OH 45435

Myotonia congenita is caused by loss-of-function mutations of the skeletal muscle chloride channel (CIC-1) and is characterized by involuntary repetitive firing of action potentials. Patients with myotonia congenita experience muscle dysfunction, mainly due to muscle stiffness. For over 40 yr, it has been accepted that a buildup of K+ in the transverse tubular system during trains of evoked action potentials is responsible for the sustained firing of spontaneous action potentials. If K+ buildup is the cause, opening K+ channels should worsen myotonia. However, our preliminary studies show that opening K+ channels using retigabine lessened myotonic firing. This strongly suggests that K+ buildup cannot be the sole cause of myotonia. We thus hypothesized that an additional depolarizing event, which remains active for seconds (the length of a myotonic run),
drives the repetitive action potentials in myotonia. A possible mechanism involves persistent inward currents (PICs), which are resistant to fast inactivation and enable repetitive firing in motor neurons. To look for a PIC in skeletal muscle, we used slow ramp protocols under voltage clamp in dissociated flexor digitorum brevis fibers of CIC-1-null (myotonic) mice. We identified a PIC in myotonic skeletal muscle with characteristics needed to drive the repetitive involuntary firing seen in myotonia congenita. We also determined that a similar PIC is present in wild-type muscle, suggesting the current has a hitherto overlooked physiological role. We suggest that a physiological role of skeletal muscle PIC may be to sustain repetitive firing during prolonged activation of muscle.

34. Kv12-Encoded Channels Selectively Regulate Nighttime Firing Rates in the Suprachiasmatic Nucleus. TRACEY O. HERMANSTYNE,1 DANIEL GRANADOS-FUENTES,2 ERIK D. HERZOG,2 and JEANNE M. NERBONNE,1 1Departments of Medicine and Developmental Biology, Washington University School of Medicine and 2Department of Biology, Washington University, St. Louis, MO 63130

Kv12.1 and Kv12.2 K⁺ channels, which generate outwardly rectifying currents in the subthreshold voltage range, are strong candidates for regulating daily rhythms in suprachiasmatic nucleus (SCN) membrane excitability. We explored the hypothesis that Kv12.1 and/or Kv12.2 contribute to the daily variations in the resting and active properties of neurons in the SCN that are critical for circadian rhythms in physiology and behavior. Short hairpin RNAs (shRNAs) selectively targeting Kv12.1 or Kv12.2 were used to acutely “knockdown” Kv12.1 or Kv12.2 expression in the adult mouse SCN. Current-clamp recordings in acute SCN slices revealed that the mean input resistances of SCN neurons at night were significantly (P < 0.05) higher in Kv12.1 (1.2 ± 0.3 GΩ) and Kv12.2 (1.8 ± 0.3 GΩ)-targeted shRNA-expressing SCN neurons, compared with WT SCN neurons (0.9 ± 0.1 GΩ). In addition, mean ± SEM nighttime action potential thresholds were significantly (P < 0.05) more hyperpolarized in Kv12.1-targeted shRNA-expressing (−32.6 ± 2.1 mV) and Kv12.2-targeted shRNA-expressing (−30.1 ± 2.2 mV) than in WT (−24.1 ± 1.8 mV) SCN neurons. Furthermore, compared with WT SCN neurons (−46.2 ± 1.4 mV), the mean ± SEM nighttime resting membrane potential was significantly (P < 0.05) more depolarized in Kv12.1-targeted shRNA-expressing SCN neurons (−37.3 ± 3.3 mV). Mean ± SEM repetitive firing rates measured at night were also significantly (P < 0.01) higher in Kv12.1 (4.9 ± 1.1 Hz) and Kv12.2 (3.6 ± 0.8 Hz)-targeted shRNA-expressing SCN neurons when compared with WT SCN neurons (0.7 ± 0.2 Hz). Parallel daytime experiments revealed no significant differences in passive membrane properties and repetitive firing rates in WT and Kv12.1- and Kv12.2-targeted shRNA-expressing SCN neurons. Taken together, these results reveal that the targeted “knockdown” of Kv12.1 or Kv12.2 selectively regulates nighttime firing rates in the SCN with very little effect on daytime firing activity. These observations suggest a critical role for Kv12.1- and Kv12.2-encoded K⁺ channels in regulating the “down-state” nighttime electrical activity in the SCN.

35. Impaired Calcium Release in Skeletal Muscle Fibers with Cav1.1 Channels Carrying the Hypokalemic Periodic Paralysis-Causing Mutation R528H. ERICK O. HERNÁNDEZ-OCHOA,1 FEN FEN WU,2 STEPHEN C. CANNON,2 and MARTIN F. SCHNEIDER,1 1Department of Biochemistry and Molecular Biology, University of Maryland School of Medicine, Baltimore, MD 21201; 2Department of Physiology, David Geffen School of Medicine, University of California, Los Angeles, Los Angeles, CA 90095

Hypokalemic periodic paralysis (HypoPP) is a genetic disease that presents with severe and recurrent episodes of weakness triggered by reduced serum potassium. Mutations at arginine residues in the S4 segments of the voltage sensor domains (VSD) in the voltage-gated calcium channel (Cav1.1) are associated with HypoPP. We developed a knock-in Cav1.1 mutant with one arginine-to-histidine substitution (R528H) located at the S4 of the second VSD to investigate abnormal muscle excitability in HypoPP. R528H mice recapitulated the disease in a gene dosage–dependent manner; muscle force was reduced and became exacerbated after low K⁺ challenge, muscle fibers displayed abnormal excitability and aberrant “gating pore” currents–ionic fluxes conducted via a crevice of the channel through which the S4 moves—that increase the susceptibility to paradoxical depolarization in low extracellular [K⁺] distinctive of HypoPP. Interestingly, we also reported that depolarization-induced Ca²⁺ transients were impaired in fibers from homozygous R528H mice. Here, we further investigate the impact of the R528H mutation on voltage-dependent Ca²⁺ release using Ca²⁺ imaging, membrane dyes, voltage-clamp, and confocal microscopy techniques. Depolarization-induced rhod-2 Ca²⁺ transients were significantly decreased in muscle fibers from homozygous R528H mice when compared with heterozygous or wild-type counterparts, as were the Ca²⁺ transients and Ca²⁺ release calculated from F/F₀ signals. The altered Ca²⁺ release was not due to differences in internal releasable Ca²⁺ or alterations in fiber and T-tubule morphology, suggesting Ca²⁺ release defects and further dysfunction of the Cav1.1. The impact of R528H mutation on Ca²⁺ release suggests the hypothesis that S4 from second VSD contributes to Cav1.1/RyR1 electromechanical coupling and Ca²⁺ release in skeletal muscle. Further mutations in the VSDs of Cav1.1 are needed to explore the mechanistic underpinnings of Cav1.1 voltage sensing and excitation-contraction coupling.

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36. Monoaminergic Synaptic Transmission at the Merkel Cell–Neurite Complex. BENJAMIN U. HOFFMAN, YOSHIHISA BABA, and ELLEN A. LUMPKIN, Department of Dermatology, Columbia University, New York, NY 10032

Of the five cardinal senses, touch is the least understood. Due to the complexity of the peripheral nervous system, mechanisms of how we encode touch stimuli have remained elusive. In the skin, gentle touch is mediated in part by Merkel cells—epidermal cells that cluster in highly touch-sensitive skin regions, including fingertips, whisker follicles, and touch domes. In complex with myelinated (Aβ) somatosensory afferents, they form high acuity receptors that mediate slowly adapting type I (SAI) responses. The Merkel cell–neurite complex detects pressure, represents shapes and edges, and guides dexterous hand movements. We hypothesized that Merkel cells release monoamine-containing vesicles at SNARE-dependent synapses to excite firing in tactile afferents. To test this hypothesis, we selectively expressed tetanus toxin light chain (TeNT) in epidermal cells in vivo in order to disrupt SNARE-mediated synaptic vesicle release. Using ex vivo skin-nerve recordings, we showed that Merkel cells require functional SNARE proteins to mediate canonical SAI responses. Indeed, in response to mid-range stimuli, TeNT-expressing Merkel cells were unable to mediate sustained firing in tactile afferents. Moreover, to show that Merkel cells package and release monoamines as neurotransmitters, we employed a fluorescent neurotransmitter analogue to label Merkel cells in semi-intact epidermal preparations. Merkel cells loaded neurotransmitter analogues into acidic vesicles in a VMAT2-dependent manner. Finally, neurotransmitters were released from Merkel cells in response to depolarizing stimulation. For the first time, these experiments reveal that Merkel cells employ SNARE-dependent, regulated release of neurotransmitters to signal to tactile afferents and set the stage to study the receptor pathways through which this information is transduced.

37. A Novel Pain Assay Using a Transgenic Mousexpressing hNa1.7 with an Inherited Erythromelalgia Mutation: Comparison with Traditional Models for Inflammatory and Neuropathic Pain. S. HOWARD,1 G. BANKAR,2 K. NELKENBRECHER,3 M. WALDBROOK,3 Z. XIE,1 K. KHAKH,1 E. CHANG,1 C. YOUNG,1 S. LIN,1 J.P. JOHNSON,1 L.E. SOJO,1 A. LINDGREM,1 N. CHAHAL,1 S. CHOWDHURY,1 S. DECKER,1 I. HEMEON,1 C.M. DEHNHARDT,1 J. CHANG,2 B. SAFINA,2 D.P. SUTHERLIN,2 D. HACKOS,2 C.L. ROBINETTE,1 and C.J. COHEN,1 Xenon Pharmaceuticals, Burnaby, BC, Canada; 2Genentech, South San Francisco, CA 94080

Loss-of-function mutations in Na1.7 result in congenital indifference to pain with little or no effect on motor or cognitive function. Moreover, gain-of-function mutations result in inherited erythromelalgia (IEM) and other painful conditions. A humanized transgenic mouse was developed for in vivo testing of target engagement of hNa1.7. BAC transgenic FVB mice incorporated hNa1.7 with the IEM mutation I848T. Transgenic mice express high levels of hNa1.7 mRNA in DRGs and olfactory bulbs, areas of high expression of the endogenous channel. A subcutaneous hind paw injection of the sodium channel agonist aconitine elicits a reproducible and quantifiable flinching/licking response for up to 60 min. The aconitine effects are dependent on the presence of the transgene since a comparable injection of aconitine into wild-type mice induces minimal nociceptive behaviors. The model provides a well-defined PK-PD relationship and facilitates the rapid evaluation of compounds. The analgesic activity of sodium channel blockers was characterized both in the transgenic IEM assay and in traditional pain assays using littermate wild-type mice. We have used two acylsulfonamide s that are potent inhibitors of [3H]GX-545 binding to Na1.7. This ligand targets a binding site on VSD4 that has a high degree of molecular selectivity among Na,8 (Ahuja et al. 2015. Science, 350:4ac5464). One compound has limited selectivity against Na1.1, Na1.2, and NaV1.6, while the second is >10-fold selective against these isoforms. However, with allowance for modest differences in IC50 for mouse versus human Na1.7, both compounds are equipotent in the IEM target engagement assay and in assays using littermate wild-type mice. The assays include inflammatory pain induced by complete Freund’s adjuvant or formalin or neuropathic pain caused by treatment with streptozocin. These comparative studies with compounds that differ in selectivity among Na,8 indicate that block of Na1.7 alone is sufficient for broad analgesic activity against inflammatory and neuropathic pain.

38. Pyramidal Neuron Subpopulation Excitability Increases with Reduced Function and Expression of Voltage-Gated Na+ Channels in the Scn1b-Null Mouse Model of Dravet Syndrome. JACOB HULL, LARISA KRUGER, HEATHER O’MALLEY, and LORI ISOM, Department of Neuroscience, University of Michigan, Ann Arbor, MI 48105

Mutations in SCN1B, encoding voltage-gated Na+ channel (VGSC) β1 subunits, result in pediatric epilepsies including Dravet syndrome (DS), an epileptic encephalopathy. While the majority of DS patients have de novo mutations in SCN1A, patients with two loss-of-function SCN1B alleles also have DS. Brain VGSCs contain one pore forming α and two modulatory β subunits. The effects of Scn1b on excitability are cell type and brain region specific, and thus studying Scn1b-null mice may help identify dysfunctional DS microcircuits. We investigated the effects of Scn1b deletion at the level of individual pyramidal neuron populations in p14-21–null
and WT mice. We found no changes in excitability of pyramidal neurons in acute slices of visual cortex layer 2/3 (n = 19, n = 9) versus WT (n = 16, n = 10) or layer 5 (n = 16, n = 8) versus WT (n = 12, n = 7). However, null layer 6 (L6) (n = 22, n = 19) versus WT (n = 19, n = 11) and subicular pyramidal neurons (n = 9, n = 4) versus WT (n = 13, n = 6) had increased firing in 1s current injections and increased input resistance (Rm). To investigate the role of somatic Na+ current density (I\text{Na}), we pulled nucleated patches from L6-null (n = 9, n = 8) and WT (n = 8, n = 8) and subicular-null (n = 4, n = 4) and WT (n = 8, n = 8) pyramidal neurons, finding 43% and 39% decreases in I\text{Na} density, respectively (P < 0.05). Action potential (AP) phase plots in 1s trains showed no change during the first AP and attenuation of both the axon initial segment and somatic components in later APs in null L6 neurons versus WT (P < 0.05). H-CTX binding to cortical membranes showed a 35% reduction in VGSCs in null (n = 5) versus WT (n = 3) (P < 0.05). Reduced I\text{Na} suggests K+ channels regulating Rm may underlie hyperexcitability in Scn1b-null neurons. Future work will test this hypothesis. Identification of affected neuron populations and the contributing ion channels may guide development of treatments for DS.

39. Discovering Epilepsy Mechanisms in Dravet Syndrome. LORI L. ISOM, University of Michigan Medical School, Ann Arbor, MI 48105

Neuronal channelopathies cause various brain disorders including epilepsy, migraine, and ataxia. Despite the development of mouse models, pathophysiological mechanisms for these disorders are poorly understood. One particularly devastating channelopathy is Dravet syndrome (DS), a severe childhood epileptic encephalopathy (EE) with a high risk of sudden unexplained death in epilepsy (SUDEP). DS is typically caused by de novo dominant mutations in SCN1A, encoding the voltage-gated Na+ channel Na1.1. Although SUDEP is the most devastating consequence of epilepsy and the leading cause of epilepsy mortality, astonishingly little is understood about its causes, and no biomarkers exist to identify at-risk epilepsy patients. Heterologous expression of mutant Na1.1 channels suggests haploinsufficiency, raising the question of how loss of Na+ channels underlying action potentials produces hyperexcitability. Data from DS mouse models indicate both decreased Na+ current (I\text{Na}) in interneurons, implicating disinhibition, and increased I\text{Na} in pyramidal cells, implicating hyperexcitability, depending on genetic background, brain area, and animal age. To understand the effects of SCN1A DS mutations in human neurons, we derived forebrain-like neurons from two DS subjects by induced pluripotent stem cell (iPSC) reprogramming of patient fibroblasts and compared them with iPSC-derived neurons from human controls. We found that DS patient-derived neurons have increased I\text{Na} density and spontaneous bursting in both bipolar- and pyramidal-shaped neurons. Because Na1.1 is also expressed in heart, a compelling idea is that altered I\text{Na} in DS cardiac myocytes, in addition to central neurons, may lead to arrhythmias and contribute to SUDEP. To test this hypothesis, we used the iPSC method to derive cardiac myocytes from fibroblasts of DS subjects. Our data suggest that a subset of DS subjects shows increased cardiac myocyte I\text{Na} and excitability, which may be predictive of SUDEP risk. In parallel studies of a DS human mutant SCN1A knock-in mouse model, we observed spontaneous seizures and SUDEP in the mice, increased ventricular cardiac myocyte I\text{Na} density, and ventricular arrhythmias at the time of SUDEP. To discover potential common mechanisms of SUDEP, we studied changes in cardiac excitability in two additional EE models, a SCN8A knock-in model of EIEE13 and a Scn1b-null model of DS. Both models showed cardiac arrhythmias with changes in I\text{Na} and calcium handling in ventricular myocytes. Parallel studies of human patient-derived SCN8A- and Scn1b-EE-linked iPSC neurons and cardiac myocytes are in progress. The ultimate goal of our work is to provide a greater understanding of the mechanisms of DS and related EEs that may lead to novel therapeutic agents for epilepsy and SUDEP prevention.

40. Regional Differences in Transient Outward Potassium (I\text{o}) Currents and the Role of I\text{o} in Human Ventricular Repolarization. ERIC K. JOHNSON,1 STEVEN SPRINGER,1 WEI WANG,1 ANNA ZHANG,1 EVELYN M. KANTER,1,2 KATHRYN A. YAMADA,1 and JEANNE M. NERBONNE,1,3 1Department of Medicine, Division of Cardiology, 2Department of Surgery, and 3Department of Developmental Biology, Washington University in St. Louis, St. Louis, MO 63130

The rapidly activating and inactivating Ca2+-independent transient outward potassium current, I\text{o}, has been described in the hearts of many mammals, where it contributes to the early phase of action potential repolarization and to the generation of normal cardiac rhythms. Dysregulation of I\text{o} has been implicated in Brugada syndrome and in hypertrophied and failing hearts and is associated with increased risk of potentially life-threatening arrhythmias. The biophysical properties and functional role(s) of I\text{o} in the human heart, however, are poorly understood. To fill this knowledge gap, voltage-clamp studies were performed in myocytes isolated from the subepicardial (LV Sub-Epi), epicardial (LV Epi), endocardial (LV Endo), and sub-endocardial (LV Sub-Endo) regions of the left ventricle. A gradient of I\text{o} densities was identified with higher densities in LV Sub-Epi and LV Epi than in LV Sub-Endo and LV Endo myocytes. The voltage dependencies and kinetics of I\text{o} activation and inactivation were similar in all LV cell types. Analysis of the kinetics of I\text{o} recovery from inactivation revealed two distinct
I_\text{to}, component: in LV Sub-Epi myocytes, the mean ± SEM time constant of recovery was 19.3 ± 0.3 ms (n = 21), reflecting expression of I_{\text{to,f}}, whereas the time constant of recovery in LV Sub-Endo myocytes was 1,067 ± 45 ms (n = 37), reflecting expression of I_{\text{to,s}}. In LV Epi (n = 42) and LV Endo (n = 39) myocytes, recovery was bi-exponential, consistent with the presence of both I_{\text{to,f}} and I_{\text{to,s}}.

To explore the role(s) of I_{\text{to}} in shaping human ventricular action potential (AP) waveforms, a mathematical model of I_{\text{to}} was generated and used in dynamic-clamp experiments, and the effects of changing I_{\text{to}} density and inactivation kinetics on AP properties were determined. Manipulation of I_{\text{to}} density had pronounced effects on notch potentials with minimal effects on other AP properties. Manipulation of I_{\text{to}} inactivation kinetics had dramatic effects on both the shapes and durations of action potentials.

41. Patient-Specific Induced Pluripotent Stems as a Platform for Precision Therapeutics in the Treatment of an Inherited Cardiac Arrhythmia. ROBERT KASS, Columbia University, New York, NY 10027

Understanding the genetic basis for differential responses to drug therapies is a key goal of precision medicine. Induced pluripotent stem cells (iPSCs) offer a unique system to investigate the pharmacology of disease processes in therapeutically and genetically relevant primary cell types in vitro. In this presentation, we report the use of patient-derived iPSCs to understand limitations of clinical regimen that had failed to control cardiac arrhythmias in a long QT syndrome patient who was found to have a de novo SCN5A LQT-3 mutation and a polymorphism in KCNH2, the gene for LQT2. Analysis of the molecular pharmacology of ion channels expressed in cardiomyocytes differentiated from these iPSCs (iPCS-CMs) revealed drug interactions that included off target inhibition of the KCNH2 (hERG) channel that could be minimized by altering heart rate. The improved in vitro pharmacologic approach correlated with improved management of arrhythmias in the patient and provides support for this approach in developing precise patient-specific clinical regimens.

42. Elucidating the Mechanisms Underlying the Regulation of Voltage-Gated Ca^{2+} Channels Using Transgenic Mice. ALEXANDER KATCHMAN,1 LIN YANG,1 and STEVEN MARX,1,2 1Division of Cardiology, Department of Medicine and 2Department of Pharmacology, College of Physicians and Surgeons, Columbia University, New York, NY 10027

Cav1.2, the sarcolemmal L-type Ca^{2+} channel, plays a key role in regulation of signaling in the brain and the heart. In the heart and brain, abnormalities in Cav1.2 function are associated with diseases, such as autism, heart failure, and hypertrophy. A detailed molecular understanding of Cav1.2 regulation in myocytes and brain has been hampered, however, by the inability to recapitulate and then dissect in heterologous expression systems key aspects of Cav1.2 function. We have developed novel tools to surmount major obstacles that have limited progress in the field and allow us to probe molecular aspects of Cav1.2 regulation, using biochemical and electrophysiological techniques, within the context of cardiomyocytes and neurons, but with the power of a heterologous expression system. Using a transgenic (TG) approach that enables selective and reliable expression of FLAG-epitope–tagged, dihydropyridine-resistant Cav1.2 channel subunits, harboring mutations at key regulatory sites or covalently linked to regulatory components, in adult cardiomyocytes and at all stages of development, we have explored the mechanisms responsible for adrenergic regulation of Cav1.2. We found that Ser1700 and Thr1704 are not required for β-adrenergic stimulation of Cav1.2 in adult cardiomyocytes. Naturally, in a protein as large as α1C, there are numerous intracellular consensus PKA phosphorylation sites, although many may be not surface exposed. Rather than continue the search for single phosphorylated residues (using in vitro techniques which has not been fruitful), we created a transgenic mouse with Ala substitutions of 17 and 22 potential PKA sites and found that Cav1.2 channels harboring these 17 and 22 Ala substitutions were functional, trafficked to the dyad, and were appropriately regulated by adrenergic stimulation, implying that these residues are not required for adrenergic regulation. We have concluded that direct phosphorylation of the α1C subunit by PKA may not be required for adrenergic regulation of Cav1.2. We have also sought to determine whether proteolytic cleavage of α1C is required for the adrenergic stimulation of Cav1.2. Since proteolytic cleavage cannot be reconstituted in heterologous expression, there is no effective way to study the process, other than in native tissues. Although deletion of the 1798NNAN motif did not alter the proteolytic cleavage of α1C, cleavage likely occurs in this general region, however, based upon the molecular weight of the truncated α1C. Within the region, there are other similar motifs including 1794NANI1797. Although the protease responsible for cleavage of α1C is not known, it has been speculated to be calpain-like, A conserved PEST sequence, a motif rich in Pro, Glu, Ser, and Thr that has been proposed to serve as substrate recognition sites for calpains, is just N-terminal to Ala1800. We have created a transgenic mouse line with deletion of both the PEST sequence (1769DTESP) and 1794NANI1794NANN1802. Data will be presented concerning these mice. In summary, using novel transgenic approaches, we have clarified key mechanisms responsible for adrenergic regulation of Ca^{2+} channels.
43. TRPV1-Mediated Ca\(^{2+}\) Influx in Arterial Smooth Muscle. MASAYO KOIDE, ELIZABETH HUGHES, ARSALAN U. SYED, SWAPNIL SONKUSARE, INESSA MANUELYAN, MARK T. NELSON, and GEORGE C. WELLMAN, Department of Pharmacology, University of Vermont College of Medicine, Burlington, VT 05405

Ca\(^{2+}\) influx through L-type voltage-dependent Ca\(^{2+}\) channels (VDCCs) has traditionally been viewed as the major Ca\(^{2+}\) entry pathway in arterial smooth muscle (ASM) to control contraction as well as modulate excitability and transcription. Here, we provide direct evidence for another major Ca\(^{2+}\) entry pathway—the TRPV1 channel—in ASM of specific vascular beds. Using mice with a genetically-encoded Ca\(^{2+}\) indicator (GCaMP5-tdTomato) linked to endogenous TRPV1 expression, we have found functional expression of TRPV1 channels in ASM throughout the external carotid artery (ECA) territory. Using “optical” patch clamping techniques (Sonkusare et al. 2012, Science. 336:597–601), we provide the first measurements of Ca\(^{2+}\) influx through single TRPV1 channels (“TRPV1 sparklets”) and show that this unitary input of Ca\(^{2+}\) dwarfs (fivefold) the flux through a single VDCC. In TRPV1-positive arteries (e.g., facial artery), physiological stimuli such as adrenergic agonists and endocannabinoids cause robust increases in ASM Ca\(^{2+}\) and vasoconstriction through TRPV1 activation. Our data indicate that when ASM TRPV1 are activated and become the dominant contributor to membrane conductance and Ca\(^{2+}\) entry, VDCC-dependent regulation of arterial diameter is lost. We propose that TRPV1-mediated vasoconstriction causes increased resistance in the ECA territory, leading to a redistribution of blood flow from ECA to the internal carotid and cerebral arteries, which do not possess functional TRPV1 channels. Further, under conditions of ASM TRPV1 activation, endogenous hyperpolarizing vasodilator mechanisms would remain intact in cerebral arteries but would be short-circuited in the external carotid territory, amplifying increases in cerebral blood flow. In summary, our data suggest that Ca\(^{2+}\) influx mediated via ASM TRPV1 channels can profoundly impact the regulation of arterial diameter and blood flow.

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44. Enhanced Astrocyte Ca\(^{2+}\) Signaling Contributes to Pathological Neurovascular Coupling After Subarachnoid Hemorrhage. MASAYO KOIDE, ANTHONY C. PAPPAS, and GEORGE C. WELLMAN, Department of Pharmacology, University of Vermont College of Medicine, Burlington, VT 05405

Neurovascular coupling (NVC) is a unique function within the brain to increase local blood flow in regions of active neurons. We have previously demonstrated an inversion of NVC, or neuronal activation-induced vasoconstriction rather than vasodilation, in brain slices from subarachnoid hemorrhage (SAH) model rats (Koide et al. 2012, Proc. Natl. Acad. Sci. 109:E1387–E1395). This pathological neuronally-evoked vasoconstriction coincided with the emergence of astrocyte endfeet high-amplitude Ca\(^{2+}\) signals (eHACS). In the present study, we examined the mechanism underlying SAH-induced eHACS. Using two-photon fluorescent and infrared-differential interference contrast microscopy, astrocyte endfeet Ca\(^{2+}\) and arteriolar diameter were recorded in brain slices from SAH model animals. We observed both high amplitude (eHACS, peak Ca\(^{2+}\) >500 nM) and control-like astrocyte Ca\(^{2+}\) signals in the absence of electrical or chemical stimulation that were abolished by cyclopiazonic acid (a sarco/endoplasmic reticulum Ca\(^{2+}\) ATPase inhibitor) or U73122 (a phospholipase C inhibitor). These data indicate that these spontaneous endfoot Ca\(^{2+}\) signals are IP\(_3\)-mediated intracellular Ca\(^{2+}\) release events. Interestingly, either the broad-spectrum purinergic receptor inhibitor, suramin, or a cocktail of P2Y receptor inhibitors blocked eHACS, but not control-like Ca\(^{2+}\) events. Further, suramin treatment restored vasodilatory NVC in brain slices from SAH model animals. These results indicate that P2Y purinergic signaling contributes to SAH-induced eHACS and inversion of NVC. Moreover, these data demonstrate that the amplitude of spontaneous astrocyte Ca\(^{2+}\) signaling can profoundly impact NVC. Abnormal astrocyte Ca\(^{2+}\) oscillations are observed in various disease states such as Alzheimer diseases, epilepsy, and small vessel diseases. Thus, investigating and targeting molecular mechanisms of abnormal astrocyte Ca\(^{2+}\) signaling may improve cerebral blood flow and provide benefit to individuals with a range of brain pathologies.

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45. Ethanol-Induced Gating Alterations in Homomeric slo1 Versus \(\beta1\)-Containing BK Channels. GURUPRASAD KUNTIMALLAPPANVAR and ALEX M. DOPICO, Department of Pharmacology, College of Medicine, The University of Tennessee Health Science Center, Memphis, TN 38163

Calcium and voltage-gated, large conductance potassium (BK) channels result from tetrameric association of slo1 proteins (BK \(\alpha\) subunits). In most mammalian tissues, BK channels consist of \(\alpha\) and \(\beta\) subunits. The latter do not form channels themselves but affect the BK phenotype (Torres et al. 2014, Front. Physiol. 5:383). Four \(\beta\) subtypes have been identified, with \(\beta1\) being predominantly expressed in smooth muscle (Brenner et al. 2000, J. Bio. Chem. 275:6453–6461). Intoxicating levels of ethanol (10–100 mM) usually increase the open probability (Po) of homomeric slo1 channels while decreasing Po of \(\beta1\)-containing BK complexes (Dopico et al. 2014, Front. Physiol. 5:466). The gating modifications leading to these differential ethanol actions on BK Po have remained
unresolved. We tested ethanol action on slo1 (cbv1) and β1 subunits cloned from rat cerebral artery myocytes and expressed in *Xenopus* oocytes. Single channel and macroscopic currents were obtained from inside-out patches before and after 1–2 min exposure to 50 mM ethanol. We used the Horrigan Aldrich (HA) allosteric model of BK channel gating to address which allosteric parameters and gating processes were altered by ethanol. Data show that ethanol-induced change in cbv1-mediated ionic current is associated with a ×9 time increase in the channel’s apparent calcium binding affinity ($K_a = 9.02 \pm 1.4$ vs. $1 \pm 0.07$), whereas there is no significant change in voltage-dependent ($V_{0.5}(j) = 155$ vs. 155.1 ± 2; $z_2 = 0.6 \pm 0.04$ vs. 0.57 ± 0.02; $D = 19.7 \pm 1.4$ vs. 19.7 ± 0.61) or intrinsic gating parameters ($L_0 \times 10^{-6} = 6.3 \times 10^{-7}$ vs. $2.1 \times 10^{-6} \pm 2.8 \times 10^{-7}$; $z_L = 0.34 \pm 0.05$ vs. 0.35 ± 0.04). On the other hand, ethanol modification of cbv1+β1-mediated current is due to reduced allosteric interactions between calcium binding-channel opening (i.e., decreased allosteric factor C by ∼50%; 27.2 ± 2.3 vs. 13.54) and calcium binding-voltage sensor activation (i.e., decreased allosteric factor E by ∼50%; 9.5 ± 1.6 vs. 4.06 ± 0.5). Thus, while ethanol increases the apparent calcium affinity of both slo1 and slo1+β1 channels, ethanol-induced disruption of allosteric coupling in slo1+β1 override drug-induced decrease in $K_a$, leading to overall decrease in $P_0$.

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46. Pharmacological and Genetic Modifiers of a Ca2,1 Channelopathy. AMY LEE, University of Iowa, Cedar Falls, IA 50614

Voltage-gated ion channels exhibit complex properties, which can be targeted in pharmacological therapies for disease. We have found that the pro-oxidant, tert-butyl dihydroquinone (BHQ), modulates Ca2,1 Ca2+ channels in ways that oppose defects in channel gating and synaptic transmission, resulting from a familial hemiplegic migraine mutation (S218L). BHQ slows deactivation, inhibits voltage-dependent activation, and potentiates Ca2+ dependent facilitation of Ca2,1 channels in transfected HEK293T cells. These actions of BHQ help offset the gain of function and reduced Ca2+ dependent facilitation of Ca2,1 channels with the S218L mutation. Transgenic expression of the mutant channels at the *Drosophila* neuromuscular junction causes abnormally elevated evoked postsynaptic potentials and impaired synaptic plasticity, which are largely restored to the wild-type phenotypes by BHQ. Our results reveal a mechanism by which a Ca2,1 gating modifier can ameliorate defects associated with a disease-causing mutation in Ca2,1. We are currently screening for genetic modifiers of “flygraine” phenotypes in S218L-expressing flies in an effort to identify new signaling pathways that may be targeted in novel therapies.

47. Using Knockout Approaches To Tease Apart Physiological Roles of Slo Family Ion Channels and the Contributions of Regulatory Subunits. CHRISTOPHER LINGLE, Washington University School Medicine, St. Louis, MO 63130

The Slo family of four mammalian genes encode large conductance K+ channels that are regulated by cytosolic soluble ions, Slo1 by Ca2+, Slo2.1 and Slo2.2 by Na+, and Slo3 by alkalization. In some cases, additional regulatory b and g subunits help define key tissue-specific physiological properties. In previous work, important advances in understanding the physiological roles of Slo1 BK-type channels have been gained from development of KO animals for the Kcnma1 (BK α subunit) gene, the Kcnmb1 (BK β1) gene, and the Kcnmb4 (BK β4) gene. Such models have provided insight into potential roles of the BK channel in hypertension, epilepsy, and motor coordination with additional roles likely to be identified. Now, new mouse models of other Slo family subunits have become available, including Kcnv1 (Slo3), Lrrc52 (Slo3 y2 subunit), Kcnmb2 (BK β2), Slo2.1, and Slo2.2. Together, the Slo3 and Lrrc52 KOs reveal a Slo3/Lrrc52 partnership is critical to mouse fertility. The BK β2 KO reveals a critical role of β2 containing BK channels in shaping repetitive firing and, unexpectedly, that the absence of β2 subunits may favor spontaneous slow-wave bursting in some cells. Slo2.2 KO results in acute sensory phenotypes and, most obviously, an enhanced initial response to itch stimuli that arises from a reduced threshold for action potential generation in DRG neurons. Although the different physiological systems influenced by these subunits are only beginning to be examined, in each case this general approach has so far revealed novel aspects of physiological roles of these ion channels which had not been previously anticipated.

48. Translational Control of Ion Channel Composition. FANG LIU,1 DAVID K. JONES,1 WILLEM DE LANGE,2 and GAIL A. ROBERTSON,1,*1Department of Neuroscience, 2Department of Pediatrics, and Cardiovascular Research Center, Wisconsin Institutes of Medical Research, University of Wisconsin School of Medicine and Public Health, Madison, WI 53705

Ion channel function depends on subunit composition and stoichiometry. How the subunits find each other and assemble into complexes with the appropriate composition is poorly understood. We observed that alternate transcripts encoding hERG1a and 1b subunits, which assemble to produce ion channels mediating cardiac repolarization, are physically associated during translation. This association was first suggested by the observation that shRNA specifically targeting either hERG1a or 1b transcripts reduced levels of both transcripts when they were co-expressed heterologously. Native hERG1a and 1b transcripts in cardiomyocytes
derived from human-induced pluripotent stem cells were similarly knocked down by either shRNA, whereas KCNE1 or RYR2 levels were unaffected. The shRNA targeted transcripts undergoing translation, as opposed to aggregates destined for degradation, as confirmed by reductions in corresponding protein levels assayed by Western blot, quantitative immunocytochemistry, and membrane current recordings. To further test the hypothesis that co-knockdown of transcripts reflects a physical association, we immunoprecipitated nascent hERG1a protein from polysomal preparations and found that both the hERG1a and 1b transcripts co-purified. This association occurred even when translation of 1b protein was prevented by mutating the translation start site, indicating that the association could not be explained by cotranslational association of the nascent proteins and is likely attributable to the action of an RNA-binding protein. The physical association of transcripts encoding different subunits provides the spatial proximity required for nascent proteins to interact during biogenesis and may represent a general mechanism facilitating assembly of heteromeric protein complexes involved in a range of biological processes.

49. cPLA\textsubscript{2a}/α\textsuperscript{−/−} Sympathetic Neurons Exhibit Increased Membrane Excitability and Loss of N-Type Ca\textsuperscript{2+} Current Inhibition by M\textsubscript{1} Muscarinic Receptor Signaling. LIWANG LIU,\textsuperscript{1} JOSEPH V. BONVENTRE,\textsuperscript{2} and ANN R. RITTENHOUSE,\textsuperscript{1} \textsuperscript{1}University of Massachusetts Medical School, Worcester, MA 01655; \textsuperscript{2}Renal Division, Brigham and Women’s Hospital, Harvard Medical School, Boston, MA 02115

Group IVa cytosolic phospholipase A\textsubscript{2} (cPLA\textsubscript{2a}) mediates GPCR-stimulated arachidonic acid (AA) release from phosphatidylcholine 4,5-bisphosphate (PIP\textsubscript{2}), located in plasma membranes. We previously found in superficial cervical ganglion (SCG) neurons that PLA\textsubscript{2} activity is required for voltage-independent N-type Ca\textsuperscript{2+} (N-) current inhibition by M\textsubscript{1} muscarinic receptors (M\textsubscript{1},R\textsubscript{s}). These findings are at odds with an alternative model, previously observed for M-type K\textsuperscript{+} (M-) current inhibition, where PIP\textsubscript{2} dissociation from channels and subsequent metabolism by phospholipase C suffices for M-current inhibition. To resolve cPLA\textsubscript{2a}’s importance, we investigated its role in mediating voltage-independent N-current inhibition of ~40% that follows application of the muscarinic agonist oxotremorine-M (Oxo-M). Using a multidisciplinary approach that combined lipid pharmacology, G\textsubscript{s} signaling, gene knockout technology, and ion channel biophysics, we found that cPLA\textsubscript{2a} is required for M\textsubscript{1}R\textsubscript{s}-mediated, voltage-independent N-current inhibition. cPLA\textsubscript{2a}’s role appears specific for voltage-independent N-current inhibition since voltage-dependent inhibition by M\textsubscript{2}/M\textsubscript{3} R signaling remained intact in cPLA\textsubscript{2a}/α\textsuperscript{−/−} SCG neurons. Moreover, M-current inhibition occurred independently of cPLA\textsubscript{2a}, documenting divergent M\textsubscript{1}R signaling mediating M-current and N-current inhibition. These findings support an emerging idea that multiple phospholipases act in highly specific ways to decrease PIP\textsubscript{2} levels at or near N-channels following M\textsubscript{1}R stimulation. To determine cPLA\textsubscript{2a}’s functional importance at the neuronal level, we compared action potential firing of cPLA\textsubscript{2a}/α\textsuperscript{−/−} and cPLA\textsubscript{2a}/α\textsuperscript{+/−} SCG neurons. Decreases in latency to first firing and in interspike interval resulted in a doubling of firing frequency in cPLA\textsubscript{2a}/α\textsuperscript{−/−} neurons. These unexpected findings identify cPLA\textsubscript{2a} as a tonically regulating neuronal membrane excitability.

50. Modeling Epileptic Encephalopathies With Patient-Derived Neurons. YU LIU, ANDREW TIDBALL, LUIS LOPEZ-SANTIAGO, YUKUN YUAN, JACY WAGNON, MIRIAM MEISLER, LORI ISOM, and JACK PARENT, University of Michigan Medical School, Ann Arbor, MI 48109

Reprogramming somatic cells to a pluripotent state via the induced pluripotent stem cell (iPSC) method offers an unparalleled approach for neurological disease modeling using patient-derived neurons. My laboratory has applied the iPSC approach to model severe childhood genetic epilepsies with patient-derived cells. I will provide some background on disease modeling with iPSCs and then discuss our recent work modeling epilepsy and encephalopathies caused by sodium channel mutations.

We initially generated patient-derived neurons to study epilepsy mechanisms in Dravet syndrome (DS), a catastrophic childhood epilepsy caused by de novo dominant mutations and haploinsufficiency of the SCN1A gene that encodes the voltage-gated sodium channel Na\textsubscript{1.1}. The talk will describe our findings of altered sodium currents and excitability in DS patient neurons. Then I will describe recent work examining potential epilepsy mechanisms in another severe childhood epilepsy caused by putative gain-of-function mutations in the SCN8A gene that encodes Na\textsubscript{1.6}. Compared with control iPSC neurons, mutant SCN8A patient-derived neurons show increased persistent sodium current and hyperexcitability. Using a multi-well multi-electrode array for drug screening, we are validating the model with drugs known to work or to be ineffective in patients with SCN8A-associated epilepsy. I will also describe related work in mouse models of these disorders. Taken together, our work suggests that the iPSC approach offers great promise for modeling childhood epileptic encephalopathies and should provide a useful platform to identify novel therapies.

51. Critical Role of Nitric Oxide in Capillary-to-Arteriole Electrical Signaling in the Brain. THOMAS A. LONGDEN and MARK T. NELSON, Department of Pharmacology, College of Medicine, University of Vermont, Burlington, VT 05405; Institute of Cardiovascular Sciences, University of Manchester, Manchester, England, UK
Brain capillary ECs are ideally positioned to detect and respond to local neuronal activation by signaling for increases in blood flow, but the role of capillaries in this context has so far received little attention. We have shown that capillary ECs are capable of detecting potassium (K⁺) ions released during neuronal activity, which leads to inward rectifier K⁺ (KᵢR) channel-mediated retrograde electrical signaling to dilate upstream PAs and increase CBF (Longden et al. 2015, J. Gen. Physiol. 146:10A). Nitric oxide (NO) is required for the proper function of multiple neurovascular coupling pathways in the brain, including K⁺-evoked cerebral blood flow responses in vivo (Dreier et al. 1995, J. Cereb. Blood Flow Metab. 15:914–919). Here, we aimed to explore whether NO contributes to capillary-to-arteriole electrical signaling in the brain.

Using the perforated configuration of the patch clamp technique, we observed capillary EC KᵢR currents sensitive to 100 μM Ba²⁺. Interestingly, incubation of cells with the NO synthase inhibitor L-NNA (100 μM) markedly reduced KᵢR channel current density. Using an in vivo cranial window model to enable visualization of microcirculatory hemodynamics in the brain, we observed that selective delivery of K⁺ to capillaries produced robust hyperemia within seconds, due to activation of capillary KᵢR channels. This response was almost abolished upon superfusion of the cortical surface with 100 μM L-NNA.

Brain capillaries constitute an active sensory web, converting changes in external K⁺ into rapid upstream electrical signaling to regulate blood flow into the brain. These results suggest that NO has a critical role in maintaining capillary KᵢR channel activity to permit ongoing electrical communication with upstream arterioles.

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52. Alteration of Gating Properties in Ca₃.1 Channel Induced by Aspartic Residues in Its Pore. IGNACIO LÓPEZ-GONZÁLEZ, EDGAR GARZA-LÓPEZ, and TAKUYA NISHIGAKI, Departamento de Genética del Desarrollo y Fisiología Molecular, Instituto de Biotecnología, UNAM Av. Universidad #2001, Col. Chamilpa, Cuernavaca, Morelos, Mexico CP 62210

The simple and classical view of voltage-gated ion channels proposes that amino acid residues of the pore selectivity filter do not have a strong influence on the gating properties of the channel. However, previous reports have questioned this classical point of view. Regarding low voltage-activated (LVA) Ca²⁺ channels, it has been demonstrated that substitutions of pore aspartate residues (D) by glutamate residues (E) in domains III and IV alters the channel gating properties with the same tendency (Talavera et al. 2003, J. Gen. Physiol. 121:529–540), suggesting certain pore symmetry. In the present report, we evaluated the role of pore (E) residues of domains I and II on the Ca₃.1 channel gating properties. Our results indicate that substitution of (E) residues in the pore loops of domains I and II by (D) residues differentially modify the gating properties of Ca₃.1 channel. The single mutant channel (DEDD, domain I) presented slight uncoupling between the activation and inactivation processes, with a more stable inactivation state and a slower recovery from inactivation without change in the deactivation kinetics. In contrast, the single mutant channel (EDDD, domain II) presented a less stable close state, allowing an easier channel transition to the open state, with inactivation kinetics similar to WT Ca₃.1 channels and slower deactivation kinetics. At last, the double mutant channel (DDDD, domains I and II) presented completely uncoupled activation and inactivation processes; faster activation, inactivation, and deactivation kinetics; and slower deactivation process than the WT Ca₃.1 channel. Taking into account our results, we can conclude that the selectivity filter residues of LVA Ca²⁺ channels have a differential influence on the gating properties of this Ca²⁺ channel subfamily, suggesting a pore pseudo-symmetry. In a speculative way, we propose these new mutations in the selectivity filter of the Ca₃.1 channel could be important in some types of epilepsies (Chen et al. 2003, Ann. Neurol. 54:29–243; McKeown et al. 2006, Acta Pharmacol. Sin. 27:799–812; Heron et al. 2007, Ann. Neurol. 62:560–568).

53. FLARE Biosensors for Multiparametric Imaging. MICHELE L. MARKWARDT, BRIAN ROSS, JENNIFER MCFARLAND, JIN ZHANG, and MARK A. RIZZO

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Optical tools designed to track activation of key cell signaling pathways have revealed the spatial and temporal dynamics of intracellular communication with unprecedented resolution. Even so, many of these studies are practically limited to tracking a single endpoint because of limitations set by the intrinsic optical properties of the sensors. This is problematic because intracellular communication pathways interact in complex ways in time and space. For example, calcium and cAMP signals frequently display oscillatory temporal dynamics and tight spatial compartmentalization that can influence one another. Thus, better tools are needed to permit quantitative measurement of multiple signaling pathways in living cells and tissues and enable studies of complex cellular processes, including how cells integrate information from competing signals and how progressive diseases affect a cell’s ability to process information. Here, we describe a novel FRET-biosensor strategy that is optimized for multiparameter measurements in living cells.
cells, both cultured and in vivo, using homotransfer fluorescence anisotropy reporters (FLAREs). The FLARE sensors are generally constructed from existing, validated heterotransfer sensors, but include FRET pairs of the same color that permit FRET detection using fluorescence polarization microscopy. Single-color FLARE biosensors for calcium, cAMP, voltage, myosin kinase, and myosin phosphorylation have been constructed in multiple colors, enabling measurement of up to three intracellular signals in a single preparation. The general principle of the method will be discussed, as well as evidence supporting the utility of this technique for in vivo imaging in living organisms. In addition, we present evidence that inducing progressive endoplasmic reticulum (ER) stress in excitable cells leads to ER calcium retention and reduced cytoplasmic calcium. This cripples cell–cell communication by decoupling GPCR signaling from activation of ER calcium channels through calcium-induced calcium release.

54. Progressive Aortic Valve Stenosis in the Mouse with a Gain-of-Function Mutation in Ca$_{1.2}$ (CACNA1C). MAIKO MATSUMI,$^1$ KUSHAL KADAKIA,$^2$ ERIC WEI,$^2$ DANIEL SINDEN,$^3$ and GEOFFREY PITT.$^1$ $^1$Weill Cornell Medical College, Cornell University, New York, NY 10065; $^2$Duke University Medical Center, Durham, NC 27710

Calcific aortic valve stenosis (CAVS) is a life-threatening disorder affecting $\sim$2% of people $>$65 years. Previously thought to be due to “wear and tear,” studies show that osteoblast-like and osteoclast-like cells are found in human CAVS and osteogenic pathways such as BMP and Wnt signaling pathways are activated in calcific valves. Importantly, genome-wide association studies recently identified CACNA1C, encoding the $\alpha$ subunit of voltage-dependent L-type calcium channel Ca$_{1.2}$, as a new CAVS susceptibility gene, and expression quantitative trait loci (eQTL) mapping suggested that increased Ca$^{2+}$ influx through the channel drove the phenotype. Beyond this genetic association, little is known about roles for Ca$_{1.2}$ in the aortic valve and how increased Ca$^{2+}$ influx through Ca$_{1.2}$ contributes to CAVS.

In this study, we exploited a Ca$^{2+}$lce reporter line to define expression patterns in valves and found Ca$^{2+}$lce is expressed primarily in the annulus of adult aortic valves.

We generated two mouse lines. For both mouse models, we evaluated valves by histology to assess CAVS and examined transcriptional responses by qPCR.

The first mouse line, A G406R knock-in mutation in Ca$^{2+}$lce, which decreases channel inactivation and leads to Ca$^{2+}$ influx, displayed chondrocyte-like transformation of cells in the attachment of the cusps to the annulus. These areas were markedly thickened compared with wild-type littermate controls and showed evidence of accelerated calcification. In the second mouse line, we activated a G406R mutant Ca$^{2+}$lce transgene specifically in valve interstitial cells (VICs) with a Cre recombinase driven by the transcription factor Scleraxis (Scx). Ectopic activation of the G406R mutant transgene in VICs by Scx-Cre resulted in marked thickening of the valve cusps. Our results indicate that increased Ca$^{2+}$ influx through Ca$_{1.2}$ in the aortic valve leads to thickened valves and calcification. Ectopic expression of the mutant Ca$_{1.2}$ in valve cells demonstrates that the role of Ca$_{1.2}$ is cell autonomous for the development of CAVs.

55. Molecular Mechanisms of K$_{ATP}$ Channel Mutations in Cantu Syndrome. CONOR MCCLENAGHAN,$^{1,2}$ MONICA SALA-RABINAL,$^{1,2}$ PAIGE COOPER,$^{1,2}$ RISHA SHAH,$^{1,2}$ HAIXIA ZHANG,$^{1,2}$ THERESA HARTER,$^{1,2}$ CHRIS EMFIGER,$^{1,2}$ BLANCHIE SCHWAPPACH,$^3$ MARIA REMEDI,$^{1,2}$ and COLIN NICHOLS.$^{1,2}$ $^1$Department of Cell Biology and Physiology, and $^2$The Centre for the Investigation of Membrane Excitability and Disease, Washington University School of Medicine, St. Louis, MO 63130; $^3$Department of Molecular Biology, University Medicine Göttingen, Germany; $^4$Division of Endocrinology, Metabolism and Lipid Research, Washington University School of Medicine, St. Louis, MO 63130

Cantu syndrome (CS) is a rare condition characterized by craniofacial dysmorphology, hypertrichosis, osteochondrodysplasia, and various cardiovascular symptoms including cardiomegaly (Cantu et al. 1982. Hum. Genet. 60:36–41). ATP-sensitive potassium channels (K$_{ATP}$ channels) are formed as hetero-octameric complexes of Kir6.x and SUR subunits. The recent identification of gain of function (GoF) mutations in ABCC9 (SUR2) and KCNJ8 (Kir6.1) in numerous CS patients implicates K$_{ATP}$ channels in the syndrome (Harakalova et al. 2012. Nat. Genet. 44:793–796; Cooper et al. 2014. Hum. Mutat. 35:809–813). How these mutations affect channel function and how this relates to the pathophysiology of CS is the subject of current investigation. Here, we demonstrate that two hitherto uncharacterized mutations, R1150W (R1154W in hSUR2) and S1050Y (S1054Y in hSUR2), result in increased currents in the presence of Mg-nucleotides yet have no effect on ATP inhibition in the absence of magnesium, demonstrating that these mutations augment Mg-nucleotide activation. Interestingly, the effect of the R1150Q mutation on nucleotide regulation appears more subtle. This mutation occurs close to an exon/intron boundary, and characterization of the recently developed CS mouse SUR2[R1150Q] indicates that the mutation results in alternative splicing of SUR2A and subsequent expression of a transcript encoding for a nonfunctional subunit (R1150Qdel exon 30). Further investigation suggests that decreased SUR2 protein expression may be compensated by increased SUR1 in ventricular myocytes, which would contribute to K$_{ATP}$ GoF due to the increased Mg-nucleotide sensitivity of the SUR1 subunit. These preliminary results point toward diverse molecular...
Mechanisms of exonic SUR2 mutations in CS, including modulation of splicing in addition to direct alteration in nucleotide sensitivity. The strategy to further delineate the effects of \( K_{\text{ATP}} \) channel mutations in CS, including biophysical characterization and transgenic mouse studies, is discussed.

56. Mechanistic Analysis of SNP effects on human BK Channel Properties. BETH A. MCNALLY, AMBER E. PLANTE, and ANDREA L. MEREDITH, University of Maryland School of Medicine, Baltimore, MD 21201

The human large conductance calcium-activated potassium channel (BK) is encoded by a single gene (KCNMA1), and we recently found six single nucleotide polymorphisms (SNPs) in the KCNMA1 coding region that alter BK channel properties under standard testing conditions (0, 1, 10, and 100 \( \mu \)M Ca). Here, we selected three of these SNPs (A138V, C495G, and R800W) and further probed potential mechanisms of their altered BK channel properties. Since an autism-linked SNP (A138V) is in close proximity to the BK channel Mg\(^2+\) coordination site, specifically residue D164 (D99 in mbr5), we examined Mg\(^2+\) gating. We found that physiological Mg\(^2+\) concentrations (1 and 3 mM) resulted in a leftward G-V shift for both A138V and WT. A138V was significantly right shifted at 1 mM Mg\(^2+\) / 10 \( \mu \)M Ca\(^2+\) (\( P < 0.05 \)) when compared with WT. Second, we examined whether the C495G SNP alters the redox/oxidation of the channel, since this residue (C430 in hSlo1) was previously shown to contribute to the rightward shifting effects of cysteine oxidation on BK channel properties. Next, we examined whether the rightward shifting effect of the R800W SNP was due to size or charge by testing R800A, R800E, and R800Q mutations, respectively. Lastly, we used the human gain-of-function mutation, D434G, which increases BK currents by altering the coupling of Ca\(^2+\) binding and channel opening to assess the distinct mechanistic effect of C495G and R800W on channel properties. The D434G leftward shift was not altered in the presence of C495G, whereas the R800W rightward shift was decreased by the presence of D434G, suggesting that R800W and D434G can exert distinct and opposite effects on channel properties. Together, these results begin to reveal the gating mechanisms acted upon by human SNP variation and suggest a disease-linked mutation could be mitigated by SNP variation at distal residues.

57. Rhythmic Regulation of BK Splice Variants in the Circadian Clock. ANDREA MEREDITH, University of Maryland School of Medicine, Baltimore, MD 21201

BK Ca\(^2+\)– activated K\(^+\) currents exhibit diverse properties across tissues. The functional variation in voltage- and Ca\(^2+\)-dependent gating underlying this diversity arises from multiple mechanisms, including alternate splicing of \( Kenma1 \), the gene encoding the pore-forming (\( \alpha \)) subunit of the BK channel, phosphorylation of \( \alpha \) subunits, and inclusion of \( \beta \) subunits in channel complexes. One physiologically-integrated system for studying the molecular basis of BK current diversity is found in the brain’s central clock, the suprachiasmatic nucleus (SCN) of the hypothalamus. We identified a novel role for BK channels in the daily patterning of neuronal activity in the SCN, correlated with a day versus night difference in BK macroscopic current levels, but the channel-based mechanisms that produce rhythmic action potential activity are not fully understood. In the SCN, alternative splicing is under circadian control, and the abundance of two \( Kenma1 \) splice variants (BK\(_{\text{SRKR}}\) and BK\(_{\alpha} \)) differs between day and night. Currents recorded from “day” BK\(_{\text{SRKR}}\) channels in HEK293 cells had a significantly right-shifted current-voltage relationship across a range of Ca\(^2+\) concentrations, slower activation, and faster deactivation, compared with “night” BK\(_{\alpha} \) channels. These effects were dependent upon phosphorylation of S642, a constitutive serine immediately preceding the SRKR insert, and were abolished by alkaline phosphatase or mutation of S642. To test this mechanism in daytime SCN neurons, alkaline phosphatase was applied intracellularly, resulting in increased BK current in response to both step and action-potential commands. The change in BK current was correlated with a decrease in native spontaneous action potential firing rate. This work identifies alternative splicing of BK channels as a genetic mechanism that may contribute to the dynamic regulation of excitability in the circadian pacemaker.

58. Defining New Mechanisms Underlying Cardiovascular Disease. PETER J. MOHLER, Davis Heart and Lung Research Institute, The Ohio State University Medical Center, Columbus, OH 43210

Our research is focused on the mechanisms underlying the targeting and regulation of membrane-associated (ion channels, transporters, and receptors) and signaling proteins in cardiac and other excitable cells. In particular, we are interested in the role of membrane-associated ankyrin and spectrin family of polypeptides in the targeting and function of ion channels and transporters as well as kinases and phosphatases. A primary focus of the laboratory is the role of the ankyrin-G-based pathway for targeting voltage-gated sodium channels to the intercalated disc of cardiomyocytes. We have discovered a direct requirement of ankyrin-G for Na channel targeting and have linked human Na channel arrhythmia mutations with loss of ankyrin-G binding, and Na channel targeting resulting in defects in Na channel function and myocyte excitability. A second line of work in the laboratory establishes that loss-of-function mutation in ankyrin-B is the basis for a human cardiac arrhythmia syndrome associated with sinus node dysfunction, repolarization defects, and polymorphic
tachyarrhythmia in response to stress and/or exercise (“ankyrin-B syndrome”). Additionally, our work revealed that reduction of ankyrin-B in mice results in reduced levels and abnormal localization of Na/Ca exchanger, Na/K ATPase, and InsP3 receptor at T-tubule/SR sites in cardiomyocytes and leads to altered Ca2+ signaling and extrasystoles that provide a rationale for the arrhythmia. These studies establish a physiological requirement for ankyrins and spectrins in localization of a variety of ion channels in excitable membranes in the heart and demonstrate a new class of functional “channelopathies” due to abnormal cellular localization of functionally related ion channels and transporters. More recently, we have developed a third line of research in the laboratory focused on the molecular mechanisms underlying kinase and phosphatase targetting in excitable cardiomyocytes. Specifically, work from our laboratory has shown the importance of CaMKII and PP2A targeting for myocyte and cardiac function.

59. Potentiation of Excitatory Synaptic Transmission in the Superficial Dorsal Horn by Low Concentrations of Kv Channel Inhibitors. TANZIYAH MUQEM,1 VITOR PINTO,2 and MANUEL COVARRUBIAS.1 1Department of Neuroscience and Farber Institute for Neurosciences, Thomas Jefferson University, Philadelphia, PA 19107; 2ICVS/3Bs, University of Minho, Braga, Portugal

Kv3.4 channels underlie a majority of the high-voltage activating K+ current in dorsal root ganglion (DRG) nociceptors and are expressed in all functional compartments of these neurons, including synaptic terminals. Kv3.4 channels in DRG neurons regulate the rate of action potential repolarization in a manner that depends on phosphorylation of the channels’ inactivation domain. Additionally, Kv3.4 channel down-regulation is implicated in the pathophysiology of chronic pain induced by spinal cord injury. We hypothesize that, through their ability to facilitate action potential repolarization, Kv3.4 channels help keep nociceptive synaptic transmission in check. Kv3.4 channel down-regulation would, therefore, promote pain transduction. To test this hypothesis, we have investigated excitatory synaptic transmission in the superficial dorsal horn under conditions that inhibit the Kv3.4 current in DRG nerve terminals. We used an ex vivo rat cervical spinal cord preparation suitable for whole-cell patch clamping of secondary neurons in the dorsal horn. Since Kv3.4 channels are hypersensitive to submillimolar concentrations of 4-aminopyridine (4-AP) and tetraethylammonium (TEA), we tested their effects on excitatory postsynaptic currents (EPSCs) in laminae I and II of the dorsal horn. We found that 500 μM TEA potentiates the EPSCs by 53% (P < 0.001, paired t test, n = 3). Similarly, 50 μM 4-AP potentiates the EPSCs by 59% (P < 0.001, paired t test, n = 3). We conclude that the Kv3.4 channel expressed in nociceptors is a plausible regulator of nociceptive synaptic transmission in the spinal cord. Currently, we are testing additional inhibitors to evaluate possible contributions of other K+ channels and exploring more specific knockdown strategies to assess the role of the Kv3.4 channel as a regulator of nociception in vivo.

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60. Potassium Channelopathy-Like Defect Underlies Early-Stage Cerebrovascular Dysfunction in a Genetic Model of Small Vessel Disease. MARK NELSON, University of Vermont, Burlington, VT 05405

Cerebral autosomal-dominant arteriopathy with subcortical infarcts and leukoencephalopathy (CADASIL), caused by dominant mutations in the Notch3 receptor in vascular smooth muscle, is a genetic paradigm of small vessel disease (SVD) of the brain. Recent studies using transgenic (Tg)Notch3R169C mice, a genetic model of CADASIL, revealed functional defects in cerebral (pial) arteries on the surface of the brain at an early stage of disease progression. Here, using parenchymal arterioles (PAs) from within the brain, we determined the molecular mechanism underlying the early functional deficits associated with this Notch3 mutation. At physiological pressure (40 mmHg), smooth muscle membrane potential depolarization, and constriction to pressure (myogenic tone) were blunted in PAs from TgNotch3R169C mice. This effect was associated with an ~60% increase in the number of voltage-gated potassium (Kv1.5) channels, which oppose pressure-induced depolarization. Inhibition of Kv1.1 channels with 4-aminopyridine or treatment with the epidermal growth factor receptor agonist heparin-binding EGF (HB-EGF), which promotes Kv1.1 channel endocytosis, reduced Kv current density and restored myogenic responses in PAs from TgNotch3R169C mice, whereas pharmacological inhibition of other major vasodilatory influences had no effect. Kv1.1 currents and myogenic responses were similarly altered in pial arteries from TgNotch3R169C mice, but not in mesenteric arteries. Interestingly, HB-EGF had no effect on mesenteric arteries, suggesting a possible mechanistic basis for the exclusive cerebrovascular manifestation of CADASIL. The metalloproteinase inhibitor, TIMP3, accumulates in the extracellular matrix of PAs from the CADASIL mouse model and in human CADASIL patients. Genetic overexpression of TIMP3 mimicked the effects of CADASIL on Kv current density and the loss of myogenic tone. Collectively, our results indicate that increasing the number of Kv1.1 channels in cerebral smooth muscle produces a mutant vascular phenotype akin to a channelopathy in a genetic model of SVD.

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61. Cantu Syndrome: Multi-Organ Complexities from KCN8 Gain-of-Function. **COLIN G. NICHOLS,**1
HAIXIA ZHANG,1 PAIGE E. COOPER,1 JIN-MOO LEE,1 ELAINE L. SHELTON,2 MICHAEL J. DAVIS,3
MARK A. LEVIN,1 MARIA S. REMEDI,1 GAUTAM K. SINGH,1 and DOROTHY K. GRANGE,1
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Cantu syndrome (CS, MIM 239850) is a complex congenital disorder with multiple cardiovascular features. Hypertrichosis is a constant feature, with coarse facial features, generalized macrosomia, and macrocephaly typically presenting at birth. Cardiovascular features include enlarged and hypercontractile hearts, hypertension, pericardial effusion, persistent patent ductus arteriosus (PDA) and fetal brain circulation, and lymphedema. In the last three years, we have demonstrated that GOF mutations in the genes encoding both the regulatory (ABCC9) and pore-forming (KCNJ8) subunits of KATP channels underlie the disease in the majority of patients, but such findings have not elucidated the mechanisms underlying the myriad syndrome features. Our development of relevant transgenic animal models and a unique research CS clinic have permitted extensive characterization of disease phenotype and have led us to unique insights to the underlying pathology of cardiac, vascular, and lymphatic features.

62. Precision Physiology and Rescue of Ion Channel Disorders: The Gate is Now Wide Open. **JEFFREY NOEBELS,** Baylor College of Medicine, Houston, TX 77030

A steady drumroll of genetic discoveries linking ion channel mutations to human disease highlights the urgent need for a deeper understanding of how altered biophysics and cell biology of channel function can be therapeutically reversed. Ion channel subunits now comprise the single largest gene family underlying disorders of heart, muscle, and brain, and the most frequently tested for precision clinical diagnosis of a broad phenotypic spectrum of cognitive disability, neuropsychiatry, epilepsy, sensorimotor dysfunction, and neurodegenerative disease. These disorders collectively constitute an enormous public health burden, with a greater number of life years diminished or lost than cancer. The significance of each variant, which in some cases spells the difference between lifelong disability or sudden death, requires accurate functional interpretation in order to stimulate drug discovery and guide the use of mutation-specific therapies.

What is missing from this brave new agenda? There is a profound lack of functional information regarding the majority of roughly 400 genes and their myriad splice forms, as well as the unexpected complexity of their coordinate regulation and interaction within cells. This deficit poses a major obstacle to successful therapeutic translation of precision diagnostic testing. The results of clinical exomes point to recurring mysteries that are impossible to ignore: seemingly slight or no changes in gating kinetics may lead to devastating disease, implicating the involvement of unknown nonpore functions. Shared subunit expression in heart and brain may lead to either unpredictable patterns of compensatory functional sparing or devastating patterns of selective vulnerability.

The arrival of genetic testing has permanently altered the ion channel basic research mandate. Focused studies of the functional biology of human channel mutations are essential to the success of precision medicine. New basic/clinical collaborative approaches are required to uncover, validate, and model meaning, either singly or in complex combinations. Fortunately, rather than distracting from the fundamental goal of defining canonical protein function, a focus on the cell and clinical biology of mutant channels accelerates discovery of unsuspected aspects of channel gating, interaction domains, location, remodeling, and epistasis. At higher levels of network organization, mutations provide insight into activity-driven wiring of developing brain circuitry. Finally, genetic editing strategies are already beginning to repair point mutations of ion channels in model systems to confirm and explore the disease mechanism. The rich biology and powerful approaches explain why ion channel mutations represent some of the most intriguing and medically essential molecular lesions to understand and treat.

63. Targeting Sodium Channel Slow Inactivation as a Novel Therapeutic Approach in Myotonia Congenita. **KEVIN NOVAK,**1 WILLIAM ARNOLD,2 ALAN SANDERSON,2 JOHN KISSEL,2 and MARK RICH,1 1Wright State University, Dayton, OH 45435; 2The Ohio State University, Columbus, OH 43210

Patients with myotonia congenita have muscle hyperexcitability due to loss-of-function mutations in the chloride channel in skeletal muscle, which causes spontaneous firing of muscle action potentials (myotonia), producing muscle stiffness. In patients, muscle stiffness lessens with exercise, a change known as the warmup phenomenon. Our goal was to identify the mechanism underlying warmup and to use this information to guide development of novel therapy. Exercise was mimicked in vitro in individual muscle fibers by stimulating 5,000 muscle action potentials at a rate of 20 Hz. Action potential morphology was assessed before and after the repeated stimulation. Significant decreases in action potential height and rate of rise (dV/dt) suggested increased sodium channel slow inactivation contributes to warm up. These data suggested that enhancing slow inactivation of sodium channels might offer effective therapy for myotonia. Lacosamide and ranolazine enhance
slow inactivation of sodium channels and are approved by the US Food and Drug Administration for other uses in patients. We compared the efficacy of both drugs with mexiletine, a sodium channel blocker currently used to treat myotonia. In vitro studies suggested that both lacosamide and ranolazine were superior to mexiletine. However, in vivo studies in a mouse model of myotonia congenita suggested that side effects could limit the efficacy of lacosamide. Ranolazine produced fewer side effects and was as effective as mexiletine at a dose that produced none of mexiletine’s hypexcitability side effects. Based on our preclinical studies, we performed an open label, proof of concept trial in patients with myotonia congenita to assess the tolerability and efficacy of ranolazine. Participants demonstrated less self-reported stiffness, improved motor performance, and had reduced electromyographic myotonia. No participant discontinued ranolazine due to side effects. These results suggest ranolazine is well tolerated at doses that significantly improve motor symptoms and lessen myotonia and support a future controlled trial.

64. Na\(_{1.1}\) in A\(_\delta\) Fibers Mediates Mechanical Pain Signaling: Relevance for Migraine Pathogenesis. JEREMIAH D. OSTEEN,\(^1\) JOHN GILCHRIST,\(^2\) FRANK BOSMANS,\(^2\) and DAVID JULIUS.\(^1\) \(^1\)Department of Physiology, University of California, San Francisco, San Francisco, CA 94143; \(^2\)Department of Physiology and Solomon H. Snyder Department of Neuroscience, Johns Hopkins University School of Medicine, Baltimore, MD 21205

Migraine headache is a common, debilitating condition characterized by recurrent attacks of headache pain, sensory sensitivity and nausea. These attacks are sometimes preceded by aura, a sensory disturbance resulting from cortical spreading depression in the brain. Subsequent migraine pain and photophobia arise from sensitization of trigeminal primary afferents, including release of pro-inflammatory peptides. A third hallmark is vasodilation of the meningeal vasculature, which increases pressure on trigeminal afferents. The etiology of migraine has been a subject of debate, since the separation of cause and effect between brain, trigeminal, and vascular changes is difficult to distinguish. Recently, the Na\(_{1.1}\) subtype of voltage-gated sodium channel has been associated with an inherited form of migraine (familial hemiplegic migraine type 3—FHM3). Na\(_{1.1}\) is widely expressed in the brain, and migraine mutations are currently thought to act through a CNS-initiated mechanism. Here, we describe a spider toxin, Hm1a, which selectively activates Na\(_{1.1}\). Following injection of Hm1a into the hind paw of a mouse, we see acute pain behaviors as well as sensitization to mechanical, but not thermal stimuli, including evidence of robust central sensitization. We find that Na\(_{1.1}\) resides in primary afferent sensory neurons, including myelinated A\(_\delta\) fibers, but not within the most well-studied nociceptor class, the unmyelinated c fibers. Interestingly, we find that the main effects of Hm1a on Na\(_{1.1}\) mirror the effects of several studied FHM3 mutations: Hm1a shifts steady-state inactivation to more depolarized potentials and inhibits slow inactivation. In addition, Hm1a targets the Na\(_{1.1}\) domain IV voltage sensor, the same location around which many FHM3 mutations cluster. Taken together, our results demonstrate a peripheral role for Na\(_{1.1}\) in the development of mechanical pain and implicate hyperexcitability of Na\(_{1.1}\)-expressing A\(_\delta\) fibers in FHM3 pathogenesis. Based on these data, Hm1a can now be used to further probe migraine both centrally and peripherally, thereby helping to untangle migraine etiology.

65. Cysteine Mutagenesis Reveals TMC1 Residues that Contribute to Mechanotransduction. BIFENG PAN,\(^3\) XIAO-PING LIU,\(^3\) YUKAKO ASAI,\(^1\) KIYOTO KURIMA,\(^2\) ANDREW J. GRIFFITH,\(^2\) and JEFFREY R. HOLT,\(^1\) \(^1\)Department of Otolaryngology, F.M. Kirby Neurobiology Center, Boston Children’s Hospital, Harvard Medical School, Boston, MA 02115; \(^2\)Molecular Biology and Genetics Section, National Institute on Deafness and Other Communication Disorders, NIH, Bethesda, MD 20892

TMC1 and TMC2 have emerged as strong candidates for the hair cell transduction channel. A growing body of evidence supports a direct role for these molecules as necessary components of the channel. Tmc1/Tmc2 mRNAs are expressed in auditory and vestibular hair cells at the right time during development. TMC1 and TMC2 proteins are localized to the tips of shorter row stereocilia. The proteins interact with other known components of the hair cell transduction complex including PCDH15 and LHFP5. Deletion of Tmc1 and Tmc2 results in complete loss of conventional hair cell transduction, and reintroduction of either Tmc1 or Tmc2 cDNA leads to restoration of mechanotransduction. Hair cells expressing the Beethoven point mutation in Tmc1 have lower single-channel conductance, lower calcium permeability, and lower sensitivity to block by dihydrostreptomycin than cells expressing wild-type Tmc1. These data are consistent with the hypothesis that TMC1 and TMC2 can function as pore-forming subunits of the transduction channel. However, the pore region within the TMC1 topology has not been definitively identified.

To investigate structure–function relationships in TMC1 we generated AAV vectors that encoded mutant TMC1 sequences, introduced at several strategic sites within the TMC1 amino acid sequence. The vectors were injected via the round window membrane into the cochleas and vestibular organs of Tmc1/Tmc2 double knockout mice. We recorded conventional mechanotransduction from AAV-transduced auditory and vestibular hair cells and assayed for changes in hair cell transduction currents and selectivity. The data provide further evidence supporting a direct role for TMC1 in hair cell transduction and suggest a revised topology.
66. G Protein–Coupled Receptor (GPCR) Signaling Underlies the Nicotine-Induced Up-Regulation of alpha 7 (α7) Nicotinic Acetylcholine Receptors (nAChRs) Expressed in Xenopus Oocytes. JAYHARSH PANCHAL, KRISTI DEBOEUF, MOHAMMAD FARIDUL ISLAM, and JOSEPH FARLEY, Neuroscience, Indiana University, Bloomington, IN 47405

α7 nAChRs are widely distributed throughout the nervous system, playing important roles in learning, memory, several disease and neurodegenerative processes, and nicotine addiction. A variety of agonists and antagonists produce functional and/or numerical up-regulation of α7 Rs in different cells, implicating multiple signaling pathways/mechanisms. Prolonged nicotine exposure can also up-regulate α7 nAChRs, which may contribute to nicotine addiction. We found approximately twofold up-regulation of murine α7 nAChRs in Xenopus oocytes following 12 h of 100 μM nicotine and extensive washout. Nicotine up-regulation was dependent upon intracellular Ca\(^{2+}\), being abolished by BAPTA-AM, and involved several Ca\(^{2+}\)-dependent enzymes (e.g., PP2B and PKC). However, up-regulation was independent of Ca\(^{2+}\) influx, being unaffected by removal of extracellular Ca\(^{2+}\). The M3-M4 loop of α7 contains a conserved G protein–binding cluster (GPBC). G protein signaling by α7 Rs has been previously shown in neurons and PC12 cells (Kabbani et al. 2013. BioEssays. 25:1025–1034). Here, we show that GPCR signaling mediates nicotine up-regulation of α7 nAChR. We observed that a substance P-analogue peptide (a putative specific inhibitor of Gαq/11 binding to GPCRs) prevented nicotine-induced up-regulation of α7 Rs. However, the substance P-analogue also significantly reduced control α7 R currents, suggesting that its effects were nonspecific. Therefore, we mutated the α7 nAChR GPBC (RMKR to AAAAA; denoted as α7 344-347A) to block interaction of Gαq with the GPBC. Receptor expression levels, peak current amplitude, and kinetics were equivalent for mutant and wild-type (wt) α7 Rs. But nicotine-up-regulation of α7 344-347A R was completely inhibited. In contrast, prolonged exposure to the cell-permeable, competitive antagonist methyllycaconitine (MLA), produced approximately twofold up-regulation of both wt and mutant α7 Rs, which were unaffected by BAPTA-AM. MLA up-regulation of mutant and wt α7 Rs may result from a chaperone-like mechanism (Lester et al. 2009. APPS J. 11:167–177). GPCR signaling of α7 Rs is critical for their nicotine-up-regulation.

67. TRPC Channel Remodeling in a Mouse Model of Essential Hypertension: The TRPC3-TRPC6 Game of Thrones. M. TERESA PEREZ-GARCIA, INES ALVAREZ-MIGUEL, PILAR CIDAD, and JOSE R. LOPEZ-LOPEZ, Department of Biochemistry and Physiology and Institute of Biology and Molecular Genetics (IBGM), School of Medicine, Universidad de Valladolid, Spain

Essential hypertension involves a gradual, sustained rise in total peripheral resistance caused by an increased vascular tone. A model has been proposed in which the combination of membrane depolarization and higher L-type Ca\(^{2+}\) channel activity conspires to generate increased Ca\(^{2+}\) influx into vascular smooth muscle cells (VSMCs), contraction and vasoconstriction. The search for culprit ion channels that can drive membrane depolarization has provided several candidates, including some members of the TRPC family. TRPC3 and TRPC6 are DAG-activated, nonselective cationic channels contributing to stretch- or agonist-induced depolarization. Conflicting information regarding changes in TRPC3 or TRPC6 functional expression associated to hypertension has been reported. However, although TRPC3-TRPC6 channels can form homo- or heteromultimers, the possibility that differences in their association pattern may change their functional contribution to vascular tone is largely unexplored.

We probe this hypothesis using a model of essential hypertension (BPH mice) and its normotensive control (BN mice). We characterize homo- and heterotetramers with electrophysiological recordings in TRPC3-, TRPC6-, or TRPC3/6-transfected CHO cells. Nonselective basal or hypotonic stimulus-activated cationic currents were elicited with all the studied combinations. However, TRPC currents were sensitive to the selective antagonist Pyr10 only when TRPC6 was present. Meanwhile, intracellular anti-TRPC3 antibody selectively blocked TRPC3-mediated currents.

In isolated mesenteric VSMCs, basal and agonist-induced currents were more sensitive to Pyr3 and Pyr10 in BPH cells. Consistently, myography studies in pressurized mesenteric arteries showed a larger Pyr3/10-induced vasodilation in BN arteries. mRNA and protein expression data (explored with qPCR and proximity ligation assay) support changes in TRPC3 and TRPC6 proportion and assembly, with a higher TRPC3 channel contribution in BPH VSMCs which could favor cell depolarization in the hypertensive phenotype. These differences in functional and pharmacological properties of TRPC3 and TRPC6 channels depending on their assembly as homo- or heterotetramers could represent novel therapeutical opportunities.

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68. The KLHL1 KO as a Model of Altered T-Type Ca Channel Function. PAULA P. PERISSINOTTI,1 YUNGUI HE,2 ELIZABETH MARTINEZ-HERNANDEZ,3 ELIZABETH A. ETHINGTON,1 ERIC ALMAZAN,1 ALISSA ZEGLIN,1,2 MICHAEL D. KOOB,3 and ERIKA S. PIEDRAS-RENTERIA.1,4 1Cell and Molecular Physiology Department and 2Stritch School of Medicine, Loyola University Chicago, Chicago, IL 60660; 3Institute for...
The physiological consequences of the global deletion of the Kelch-like 1 protein (KLHL1) in mice will be discussed. KLHL1 is a neuronal actin-binding protein that modulates Ca\textsubscript{v}2.1 P/Q-type and Ca\textsubscript{v}3.2 T-type channel activity. Decreased expression and function of these two channel types is seen when KLHL1 is acutely down-regulated using specific shRNA; however, the KLHL1 KO mouse displays tissue-specific Ca current homeostasis that can result in normal, decreased, or increased Ca currents. Altered Ca channel function and Ca current levels, neuronal excitability changes, and synaptic function are detected in this model.

Here, we will discuss our most recent findings on the KLHL1 KO mouse as a model to assess altered T-type channel expression. The hypothalamus from KO mice exhibits slightly decreased Ca\textsubscript{v}3.2 expression as expected for this model, however Ca\textsubscript{v}3.1 levels are highly up-regulated, resulting in increased LVA channel activity, increased excitability of pro-opiomelanocortin–positive neurons, and T type channel-mediated resistance to the satiety hormone leptin, as well as abnormal responses after a 20-h period of food deprivation. In contrast, KO dorsal root ganglia display decreased LVA Ca\textsubscript{v}3.2 expression without additional compensations, resulting in decreased LVA currents, decreased DRG excitability, and increased threshold to mechanical pain. In summary, the KLHL1 KO mouse model is an excellent system to explore how moderate Ca channel function changes affect neuronal tissues. Our results also establish T-type channels as possible targets for obesity treatment.

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69. The Role of Ca\textsubscript{v}1.2 L-Type Ca\textsuperscript{2+} Channels in Physiology and Disease. GEOFFREY PITT, Weill Cornell Medicine, New York, NY 10065

Calcium channels modulate the final signal transduction step in excitable cells such as neurons, hormone-secreting cells, and cardiac myocytes and thus are central for their function. The Ca\textsubscript{v}1.2 L-type Ca\textsuperscript{2+} channel encoded by Cacna1c is subject to temporally regulated and tissue-specific alternative splicing. We have generated several new models including gain-of-function and loss-of-function Ca\textsubscript{v}1.2 mouse models, some of which affect only one of two alternatively spliced and mutually exclusive exons. These various models display tissue-specific phenotypes that reflect the alternative splicing. The phenotypes include various behavioral abnormalities that mimic certain neuropsychiatric diseases, cardiac arrhythmias, developmental abnormalities, and disorders of hormone regulation. Not only do they reveal novel and unexpected roles for Ca\textsubscript{v}1.2 in physiology and disease, but they also uncover exon-specific functions that demonstrate the complex regulation of channel splicing and the exon-specific contributions to disease.

70. Single Nucleotide Polymorphisms (SNPs) Alter Current from Human BK Channels. AMBER E. PLANTE, BETH A. MCNALLY, MICHAEL H. LAI, and ANDREA L. MEREDITH, University of Maryland School of Medicine, Baltimore, MD 21201

One mechanism with the potential to alter channel properties is natural genetic variation via nonsynonymous single nucleotide polymorphisms (SNPs). We identified six SNPs (A138V, C495G, N599D, R800W, R640Q, and R645Q) in the large-conductance, Ca\textsuperscript{2+}-activated potassium (BK) channel gene (Kcnma1) predicted to alter channel structure and function. To determine whether these SNPs alter BK currents, we introduced each SNP into a human muscle-type (BK\textsubscript{OMM}) and a brain-type (BK\textsubscript{VR}) BK splice variant. Functional properties were investigated in HEK293T cells using standard voltage-clamp protocols. C495G and R800W affected the conductance-voltage relationship across multiple Ca\textsuperscript{2+} conditions—both BK\textsubscript{OMM} and BK\textsubscript{VR}. C495G and R800W consistentlyproduced leftward and rightward shifts in the V\textsubscript{1/2} values, respectively, that were similar across both splice variant backgrounds. Expressing C495G/R800W in parallel produced a right-shifted effect similar to R800W, but unlike R800W alone, this rightward shift was reduced at 10 μM Ca\textsuperscript{2+}. In contrast to C495G and R800W, two additional SNPs, A138V (an autism-linked SNP) and N599D, had variable effects on current properties. A138V and N599D produced rightward shifts in the V\textsubscript{1/2} in 0 Ca\textsuperscript{2+}, but N599D also produced a leftward shift at 10 μM Ca\textsuperscript{2+}. Next, to test whether SNP effects persist with posttranslational modifications, we treated patches with alkaline phosphatase, which resulted in increased BK channel activity attributed to dephosphorylation of Serine 642 in these conditions. We found the leftward shift due to C495G and rightward shift due to R800W persisted in dephosphorylating conditions. A combination of two SNPs (R640Q/R645Q) predicted to disrupt phosphorylation of Serine 642 produced partial effects on V\textsubscript{1/2} values at intermediate Ca\textsuperscript{2+} concentrations. These results implicate naturally occurring human genetic variation as a potential modulator of BK channel properties across splice variant backgrounds. Furthermore, SNPs exert effects on the conductance-voltage relationship of the channel that are additive with each other and with posttranslational modifications.
71. Role of Chloride Intracellular Channels (CLICs) Proteins in Maintaining Cardiac Mitochondrial Physiology. DEVASENA PONNALAGU, SHUBHA GURURAJA RAO, AHMED TAFAIRUL HUSSAIN, and HARPREET SINGH, Department of Pharmacology and Physiology, Drexel University College of Medicine, Philadelphia, PA 19102

Chloride (Cl) intracellular channels (CLICs) are a unique class of ion channel proteins existing in both soluble and integral membrane form. CLICs are widely conserved in prokaryotes as well as eukaryotes. Six paralogs of CLICs (CLIC1-CLIC6) are known in mammals. They are multifunctional proteins maintaining renal physiology, apoptosis, and tumorigenesis. In addition, using pharmacological inhibitor (IAA-94), CLICs have been implicated in ablation of myocardial infarction (MI) due to ischemia-reperfusion (IR) injury, but the molecular identity and mechanism of cardioprotection was not elucidated. This intrigued us toward understanding the significance of each CLIC in maintaining cardiac physiology and their respective role in cardioprotection from IR injury. In this study, we show that CLIC1, CLIC4, and CLIC5 are the predominant paralogs present in the cardiac tissue. CLIC4 and CLIC5 localize to the mitochondria of adult cardiomyocytes as well as Percoll-purified mitochondria, but not CLIC1, indicating CLIC4 and CLIC5 as mitochondrial channel proteins. We also observe unique distribution of CLIC4 and CLIC5 in mitochondria as CLIC4 is enriched in outer mitochondrial membrane and CLIC5 in inner mitochondrial membrane (IMM). These results for the first time established the molecular component of IMM Cl channel. Functionally, we have observed that CLIC5 knockout (KO) mice show significant increase in ROS production (P ≤ 0.05, n = 3) as compared with wild-type mice, whereas CLIC1 and CLIC4 mice did not show any significant difference in the ROS production. Interestingly, only CLIC4 KO mice showed a difference (~10%, n = 5) in the cardiac mitochondrial calcium retention capacity as compared with wild-type mice. These results strongly establish CLIC4 and CLIC5 as mitochondrial ion channels and also their significance in maintaining cardiac mitochondrial function. As CLICs are implied to play a role in cardioprotection from IR injury, role of CLIC4 and CLIC5 in cardioprotection needs to be further investigated.

72. Regulation of Neurogenesis and Astrocyte Function by Store-Operated Calcium Channels. MURALI PRAKRIYA, Northwestern University, Chicago, IL 60611

Store-operated calcium channels (SOCs) are a major pathway for calcium signaling in virtually all animal cells and serve a wide variety of functions ranging from gene expression, motility and secretion to tissue and organ development and the immune response. SOCs are activated by the depletion of Ca^{2+} from the endoplasmic reticulum (ER), triggered physiologically through stimulation of a diverse set of metabotropic surface receptors. The identification of the STIM proteins as ER Ca^{2+} sensors and the Orai proteins as store-operated channels 10 yr ago has enabled rapid progress in understanding the mechanisms of SOC function and their physiological roles in various tissues. In this talk, I will discuss our recent work in using genetically modified mice for probing the physiological roles of SOCs in regulating neurogenesis and gliotransmitter release from astrocytes.

73. HIV-Associated Cardiovascular Disease: Role of Connexin43. LISA PREVEDEL,1,2 CAMILLA MOROCHO,1 MICHAEL V. BENNETT,3,8 SUSAN MORGELLO,4,5 and ELISEO A. EUGENIN,1,2 1Public Health Research Institute (PHRI), Newark, NJ 07103; 2Department of Microbiology, Biochemistry, and Molecular Genetics, Rutgers New Jersey Medical School, Rutgers the State University of NJ, Newark, NJ 07103; 3Department of Neuroscience, Albert Einstein College of Medicine, Bronx, NY 10461; and 4Department of Pathology and Neuroscience, Icahn School of Medicine at Mount Sinai, New York, NY 10029; *presenting person

Currently, HIV-infected individuals are living longer due to effective antiretroviral treatment (ART). Despite block of viral replication, several non-AIDS comorbidities, including heart disease, have become a major cause of death in the HIV-infected population. However, the mechanisms involved are poorly described. Here we report changes in the expression and distribution of Connexin43 (Cx43) in vast areas of the heart in correlation with loss of cardiac function using postmortem human heart tissue obtained from HIV-infected and uninfected individuals. Cx43 is the main component of gap junctions in cardiomyocytes at the intercalated disks and mediates propagation of the cardiac action potential. We found that Cx43 expression is increased in HIV-infected individuals and its localization is altered, which may contribute to the increased heart disease. Cx43 expression was examined by qRT-PCR and Western blot, and protein distribution was evaluated by immunolabeling and confocal microscopy. Van Kossa staining for calcium showed increased calcium deposition in areas with compromised Cx43 expression and localization in association with altered structure evaluated by electron microscopy. Cx43 expression was up-regulated at both mRNA and protein levels and Cx43 was mislocalized to the lateral membranes of the cardiomyocytes in addition to its normal presence at the intercalated disks. Regions of the tissue with mislocalized Cx43 also exhibited calcium overload, mitochondrial degeneration/proliferation, and compromise of myofibrils. All of these changes were independent of viral replication, CD4 counts, inflammation, and previous or ongoing ART. Our results demonstrate up-regulation of Cx43 in the hearts of HIV individuals. Furthermore, these areas also
have calcium and mitochondrial dysregulation as well as myofibril compromise, despite successful block of HIV replication. We propose that persistent viral DNA expresses HIV-tat protein, which alters Cx43 expression and distribution, leading to chronic heart damage and increased sensitivity to additional stress.

74. Electrical Remodeling in the Sinoatrial Node Associated with Diabetes and Aging. CATHARINE PROENZA, University of Colorado School of Medicine, Aurora, CO 80045

Cardiac pacemaking is driven by the generation of spontaneous action potentials (APs) by specialized pacemaker myocytes in the sinoatrial node of the heart. Sinoatrial node dysfunction limits cardiac output, VO$_2$, and functional independence for older people and for people suffering from diabetes. Development of new treatments for sinoatrial node dysfunction requires understanding of the underlying molecular mechanisms that are altered by aging or disease. To this end, mice are an established model system for both aging and diabetes, the ion channels that control pacemaking are highly conserved in mice and humans, and we have found that aging and diabetes slow maximum and intrinsic heart rates in mice as they do in humans. Slower heart rates in both aged and diabetic mice resulted from slower spontaneous AP firing rates in isolated sinoatrial node myocytes (SAMs). However, we have found that aging and diabetes are associated with different changes in AP waveform parameters and membrane currents. Aging slows sinoatrial myocyte AP firing rate mainly via hyperpolarization of the maximum diastolic potential and slowing of the spontaneous diastolic depolarization rate. In contrast, diabetes slows AP firing rate mainly by increasing AP duration. In accordance, we have found that aging is associated with a hyperpolarizing shift in the voltage dependence of the funny current (I$_f$), whereas diabetes is associated with a reduction in the transient outward current (I$_{to}$) in sinoatrial myocytes. Thus, I$_f$ and I$_{to}$ represent distinct potential pharmacological targets for treatment of sinoatrial node dysfunction in aging or diabetes, respectively.

75. Intracellular Fibroblast Growth Factor 14 Has Differential Effects on the Intrinsic Excitability of Hippocampal Pyramidal Neurons and Cerebellar Purkinje Neurons. JOSEPH L. RANSDELL,1 YARIMAR CARRASQUILLO,2,* MARIE K. BOSCH,2 DAVID M. ORNITZ,2 and JEANNE M. NERBONNE,1,2 1Department of Medicine and 2Department of Developmental Biology, Washington University School of Medicine, St. Louis, MO 63110; *Present address: Division of Intramural Research, National Center for Complementary and Integrative Health (NCCIH), Bethesda, MD 20892

Intracellular fibroblast growth factors (iFGF11–14) are a unique subgroup of FGF proteins that are functionally distinct from canonical FGFs. iFGFs function intracellularly and are known to bind to the pore-forming α (α) subunits of voltage-gated sodium (Nav) channels, although recent reports suggest that iFGFs may interact with additional types of voltage-gated ion channels. iFGF14 is expressed throughout the CNS and is the locus of mutations responsible for spinal cerebellar ataxia 27 (SCA27), an autosomal-dominant disorder characterized by ataxia and cognitive decline. Mice harboring a targeted disruption in the Fgf14 locus (Fgf14$^{-/-}$) recapitulate many of the phenotypes seen in SCA27. Whole-cell recordings from Fgf14$^{-/-}$ hippocampal CA1 pyramidal neurons in acute slices revealed that the loss of iFGF14 does not affect evoked firing frequency. Analysis of phase plots, however, revealed a hyperpolarizing shift in the voltage dependence (dV/dt) of the rising phase of the action potential. Voltage-clamp experiments demonstrated that the voltage dependence of activation of the Nav conductance was significantly (P < 0.01) hyperpolarized in Fgf14$^{-/-}$ pyramidal neurons (V$_{1/2}$ = −39 mV), compared with WT (V$_{1/2}$ = −30 mV) pyramidal neurons. These results contrast with the effects of iFGF14 in cerebellar Purkinje neurons, in which loss of iFGF14 caused a dramatic reduction in spontaneous firing, and the voltage dependence of steady-state inactivation of the Nav currents was significantly (P < 0.001) hyperpolarized (V$_{1/2}$ = −73 mV), compared with WT (V$_{1/2}$ = −55 mV) Purkinje neurons. Importantly, immunohistochemical experiments revealed that the loss of iFGF14 does not affect Nav subunit localization at the axon initial segments in either hippocampal CA1 pyramidal or cerebellar Purkinje neurons. Taken together, these results demonstrate that expression of iFGF14 has cell type–specific effects: iFGF14 regulates the voltage dependence of activation of the Nav currents in hippocampal CA1 pyramidal neurons and regulates the voltage dependence of steady-state inactivation of the Nav currents in cerebellar Purkinje neurons.

76. Fast Voltage-Gated Sodium Channel Activity in Huntington’s Disease. ERIC J. REED,1 MARK M. RICH,2 and ANDREW A. VOSS,1 1Department of Biological Sciences and 2Department of Neuroscience, Cell Biology, and Physiology, Wright State University, Dayton, OH 45435

Huntington’s disease is caused by expanded CAG repeats in the huntingtin gene and results in cognitive problems as well as muscle weakness, chorea, rigidity, and dystonia. Most of the research on Huntington’s disease has focused on neurodegeneration. Recent studies have begun to find peripheral defects that may help explain the debilitating motor symptoms of the disease. For example, we have shown that skeletal muscle from transgenic R6/2 Huntington’s disease mice is hyperexcitable because of decreases in the resting chloride and potassium currents (Waters et al. 2013. Proc. Natl. Acad.
Sudden cardiac death (SCD) is a major cause of mortality, affecting more than a quarter million people annually in the United States. Although there are a number of causes for SCD, inherited genetic mutations account for a substantial proportion of deaths in victims under the age of 40 yr. Rare forms of inheritable cardiac disease underlying SCD include Long-QT3 (LQT3) and Brugada syndromes (BrS1), both of which arise as a consequence of channelopathies caused by mutations in the SCN5a gene that encodes the cardiac voltage-gated sodium channel, NaV1.5. One NaV1.5 mutant, E1784K, is among several that cause a mixed channelopathy, having the characteristics of both LQT3 and BrS1. We studied E1784K channels expressed in Xenopus oocytes and mammalian cells and found that several physiological factors, all of which are normal by-products of intense exercise, exacerbate the biophysical defects caused by the mutation itself. Voltage clamp recordings of ion and gating currents show changes in a range of biophysical properties in E1784K, compared with wild-type NaV1.5 channels, when temperature is raised, extracellular pH is reduced, or cytosolic Ca$^{2+}$ is elevated. These biophysical changes are predicted to be arrhythmogenic. We incorporated our biophysical results into a ventricular action potential model and found that, at high heart rates, the effects of temperature, pH, or calcium are individually arrhythmogenic. These results lead us to the conclusion that catastrophic arrhythmias may be triggered by intense exercise in individuals carrying the SCN5a mutation underlying the E1784K form of mixed syndrome.

78. CRISPR/Cas-Mediated Genome Engineering.

ROB TAFT, The Jackson Laboratory, Bar Harbor, ME

CRISPR (clustered regularly interspaced short palindromic repeats) is a powerful genetic tool, giving scientists unprecedented precision to alter genomic DNA—from generating transgens to correcting mutations. The main CRISPR system includes an enzyme, Cas9, that cuts DNA. This system is programmable and can be directed to precise sequences in the genome via an RNA-based guide molecule. This genome-editing technology has developed rapidly and has already been applied to a range of biological systems and disease areas since the first work in mouse and human cells less than three years ago. The power of CRISPR techniques stem not only from their precision, but also from their ease of use, most notably by significantly reducing the time to produce transgenic alleles. The promises of precise, combinatorial, efficient methods for genomic engineering are exciting, and applying CRISPR technology to a variety of model organisms is expected to quickly advance scientific understanding of disease mechanisms by allowing researchers to ask complex questions and find answers much faster than with traditional gene targeting approaches. This talk will cover how CRISPR/Cas has been used successfully in mice to generate endogenous knock-in alleles, conditional (“floxed”) mutations, as well as multiple mutations in a single generation. The discussion will focus on how this technology compares with similar technologies and technical challenges and practical considerations.

79. Selective Antagonists of Na$_{v}$1.6 Prevent Electrically Induced Seizures in a Mouse Model of EIEE13.

PARISA TARI, CELINE DUBE, KULDIP KHAKH, ELAINE CHANG, NOAH SHUAT, CLINT YOUNG, SOPHIA LIN, ZHIWEI XIE, RICHARD DEAN, ANDREA LINDGREEN, LUIS SOJO, ABID HASAN, WEI GONG, MICHAEL GRIMWOOD, THILO FOCKEN, CHARLES J. COHEN, and J.P. JOHNSON JR., Xenon Pharmaceuticals, Burnaby, BC V5G 4W8, Canada

CNS neurons express three voltage-gated sodium channels, Na$_{v}$1.1, Na$_{v}$1.2, and Na$_{v}$1.6. Na$_{v}$1.1 is the dominant isoform in inhibitory interneurons, and block of Na$_{v}$1.1 is believed to be proconvulsant since patients with heterozygous null mutations in the SCN1A gene encoding Na$_{v}$1.1 have early infantile onset epileptic encephalopathy 6 (EIEE6, a.k.a. Dravet syndrome). Na$_{v}$1.2 and Na$_{v}$1.6 are expressed in excitatory neurons, and mutations in the SCN8A gene encoding Na$_{v}$1.6 cause another catastrophic epilepsy, EIEE13. EIEE13 patients seize as early as the first day postpartum. Patients exhibit
cognitive and physical delay including ataxia and muscular hypotonia. Many patients never gain the ability to speak or walk. Most EIEE13 mutations identified are gain of function, so selective inhibition of Na\(_{1.6}\) is a promising interventional approach. Consistent with this idea, some EIEE13 patients respond to nonselective Na\(_{V}\) inhibitors, such as phenytoin. Inhibiting Na\(_{1.6}\), while sparing Na\(_{1.1}\), and thus inhibitory interneuron function, should be more effective and better tolerated. Recombinant mice with a patient-identified mutation (N1768D) display symptoms of epilepsy (Wagnon et al. 2015, H. Mol. Genet. 24:506–515). Mice have reduced seizure thresholds, spontaneous seizures, and sudden death. We evaluated the ability of Na\(_{1.6}\) inhibitors to prevent seizures in a 6-Hz psychomotor seizure assay in N1768D mice and found them effective. Drugs with little selectivity for Na\(_{1.6}\), novel compounds that block Na\(_{1.6}\) and Na\(_{1.2}\), but spare Na\(_{1.1}\), and novel Na\(_{1.6}\) selective compounds all prevented seizures in N1768D mice. In vitro inhibition of Na\(_{1.6}\) combined with brain exposure was predictive of in vivo efficacy. Block of additional sodium channels by less selective compounds did not impact efficacy, suggesting the assay is a good surrogate for evaluating Na\(_{1.6}\) inhibition and occupancy. Thus, directly addressing the underlying mechanism of disease by selective block of Na\(_{1.6}\) can improve seizure resistance in this EIEE13 model.

Sponsor: Charles Cohen

80. The Cardiac Sodium Channel Nav1.5 Occurs at Three Distinct Pools in Cardiomyocytes. SARAH VERMIJ, DIANA SHY, JEAN-SÉBASTIEN ROUGIER, and HUGUES ABRIEL, Department of Clinical Research, University of Bern, Bern, Switzerland

The voltage-gated cardiac sodium channel Nav1.5 is responsible for the rapid upstroke of the cardiac action potential. Mutations in its gene SCN5A are associated with many severe cardiac disorders, including Brugada and congenital long-QT syndrome. The phenotypic diversity may be explained by the suspected three pools of Nav1.5: at the intercalated disc (ID), in the lateral membrane (LM), and in the T-tubular membrane (TTM). These pools seem to be differentially regulated: mice that lack Na\(_{1.6}\) at the LM, leading to a reduction in sodium current. This project aims to correlate Nav1.5 to markers of the three pools in isolated cardiomyocytes by immunofluorescent stainings and the proximity ligation assay (PLA) DuoLink.

In isolated cardiomyocytes, we showed that Nav1.5 co-localized with the ID makers connexin43 (Cx43) and plakophilin-2 (PKP2) at the ID and with syntrophin and dystrophin at the LM. PLA experiments confirm that Nav1.5 is within close proximity (<40 nm) with Cx43, PKP2, and syntrophin. Stainings with the TTM marker Bin1 and Nav1.5 showed a clear overlap in fluorescence profiles. The distance between fluorescence peaks is ~2 μm, specific for T-tubules. Interestingly, preliminary data suggest that the overlap between Bin1 and Nav1.5 is stronger in ASIV than in wild-type mice.

To conclude, our data strongly suggest that Nav1.5 is present in three pools in cardiac cells: at the ID, the LM, and the TTM. Moreover, we showed that each pool is differentially regulated by specific interacting proteins: PKP2 and Cx43 at the ID and syntrophin and dystrophin at the LM. Protein interactions at the TTM remain unknown. Next, we will use super-resolution microscopy to investigate Nav1.5 clusters at the LM and the TTM and localization of interacting proteins.

81. Gain-of-Function Mutations of BK Potassium Channels and Pro-Excitatory Effects. BIN WANG, VLADISLAV BUGAY, LING LING, HUI-HSUI CHUANG, ADELINE ORTS-DEI, IMMMAGINE, JASON PUGH, DAVID B. JAFFE, and ROBERT BRENNER, University of Texas Health Sciences Center, San Antonio, TX 78229

While ion channel drugs have long been the mainstay of epilepsy therapies, genetics sometimes reveal paradoxical effects of ion channels that caution simplistic interpretation and drug targeting. Large conductance calcium- and voltage-activated potassium (BK) channels present a key example where a potassium channel gain-of-function mutation can be pro-excitatory, and blockers of BK channels protect against seizures. Using a number of BK channel gain-of-function mouse models, we have attempted to understand the mechanisms underlying pro-excitatory effects of BK channels in dentate gyrus neurons of the hippocampus. We indeed observed electrographic seizures in a mouse model of the human D434G (D369G in mouse), BK gain-of-function epilepsy mutation, and also in genetic knockout of the inhibitory β4 subunit. Pharmacological studies in the β4 knockout suggest that pro-excitatory effects of BK channels are dependent on spike-evoked ryanodine receptor calcium release. Ryanodine receptors were found to feed calcium to fast-gated BK channels, which in-turn increase the fast-afterhyperpolarization (fAHP) and shorten the interspike interval. The pro-excitatory ryanodine receptor, BK channel functional coupling was also revealed in the epileptic gain-of-function R2474S ryanodine receptor mutation mice. R2474S mice were found to have a larger BK-dependent fAHP and increased spike frequency. These findings indicate that fast-gated BK channels promote excitability in excitatory neurons by increasing the fAHP amplitude, which in turn shortens the time to the subsequent spike. Using dynamic clamp to systematically alter the fAHP amplitude during spiking, we confirmed the cause and effect between fAHP amplitude and interspike interval. In summary, selective functional coupling of fast-gated BK channels with ryanodine receptors suggest that BK channels may serve as effectors of endoplasmic reticulum calcium dysregulation in epilepsy and other diseases.
82. Limb-Girdle Muscular Dystrophy 2L Is Caused by a Defect in Phospholipid Scrambling Mediated by ANO5. JARRED M. WHITLOCK, KUAIYU, TUANYUAN CUI, and H. CRISS HARTZELL, Department of Cell Biology, Emory University School of Medicine, Atlanta, GA 30322

Limb girdle muscular dystrophies (LGMDs) are the fourth most common inherited muscle disorder. Most LGMDs are caused by recessive mutations in one of 23 different genes. LGMD2L, one of the most common LGMDs, usually presents after the third decade and is characterized by asymmetric lower limb weakness/atrophy, elevated serum creatine kinase levels, and exercise intolerance. LGMD2L is linked to mutations in ANO5/TMEM16E, a member of the Anoctamin/TMEM16 superfamily of ion channels and regulators of Ca\(^{2+}\)-dependent phospholipid scrambling (Ca\(^{2+}\)-PLS). We propose that ANO5 regulates Ca\(^{2+}\)-PLS that plays a role in muscle repair. PLS is a process where phosphatidylserine, a major signaling molecule molecule normally sequestered in the inner leaflet of the plasma membrane, is translocated to the external leaflet and exposed to the extracellular space. The fusion of myoblasts with one another and with muscle fibers plays a major role in repair of muscle fibers that have been damaged by mechanical stress during activity. Phosphatidylserine exposure occurs coincidently with myoblast fusion and participates in this process. Here we demonstrate that ANO5 expression elicits Ca\(^{2+}\)-PLS and a nonselective ionic current associated with lipid translocation. An Ano5\(^{-/-}\) knockout mouse recapitulates many features of human LGMD2L, in particular defective muscle repair. Myoblasts isolated from this mouse exhibited diminished myoblast fusion in vitro and do not exhibit Ca\(^{2+}\)-PLS or Ano5 ionic currents, while wild-type myoblasts exhibit both activities. To determine whether Ca\(^{2+}\)-PLS is disrupted in human LGMD2L, skin fibroblasts were obtained from an LGMD2L patient and unaffected controls. Sequencing demonstrated ANO5 mutations in both ANO5 alleles of the patient. Western blot showed greatly diminished ANO5 expression. The LGMD2L patient fibroblasts exhibited markedly reduced Ca\(^{2+}\)-PLS, with >75% reduction in the number of cells expressing phosphatidylserine. Our findings demonstrate that ANO5 is essential for proper Ca\(^{2+}\)-PLS signaling and that this process is perturbed in patients with LGMD2L. We propose that defective Ca\(^{2+}\)-PLS reduces the ability of myoblasts to fuse during the process of muscle repair.

83. BK Channels Are Activated by Distinct Calcium Sources During Day and Night in SCN Neurons. JOSHUA P. WHITT and ANDREA L. MEREDITH, University of Maryland School of Medicine, Baltimore, MD 21201

BK K\(^{+}\) channels are regulated by membrane depolarization and increases in local intracellular Ca\(^{2+}\) concentration ([Ca\(^{2+}\)]\(_{i}\)). BK channels can couple to the voltage-gated Ca\(^{2+}\) channels L, P, Q, and N-type, as well as to release from intracellular Ca\(^{2+}\) stores. In the suprachiasmatic nucleus (SCN), both [Ca\(^{2+}\)] and voltage-gated Ca\(^{2+}\) channel (VGCC) currents are greater during the day. To determine whether diurnal regulation of [Ca\(^{2+}\)] was relevant for BK channel activation in SCN, we used a pharmacological approach to identify the Ca\(^{2+}\) sources for BK current activation during the day and night in acute SCN slices. We found a 78% reduction in BK current with 10 μM nimodipine during the day, with less effect at night (10% reduction), suggesting that L-type VGCCs are the primary Ca\(^{2+}\) source for BK activation during the day. Furthermore, 71% of neurons in the SCN express inactivating BK currents (BK\(^{\text{a}}\)) during the day. Nimodipine abolished all the BK currents in the SCN, while 5 μM (S-) Bay-K 8644, an L-type VGCC current activator, produced 100% BK currents during the day. In contrast, at night, when nimodipine has little effect, we found a significant decrease in BK current using 10 μM dantrolene, blocking Ca\(^{2+}\) release from RyRs (63%), as well as 5 μM thapsigargin, depleting Ca\(^{2+}\) from intracellular stores (64%), suggesting that nighttime BK activation is driven by RyR-mediated store release. Lastly, 3 μM ω-conotoxin MVIIC reduced currents to a similar extent between day and night (24% and 22%, respectively), suggesting that a fraction of BK channels maintain stable Ca\(^{2+}\) channel coupling. These data demonstrate diurnal regulation of the functional coupling between BK channels and LTCCs and suggest diurnal changes in Ca\(^{2+}\) coupling could contribute to BK current inactivation in the SCN and the role of BK channel inactivation in circadian rhythmicity.

84. High-Fat Diet Improves Blood Glucose Control in a Mouse Model of Human Neonatal Diabetes: Protection by Fat? ZIHAN YAN, ALECIA WELSCHER, and MARIA S. REMEDI, Department of Medicine, Division of Endocrinology, Metabolism, and Lipid Research and Department of Cell Biology and Physiology, Washington University in St. Louis, St. Louis, MO 63130

Chronic hyperglycemia has been proposed to cause pancreatic β-cell dysfunction and reduction of β-cell mass (glucotoxicity) in diabetes. Gain-of-function (GOF) mutations in the K\(_{\text{ATP}}\) channel have been identified as causal of human neonatal diabetes (NDM). Mice expressing K\(_{\text{ATP}}\)-GOF channels in β-cells demonstrate severe diabetes and low circulating plasma insulin levels due to lack of insulin secretion. As disease progresses, insulin content and β-cell mass dramatically decrease, all secondary consequences of glucotoxicity, also seen in other forms of diabetes. However, comorbidity of hyperglycemia and elevated lipids (glucolipotoxicity) is also critical in diabetes. To test the contribution of elevated lipids in NDM, K\(_{\text{ATP}}\)-GOF mice were exposed to a high-fat diet (HFD) at disease onset. Surprisingly, HFD fed K\(_{\text{ATP}}\)-GOF mice demonstrate resistance to diet-induced obesity accompanied by a markedly divergent
phenotype: (1) remission of diabetes with normalization of fast and fed blood glucose (remitting) and (2) development of severe diabetes (nonremitting). Strikingly, the dramatic secondary loss of insulin content and β-cell mass observed in chow diet–fed K<sub>ATP</sub>-GOF is avoided in HFD-fed mice. Unexpectedly, K<sub>ATP</sub>-GOF remitting mice demonstrated a marked increase in peripheral insulin sensitivity, which precedes the normalization of blood glucose and may explain the lower circulating glucose levels. Importantly, remitting mice show reduced plasma lipids, decreased food intake, and increased energy expenditure and oxygen consumption. These results suggest that restriction of dietary carbohydrates and caloric replacement by fat may induce metabolic changes that are beneficial in terms of decreasing glucotoxicity in K<sub>ATP</sub>-induced diabetes. Dietary recommendations for blood glucose control and their influences in development and progression of diabetes remain controversial. A recent report demonstrated that high-fat dairy products reduce the risk of development of type-2 diabetes. Together, our results suggest that HFD might also be protective in certain forms of diabetes, especially if this is not accompanied by weight gain and obesity.

85. Quantitative Imaging of Genetically-Encoded Metabolic Sensors in Mouse Brain. GARY YELLEN, Harvard Medical School, Boston, MA 02115

Cellular metabolism is no mere “housekeeping” function: it responds dynamically to energy demand and to changes in fuel supply, with important consequences for cellular function. To understand the dynamics of metabolism in living cells, we have developed several genetically encoded fluorescent biosensors for key metabolites—ATP and NADH—and are imaging these in neurons and astrocytes of mice, both in acute brain slices and in vivo. We have observed marked metabolic changes in response to neuronal stimulation, both at naturalistic levels and at pathophysiological levels such as those observed in epilepsy models. We are also investigating a mouse model of metabolic resistance to epileptic seizures, in which blunted glucose metabolism produces substantial resistance to induced seizures.

86. Cardiovascular Phenotypes of a Mouse Model of Cantu Syndrome. HAIXIA ZHANG,1,2 PAIGE COOPER,1,2 CHRISTOPHER EMFINGER,1,2 THERESA HARTER,1,2 MONICA SALA-RABANAL,1,2 SCOT MATROVICH,3 BLANCHE SCHWAPPACH,4 CONOR MCCLENAUGHAN,1,2 ZIHAN YAN,4 ROBERT MECHAM,1 MARIA REMEDI,1 and COLIN G. NICHOLS,1,2 1Department of Cell Biology and Physiology, 2Center for the Investigation of Membrane Excitability Diseases, 3Department of Internal Medicine, and 4Department of Endocrinology, Washington University School of Medicine, St. Louis, MO 63110; and 5Department of Molecular Biology, University Medicine Göttingen, Göttingen, Germany

Cantu syndrome (CS) is a rare disorder characterized by congenital hypertrichosis, distinctive facial features, and cardiovascular defects; the genetic causes of CS are gain-of-function mutations (GOF) of K<sub>ATP</sub> channels. To understand the underlying mechanisms for CS, we used CRISPR/Cas9 genome engineering to generate CS mice carrying the SUR2[R1150Q] mutation (equivalent to human R1154Q—the most frequent mutation in CS patients). CS mice show lower blood pressure (BP), enlarged hearts, and enhanced heart output, which recapitulate cardiovascular phenotypes of CS patients. As expected for a GOF mutation, K<sub>ATP</sub> channels of CS mouse ventricular myocytes show decreased ATP inhibition and increased MgADP stimulation. Interestingly, pinacidil (a SUR2 specific K<sub>ATP</sub> opener) fails to lower BP in CS mice, while it significantly lowers BP in WT mice. Consistently, pinacidil activated current is much smaller in CS cardiomyocytes than in WT cardiomyocytes. However, pinacidil activates SUR2[R1150Q]/Kir6.2 recombinant channels as well as SUR2/Kir6.2 channels in Cosm6 cells, suggesting decreased sensitivity to pinacidil in CS mice is not directly derived from the SUR2[R1150Q] mutation itself. Surprisingly, cDNA sequencing from the hearts of CS mice reveals a deletion of 93 bases following R1150Q in one sequence of SUR2 cDNA, which may imply an alternative splicing event following R1150Q. In addition, SUR2A is down-regulated in CS mouse ventricles with a compensatory up-regulation in SUR1 (a more active subunit in K<sub>ATP</sub> channels), which explains the decreased sensitivity to pinacidil in CS mouse. In conclusion, the SUR2[R1150Q] CS mouse recapitulates the cardiovascular phenotypes of CS patients, and provides a promising mouse model to study the mechanisms and treatment of CS. SUR2[R1150Q] GOF and SUR1 up-regulation may both contribute to the phenotypes in CS mouse. Future studies are required to investigate the phenotypes other than cardiovascular abnormalities in CS mice, as well as the splicing events caused by SUR2[R1150Q].

87. SWELL1 Is a Regulator of Adipocyte Insulin Signaling and Glucose Homeostasis. YANHUI ZHANG,1 LITAO XIE,1 SUSHEEL GUNASEKAR,1 DAN TONG,1 ANIL MISHRA,1 WILLIAM J. GIBSON,2 CHUANSONG WANG,3 TREVOR FIDLER,1 BRODIE MARTHALER,1 ALOYSIUS KLINGELHUTZ,3 E. DALE ABEL,4 JESSICA SMITH,6 ISAAC SAMUEL,5 LEI CAO,3 and RAJAN SAH,1,4 1Department of Internal Medicine, Division of Cardiovascular Medicine, University of Iowa, Carver College of Medicine, Iowa City, IA, 52242; 2Harvard Medical School, Boston, MA, 02115; 3Department of Molecular Virology, Immunology and Medical Genetics, The Ohio State University, Columbus, OH 43210; 4Fraternal Order of the Eagles Diabetes Research Center, Iowa City, IA 52242; and 5Department of Microbiology and 6Department of Surgery, University of Iowa, Carver College of Medicine, Iowa City, IA 52242
Obesity is characterized by a tremendous increase in adipose tissue that is in large part due to massive volumetric expansion of the constituent adipocytes (Farnier et al. 2003. Int. J. Obes. Relat. Metab. Disord. 27:1178–1186). There is a longstanding concept that metabolic disease in obesity is associated more with adipocyte size than number (Farnier et al. 2003. Int. J. Obes. Relat. Metab. Disord. 27:1178–1186; Heinonen et al. 2014. Int. J. Obes. (Lond.). 38:1423–1431; Lonn et al. 2010. FASEB J. 24:326–331; Salans et al. 1968. J. Clin. Invest. 47:153–165; Weyer et al. 2000. Diabetologia. 43:1498–1506). In support of this concept, recent studies have highlighted a connection between adipocyte size and membrane tension with adipocyte signaling and adipogenesis (Ben-Or Frank et al. 344:634–638) and phosphoinositide 3-kinase (PI3K)/Akt phosphorylation (Kumar et al. 2014. J. Exp. Med. 211:929–942). Furthermore, recent studies have uncovered to determine the expression of cardiac sodium channel, and optical mapping studies were performed to determine arrhythmia inducibility in situ and to map conduction velocity and action potential duration.

Results: Overexpression of HuR by AAV injection increased cardiac sodium channel α subunit expression by 119.2% by Western blot. In addition, ectopic HuR also increased endogenous HuR protein level. Our optical mapping study revealed that HuR overexpression reduced arrhythmic inducibility, prevented reentry, increased conduction velocity, and decreased action potential rise time.

Conclusions: Our study results indicate that RNA-binding protein HuR can positively regulate cardiac sodium channel SCN5A expression in vivo and can reduce arrhythmia risk in heart failure. Therefore, manipulation of HuR expression may be a useful strategy to treat HF-associated arrhythmias.

88. Regulation of Cardiac Sodium Channel in Heart Failure by RNA-Binding Protein HuR. ANYU ZHOU,1,2 HONG LIU,1,2 GUANGBIN SHI,1 and SAMUEL C. DUDLEY, JR.1,2 1The Cardiovascular Institute, Rhode Island Hospital, Providence, RI 02903; 2The Warren Alpert Medical School, Brown University, Providence, RI 02903

Introduction: In patients, deletions or loss-of-function mutations of cardiac sodium channel α subunit gene SCN5A have been associated with a wide range of arrhythmias. The expression of SCN5A has been shown to decrease in failing hearts and the reduced expression of SCN5A is associated with the arrhythmia risk in heart failure. mRNA levels are determined by the balance between transcription and mRNA degradation. While transcriptional regulation of SCN5A expression has been extensively studied, little is known regarding the regulation of SCN5A mRNA degradation. We have shown that RNA-binding protein, HuR, positively regulates SCN5A mRNA expression by protecting SCN5A mRNA from degradation in vitro. To examine whether HuR modulates cardiac sodium channel expression in heart failure, we conducted this study.

Methods: A mouse model of myocardial infarction was created by the permanent ligation of the left anterior descending coronary artery. Mouse HuR coding sequences were cloned into an associate adenoviral (AAV) vector driven by cardiac troponin (cTNT) promoter. Recombinant AAV were packaged into infectious serotype 9 AAV particles. AAV-HuR AAV particles were injected into the MI mouse via the right jugular vein 2 wk after left anterior descending coronary artery occlusion. 2 wk after injection, mouse hearts were collected to determine the expression of cardiac sodium channel, and optical mapping studies were performed to determine arrhythmia inducibility in situ and to map conduction velocity and action potential duration.
Methods: The NaV1.5 α subunit contains four domains (DI–DIV), each with six membrane-spanning segments (S1–S6). S1–S4 form the VSDs that convert voltage into channel gating. Previously, we created four DNA constructs, each with a cysteine engineered into the extracellular S4 of a single channel domain. RNA from these constructs was injected into Xenopus oocytes and expressed channels were labeled at this cysteine with TAMRA-MTS fluorophores. Ionic current and fluorescence emission that reflected changes in VSD conformation were simultaneously recorded using the cut-open oocyte configuration.

Results: In comparison with α alone, β1 and β3 cause a depolarizing shift in channel inactivation, and correlating shift in DIV-VSD activation, while β3 also affects DIII-VSD activation. DI and DII were not significantly affected. Class Ib antiarrhythmics, such as lidocaine and ranolazine, modulate DIII-VSD activation while DII and DIII were simultaneously recorded using the cut-open oocyte configuration.

Conclusions: These results suggest that class I drug responses will be significantly altered by the β-subunits, providing a potential mechanism for chamber specificity that depends on β-subunit expression. Inherited mutations carried by the β-subunits are likely to strongly influence the patient response to therapy by these molecules.

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Conclusions: These results suggest that class I drug responses will be significantly altered by the β-subunits, providing a potential mechanism for chamber specificity that depends on β-subunit expression. Inherited mutations carried by the β-subunits are likely to strongly influence the patient response to therapy by these molecules.

Multiple types of cells including human cardiomyocytes exhibit two stable levels of resting potentials in physiologically or pathologically identical tonic conditions. In subphysiological extracellular K⁺ concentrations ([K+]o), which occur in pathological hypokalemia, human cardiomyocytes can show both hyperpolarized and depolarized resting potentials, associating with cardiac arrhythmia. Resting potentials of human cardiomyocytes and cardiomyocytes and cardiac Purkinje fibers can either spontaneously shift from hyperpolarization to depolarization in a subphysiological [K+]o, or fluctuate between two levels in a nongraded manner. Injection of small current pulses can switch resting potentials between the two levels. However, the mechanism underlying this well-known phenomenon is not well understood. We recently demonstrate that in subphysiological [K+]o, K2P1 two-pore domain K⁺ channels dynamically change ion selectivity, become nonselective cation channels, and conduct inward leak cation (mainly Na⁺) currents. Here, we show that under hypokalemia K2P1 channels contribute to two stable levels of resting potentials of cardiomyocytes. We employ two models of cardiomyocytes: human cardiomyocytes derived from induced pluripotent stem cells (hiPSC-CMs) that highly express K2P1 channels and mouse HL-1 cardiomyocytes that do not express K2P1 channels nor exhibit two levels of resting potentials in subphysiological [K+]o. In subphysiological [K+]o, “matured” hiPSC-CMs and mouse HL-1 cardiomyocytes with ectopic expression of K2P1, channels can exhibit two stable levels of resting potentials and N-shaped current-voltage relationships that cross the voltage axis three times, and the first and third zero-current potentials determine the two levels of resting potentials. Removal of K2P1-like inward Na⁺ currents or knockdown of K2P1 expression eliminates two levels of resting potentials in hiPSC-CMs. These results elucidate the mechanism that results in two levels of resting potentials of human cardiomyocytes in hypokalemia and demonstrate a new mechanism regulating excitability.
INDEX OF AUTHORS

Names in bold indicate a speaker summary; the others are submitted abstracts

*Keynote speaker

Abriel, H., 1
Abdelsayed, M., 77
Abel, E.D., 87
Abriel, H., 8, 80
Almazán, E., 68
Alvarez-Miguel, I., 67
Angelini, M., 2
Arnold, W., 63
Asai, Y., 36
Baba, Y., 37
Babich, J., 13
Balse, E., 16
Bankar, G., 37
Baraban, S.C., 30
Bassett, S., 12
Ben-Johny, M., 28
Bennett, M.V., 3, 73
Beuriot, A., 16
Bijvelds, M.J.C., 4
Bonev, A., 25
Bonventre, J.V., 49
Bosch, M.K., 75
Bose, S.J., 4
Bosmans, F., 64
Bot, A.G.M., 4
Brenner, R., 81
Bugay, V., 81
Cai, Z., 4
Cannon, S.C., 5, 35
Cao, L., 87
Carrasquillo, Y., 75
Carver, C.M., 6
Catterall, W., 7
Chahal, N., 37
Chang, E., 37, 79
Chang, J., 37
Chen, H., 90
Chen, K., 90
Chevalier, M., 8
Chowdhury, S., 37
Chuang, H.-H., 81
Chung, D.Y., 9
Cidad, P., 67
Cohen, A., 10
Cohen, C.J., 37, 79
Collier, D., 11, 25
Contreras, J.E., 22
Cooper, P., 55, 86
Cooper, P.E., 61
Coulombe, A., 16
Covarrubias, M., 59
Cui, Y., 82
Dabbertrand, F., 27
Dalziel, J.E., 12
Davis, M.J., 61
De Jonge, H.R., 4
De Lange, W., 48
Dean, R., 79
Deboeuf, K., 20, 66
Decker, S., 37
Dehnhardt, C.M., 37
Dick, I.E., 13
Dilanian, G., 16
Dopico, A.M., 45
Dube, C., 79
Dudley, S.C., Jr., 88
Earley, S., 14
Edwards, J.C., 15
Eichel, C., 16
Emfinger, C., 17, 18, 55, 86
Ethington, E.A., 68
Eugenin, E.A., 3, 73
Farley, J., 19, 20, 66
Fidler, T., 87
Finol-Urdaneta, R.K., 21
Focken, T., 79
Fraidenrach, D., 22
Franck, J., 23
Franklin, B.M., 24
Fraser, K., 12
Freeman, K., 11, 25
French, R.J., 21
Garza-López, E., 52
Gaudet, R., 21
George, A., 26
Gibson, W.J., 87
Gilchrist, J., 64
Gong, W., 79
Gonzales, A.L., 27
Gonzalez, P., 22
Gonzalez-Perez, V., 28
Gouzer, G., 29
Granados-Fuentes, D., 34
Grange, D.K., 61
Griffith, A.J., 65
Grimwood, M., 79
Grone, B.P., 30
Gunasekar, S., 87
Hackos, D., 37
Harraz, O., 25
Harraz, O.F., 31
Harris, A., 4
Harter, T., 55, 86
Hartle, C.M., 32
Hartzell, H.C., 82
Hasan, A., 79
Hatem, S., 16
Hawash, A., 33
He, Y., 68
Hemeon, I., 37
Hermanstyn, T.O., 34
Hernández-Ochoa, E.O., 35
Herzog, E.D., 34
Hoffman, B.U., 36
Holt, J.R., 65
Howard, S., 37
Hughes, E., 43
Hull, J., 38
Hussain, A.T., 71
Hyrc, K., 17
Islam, M.F., 20, 66
Isom, L., 38, 50
Isom, L.L., 39
Jaffe, D.B., 81
Johnson, E.K., 40
Johnson, J.P., 37
Johnson, J.P., Jr., 79
Jones, D.K., 48
Julius, D., 64
Kadakia, K., 54
Kanter, E.M., 40
Karagueuzian, H.S., 2
Kass, R., 41
Katchman, A., 42
Khakh, K., 37, 79
Kissel, J., 63
Klingelhutz, A., 87
Koide, M., 43, 44
Koob, M.D., 68
Kotlikoff, M.I., 27

39a
Kruger, L., 38
Kuntamallappanavar, G., 45
Kurima, K., 65
Lai, M.H., 70
Lee, A., 46
Lee, J.-M., 61
Levin, M.A., 61
Limpitikul, W.B., 13
Lin, S., 37, 79
Lindgreen, A., 79
Lin, S., 37, 79
Lindgren, A., 37
Ling, L., 81
Lingle, C., 47
Lingle, C.J., 28
Liu, F., 48
Liu, H., 88
Liu, J., 4
Liu, L., 49
Liu, X.-P., 65
Liu, Y., 50
Liu, Z., 90
Longden, T.A., 27, 51
López-González, I., 52
Lopez-Lopez, J.R., 67
Lopez-Santiago, L., 50
Louault, F., 16
Lumpkin, E.A., 36
Manuelyan, I., 43
Markwardt, M.L., 53
Martínez-Hernández, E., 68
Marx, S., 42
Matkovich, S., 86
Matsui, M., 54
McArthur, J.R., 21
McClanaghan, C., 55, 86
McFarland, J., 53
McNally, B.A., 56, 70
Mecham, R., 86
Meisler, M., 50
Meredith, A., 57
Meredith, A.L., 56, 70, 83
Mirshahi, T., 32
Mishra, A., 87
Mohler, P.J., 58
Moore, B.S., 32
Morgello, S., 73
Morrocho, C., 73
Muqeem, T., 59
Mutolo, M.J., 4
Nafzger, S., 8
Nelkenbrecher, K., 37
Nelson, M., 11, 25, 60
Nelson, M.T., 27, 31, 43, 51
Nerbonne, J.M., 34, 40, 75
Nichols, C., 17, 18, 55
Nichols, C.G., 61, 86
Nishigaki, T., 52
Noebels, J., 62, *
Novak, K., 63
O’Malley, H., 38
Oka, F., 9
Olcese, R., 2
Ornitz, D.M., 75
Orts-Del Immagine, A., 81
Osborn, J.L., 24
Osteen, J.D., 64
Palmi-Pallag, T., 4
Pan, B., 65
Panchal, J., 66
Pantazis, A., 2
Pappas, A.C., 44
Parent, J., 50
Perez-Garcia, M.T., 67
Perissinotti, P.P., 68
Peters, C.H., 77
Pezhouman, A., 2
Piedras-Renteria, E.S., 68
Pinto, V., 59
Pitt, G., 54, 69
Plante, A.E., 56, 70
Ponnalagu, D., 71
Prakriya, M., 72
Prevedel, L., 73
Proenza, C., 74
Pugh, J., 81
Qin, T., 9
Ramachandran, J., 22
Ransdell, J.L., 75
Rao, S.G., 71
Reed, E.J., 76
Reissaus, C., 17
Remedi, M., 55, 86
Remedi, M.S., 17, 18, 61, 84
Rich, M., 33, 63
Rich, M.M., 76
Rittenhouse, A.R., 49
Rizzo, M.A., 53
Robertson, G.A., 48
Robinette, C.L., 37
Ross, B., 53
Rougier, J.-S., 8, 80
Roy, N.C., 12
Ruben, P.C., 77
Ryan, T., 29
Sackheim, A., 11
Sadeghian, H., 9
Safina, B., 37
Sah, R., 87
Sala-Rabanal, M., 86
Sala-Rabinal, M., 55
Samuel, I., 87
Sanderson, A., 63
Savalli, N., 2
Schneider, M.F., 35
Schubert, A., 89
Schwappach, B., 55, 86
Shah, R., 55
Shapiro, M.S., 6
Shelton, E.L., 61
Sheppard, D.N., 4
Shi, G., 88
Shirokova, N., 22
Shu, B., 79
Shui, B., 27
Shy, D., 80
Silva, J., 89
Sinden, D., 54
Singh, G.K., 61
Singh, H., 71
Smith, J., 87
Sojo, L., 79
Sojo, L.E., 37
Sonkusare, S., 25, 43
Springer, S., 40
Sutherlin, D.P., 37
Syed, A.U., 43
Taft, R., 78
Tari, P., 79
Tidball, A., 50
Tikhonov, D., 21
Tong, D., 87
Vermij, S., 8, 80
Villalba, N., 11, 25
Voss, A., 33, 76
Voss, S.R., 24
Wagnon, J., 50
Wallbrook, M., 37
Wang, B., 81
Wang, C., 87
Wang, W., 40
Wang, Y., 17
Wei, E., 54
<table>
<thead>
<tr>
<th>Name</th>
<th>Pages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weiss, J.N.</td>
<td>2</td>
</tr>
<tr>
<td>Wellman, G.C.</td>
<td>43, 44</td>
</tr>
<tr>
<td>Welscher, A.</td>
<td>17, 18, 84</td>
</tr>
<tr>
<td>Whitlock, J.M.</td>
<td>82</td>
</tr>
<tr>
<td>Whitt, J.P.</td>
<td>83</td>
</tr>
<tr>
<td>Wu, F.F.</td>
<td>35</td>
</tr>
<tr>
<td>Xie, L.</td>
<td>87</td>
</tr>
<tr>
<td>Xie, Z.</td>
<td>37, 79</td>
</tr>
<tr>
<td>Xue, M.</td>
<td>29</td>
</tr>
<tr>
<td>Yamada, K.A.</td>
<td>40</td>
</tr>
<tr>
<td>Yan, Z.</td>
<td>17, 18, 84, 86</td>
</tr>
<tr>
<td>Yang, L.</td>
<td>42</td>
</tr>
<tr>
<td>Yang, W.</td>
<td>13</td>
</tr>
<tr>
<td>Yazawa, M.</td>
<td>13</td>
</tr>
<tr>
<td><strong>Yellen, G.</strong></td>
<td>85</td>
</tr>
<tr>
<td>Young, C.</td>
<td>37, 79</td>
</tr>
<tr>
<td>Young, W.</td>
<td>12</td>
</tr>
<tr>
<td>Yu, K.</td>
<td>82</td>
</tr>
<tr>
<td>Yuan, Y.</td>
<td>50</td>
</tr>
<tr>
<td>Yue, D.T.</td>
<td>13</td>
</tr>
<tr>
<td>Zeglin, A.</td>
<td>68</td>
</tr>
<tr>
<td>Zhang, A.</td>
<td>40</td>
</tr>
<tr>
<td>Zhang, H.</td>
<td>55, 61, 86</td>
</tr>
<tr>
<td>Zhang, J.</td>
<td>53</td>
</tr>
<tr>
<td>Zhang, Y.</td>
<td>87</td>
</tr>
<tr>
<td>Zhorov, B.</td>
<td>21</td>
</tr>
<tr>
<td>Zhou, A.</td>
<td>88</td>
</tr>
<tr>
<td>Zhou, M.</td>
<td>90</td>
</tr>
<tr>
<td>Zhu, W.</td>
<td>89</td>
</tr>
<tr>
<td>Zuo, D.</td>
<td>90</td>
</tr>
</tbody>
</table>
TRPV4 Inhibitor Improves Pulmonary Function and Oxygen Saturation in a Pig Translational Model of Chemically Induced Acute Lung Injury. SATYA ACHANTA,1 BOYI LIU,1 MICHAEL A GENTILE,1 DAVID J BEHM,2 THERESA J ROETHKE,2 MICHAEL D GUNN,1 IRA M CHEIFETZ,1 and SVEN-ERIC JORDT,1
1Duke University School of Medicine, Durham, NC 27703; 2GlaxoSmithKline Pharmaceuticals, King of Prussia, PA 19406

The treatment of chemically induced acute lung injury (ALI) remains challenging because of the lack of mechanism-based therapeutic approaches. Chlorine (Cl2) gas is a severe chemical threat with frequent exposures in domestic and occupational environments and in transportation accidents. Cl2 was repeatedly used as a chemical weapon, first in World War I and most recently in Syria, with hundreds of casualties reported there. Our recent studies in a mouse model of Cl2 exposure have shown that transient receptor potential vanilloid 4 (TRPV4), an ion channel expressed in pulmonary tissues, is a crucial mediator of chemically induced ALI. TRPV4 inhibitors ameliorated Cl2 induced ALI in mouse models (Balakrishna S. et al., 2014). In the present study, we asked whether the beneficial therapeutic effects of TRPV4 inhibition would translate to a large animal model of Cl2 exposure, the pig.

Specific pathogen free Yorkshire pigs were mechanically ventilated under anesthesia and exposed to Cl2 gas (≤ 240 ppm for 1 h). A novel TRPV4 inhibitor, GSK691 or vehicle was administered post-Cl2 exposure. Methacholine airway challenge was done. Total and differential leukocyte cell counts in bronchoalveolar lavage fluid (BALF) were determined. Lung injury markers were characterized by ELISA and quantitative PCR. Histopathological analysis and scoring was done following American Thoracic Society guidelines.

Treatment with GSK691 improved oxygenation and respiratory physiological parameters compared to vehicle group. GSK691 treatment reduced total BALF cell and neutrophil counts. Vascular leakage and edema were decreased in GSK691 treated pigs. Pro-inflammatory cytokine markers were decreased and histopathology was improved by GSK691.

These results suggest that TRPV4 inhibitors are potential new countermeasures for Cl2 induced ALI; and pave the way for further testing of TRPV4 inhibitor drug candidates following the US FDA’s animal rule (21 CFR 314.600) as an efficacious countermeasure to treat Cl2 induced ALI in human patients.

Additional Abstracts

91. TRPV4 Inhibitor Improves Pulmonary Function and Oxygen Saturation in a Pig Translational Model of Chemically Induced Acute Lung Injury. SATYA ACHANTA,1 BOYI LIU,1 MICHAEL A GENTILE,1 DAVID J BEHM,2 THERESA J ROETHKE,2 MICHAEL D GUNN,1 IRA M CHEIFETZ,1 and SVEN-ERIC JORDT,1
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92. The Action of Retigabine on KV7 Channels Activity Requires PI(4,5)P2. AARON CORBIN-LEFTWICH1,2 and CARLOS A. VILLALBA-GALEA,3 1Department of Physiology and Biophysics, Virginia Commonwealth University School of Medicine, Richmond, VA 23298, 2Department of Biology, University of Richmond, Richmond, VA 23173 3University of the Pacific, Stockton, CA 95211

Retigabine is a KV7 channel agonist used to treat hyperexcitability disorders in humans (Rudzinski et al., 2016). Although the mechanisms of action remains unclear, it is thought that Retigabine facilitates M-current activation by shifting the voltage dependence of the heteromeric KV7.2/KV7.3 channel to more negative potentials (Schenzer et al., 2005; Wuttke et al., 2005; Gunthorpe et al., 2012; Kim et al., 2015). This notion, however, has been recently challenged as the pore domain was identified as the drug’s target (Schenzer et al., 2005; Wuttke et al., 2005; Gunthorpe et al., 2012; Kim et al., 2015), suggesting that Retigabine could, instead, stabilize already-activated channels without affecting activation. Supporting this latter idea, we have recently shown that the heteromeric KV7.2/KV7.3 channel has at least two open modes, (OPEN1 and OPEN2), that OPEN2 is more stable than OPEN1, and that Retigabine further stabilizes OPEN2 (Corbin-Leftwich et al., 2016). To further understand this process, we evaluated the role of PI(4,5)P2 in KV7 channels modal behavior. KV7 channels are subject to muscarinic regulation since these channels require PI(4,5)P2 to be active. Thus, pinpointing a potential functional link between Retigabine and PI(4,5)P2 would be essential to comprehend the action of anticon-
vulsants in patients. Using pharmacological and enzymatic approaches, we have found that the stability of the KV7.2/KV7.3 OPEN2 mode exhibits a higher dependence on PI(4,5)P2 than the OPEN1 mode, suggesting that the affinity for phosphoinositides of this channel changes during activation. Per these findings, we hypothesized that the action of Retigabine would depend on the concentration of PI(4,5)P2 since Retigabine targets the OPEN2 mode. Indeed, we found that decreasing the PI(4,5)P2 concentration impairs the ability of Retigabine to further stabilize the OPEN2 mode. Therefore, we conclude that PI(4,5)P2 is required for the action of Retigabine on KV7 channels and can be subject to muscarinic regulation.

93. Role of S6 in Slo2 K⁺ channel closure. M. HUNTER GIESE,¹ ³ ALISON GARDNER,¹ ANGELA HANSEN,¹ MICHAEL C. SANGUINETTI¹,² ¹Nora Eccles Harrison Cardiovascular Research & Training Institute, Salt Lake City, UT 84112; ²Department of Internal Medicine, Division of Cardiovascular Medicine, University of Utah, Salt Lake City, UT 84112; ³Department of Physiology and Cellular Biophysics, Columbia University, New York, NY 11222

In contrast to Kv channels, Slo2 channels lack a functional voltage sensor domain which prevents their gating from being coupled to transmembrane voltage. Under normal conditions Slo2 channels have a very low open probability (P_o) but are readily activated by fenamates or elevated [Na]. The molecular mechanisms underlying activation and closure of Slo2 channels remains poorly understood. Previous work using drug accessibility and site-directed mutagenesis suggested the pore helix (PH) and/or selectivity filter (SF) were responsible for Slo2 channel closure. Recently, a structure for chicken Slo2.2 solved in the absence of Na⁺ identified a constriction formed by the S6 helices at Met333 as the activation gate of the channel (Hite et al. 2015. Nature. 527:198-203). Inconsistent with this model, mutation of the conserved Met residue in human Slo2.1 or Slo2.2 to the negatively charged Glu did not produce constitutively active channels. In Slo2.1, Leu267 and Leu270 are in close proximity to the SF/PH and are predicted to line the central cavity. To investigate the role of upper S6 in Slo2 channel closure, we substituted Leu267 and Leu270 with 15 different residues. Constitutive activity of Leu267 mutations depended upon volume and hydrophobicity with L267H producing maximal activity. The double mutant L267N/L270N was fully constitutively active, consistent with closure involving dewetting of the central cavity near the PH/SF.

94. Aldosterone and β-adrenergic activation of colonic K⁺ secretion requires BK channels (KCa1.1). SUSAN HALM and DAN HALM, Department of Neuroscience, Cell Biology & Physiology, Wright State University Boonshoft School of Medicine, Dayton, OH 45435

In the distal colonic epithelium, aldosterone (aldo) and epinephrine (epi) activate electrogenic K⁺ secretion via a cellular mechanism involving apical membrane K⁺ channels and basolateral membrane Cl⁻ channels. Epi [10 μM] activated a short-circuit current (epiI_sc) in mouse (C57BL/6J) ex vivo distal colonic mucosa consistent with transient transepithelial secretion of Cl⁻ and sustained K⁺ secretion. Prior addition of peptide-YY [0.3 μM] suppressed transient epiI_sc revealing sustained epiI_sc within 5 min after epi addition. Aldo [1 μM] activated a sustained I_sc consistent with K⁺ secretion after a delay of ~20 min. Subsequent epi addition further stimulated I_sc in a non-additive manner suggesting that aldo signaling converges on the epi signaling pathway producing K⁺ secretion. Secretory activation also was measured in age-matched litter-mates lacking the slo-gene product (BK-knockout). epiI_sc in BK⁰⁰ distal colon was a small transient with a steady-state similar to basal, and aldoI_sc remained similar to pre-stimulation I_sc. Addition of the BK
Blockers paxilline or iberiotoxin (IbTx) inhibited \( \text{I}_{\text{sc}} \) in wild-type colon supporting further a requirement for the BK channel in the apical membrane. Low sensitivity to IbTx suggested involvement of the \( \beta \)-subunits \( \text{kcnmb1} \) and/or \( \text{kcnmb4} \). Basal \( \text{I}_{\text{sc}} \) was more positive in \( \text{BK}^{\text{KO}} \) colon than wildtype, consistent with ongoing \( \text{Cl}^- \) secretion. Addition of the \( \text{Ca}^{2+} \)-activated \( \text{Cl}^- \) channel blocker CaC-Cinh-A01 [30 \( \mu \text{M} \)] in wild-type increased \( \text{basalI}_{\text{sc}} \) with a resulting \( \text{I}_{\text{sc}} \) similar to \( \text{basalI}_{\text{sc}} \) in \( \text{BK}^{\text{KO}} \). Partial inhibition of \( \text{basalI}_{\text{sc}} \) by the CFTR \( \text{Cl}^- \) channel blocker CFTRinh-172 [30 \( \mu \text{M} \)] suggested an unmasking of basal \( \text{Cl}^- \) secretion in \( \text{BK}^{\text{KO}} \) colon. These results indicate a near complete dependence of electrogenic \( K^+ \) secretion on the presence of the BK \( K^+ \) channel protein in the apical membrane of the mouse distal colonic epithelium. [NIH DK65845; WSU-BSOM Seed Grant]

95. Survival and growth of C57BL/6J mice lacking the BK channel (\( \text{K}_{\text{c}1.1} \)). SUSAN HALM and DAN HALM, Department of Neuroscience, Cell Biology & Physiology, Wright State University Boonshoft School of Medicine, Dayton, OH 45435

The BK potassium channel contributes to \( K^+ \) flow and the electrical behavior of many cell types. Mice made null for the gene (\( \text{Kcnma1}, \text{slo}-1 \)) producing the BK channel exhibit numerous deficits in physiological functions. Initial BK-null het breeding pairs in a C57BL/6J background strain were obtained from Dr. Andrea Meredith at the University of Maryland. A colony derived by breeding mice heterozygotically null for \( \text{Kcnma1} \) (\( \text{BK}^{\text{KO}} \)) had litter sizes of \(~8\) pups. For the period of maternal care (P0 – P21) pup death peaked at P1 with a second less severe interval of pup death peaking near P13. The later deaths of BK\(^{\text{KO}}\) pups (~P13) were greater than Mendelian expectations. Death of colony mice after weaning was rare. Pup death at ~P1 was twice as likely during the 20 month construction of a building adjacent to the animal facility compared with the period after cessation of construction. Births during construction were not consistent with Mendelian predictions indicating the likelihood of a specific disadvantage induced by this environmental stressor. After weaning, weight gain was lower for BK\(^{\text{KO}}\) mice compared with wild-type littermates: 5 grams smaller for male BK\(^{\text{KO}}\) mice and 4 grams smaller for female BK\(^{\text{KO}}\) mice. Body composition determined by quantitative magnetic resonance (QMR) indicated a higher fat proportion for wild-type female mice compared with males, as well as a higher water proportion. Both male and female BK\(^{\text{KO}}\) mice showed higher fat proportions than wild-type, with the female BK\(^{\text{KO}}\) mice exhibiting a larger variation. Together these results indicate that BK\(^{\text{KO}}\) mice suffer disadvantages that produce a tendency for early death and exhibit a general metabolic difference that leads to a smaller body mass and distinct composition. [NIH DK65845; WSU-BSOM Seed Grant]

96. CaBP1/caldendrin is an essential regulator of hippocampal synaptic plasticity and spatial memory. JASON HARDIE, TIAN YANG, JEREMIAH BRITT, GRANT STALKER, ANDREW PIEPER, AMY LEE, University of Iowa, Iowa City, IA 52242

CaBP1/caldendrin is an EF-hand Ca\(^{2+}\) binding protein that is highly expressed in the brain and strongly potentiates Ca\(^{2+}\) currents through Ca\(_{v}1\) L-type channels in heterologous expression systems. However, the physiological functions of CaBP1/caldendrin are unknown. Here, we used mice lacking expression of CaBP1/caldendrin (C-KO) to probe the role of this protein in the nervous system. Compared to wild-type (WT) mice, C-KO mice were deficient in learning and memory acquisition in the Morris water maze test. Consistent with this deficit, excitatory postsynaptic potentials in field recordings of hippocampal slices from C-KO mice showed less long-term potentiation in response to a 1 sec-
ond, 100-Hz tetanus than in recordings from WT mice (11.8±8% vs. 44.1±11.2% increase at time 60-120 min post-potentiation, p=0.043, t-test). Moreover, the Ca\textsubscript{1.1}-dependent phosphorylation of the transcription factor CREB was reduced in neuronal cultures from C-KO compared to WT mice. We conclude that CaBP1/calendrin regulates processes underlying synaptic plasticity and learning and memory, which may depend in part on its control of Ca\textsubscript{1.1}-mediated Ca\textsuperscript{2+} signaling.

97. The F-actin binding protein TRIOBP-1 regulates hERG K\textsuperscript{+} channels. ASHLEY A. JOHNSON,\textsuperscript{1} DAVID K. JONES,\textsuperscript{2} ELON C. ROTI ROTI,\textsuperscript{2} REBECCA UELMEN,\textsuperscript{2} GAIL A. ROBERTSON,\textsuperscript{2} MATTHEW C. TRUDEAU,\textsuperscript{1}  
\textsuperscript{1}Department of Physiology, School of Medicine, University of Maryland, Baltimore, MD 21201; \textsuperscript{2}Department of Neuroscience, University of Wisconsin, Madison, WI 53705

The voltage-gated potassium channel hERG is the primary, pore-forming subunit of the rapidly activating delayed rectifier potassium channel current (I\textsubscript{Kr}) in the heart. The physiological role of cardiac I\textsubscript{Kr} is to repolarize the ventricular action potential. The magnitude of native I\textsubscript{Kr} current is critical for repolarization, as highlighted by genetic mutations in the hERG gene that decrease I\textsubscript{Kr}, but the specific protein-protein interactions that regulate I\textsubscript{Kr} current (and the number of hERG channel subunits) at the plasma membrane are not well understood. We previously used the entire cytoplasmic C-terminal portion of hERG to conduct a yeast two-hybrid screen of a human cardiac library for proteins that might regulate hERG channel function, localization, stability, or surface expression. We identified TRIOBP-1, an F-actin binding protein associated with cytoskeletal dynamics, as a putative hERG interacting protein. Here, we tested for a direct functional interaction between TRIOBP-1 and hERG channels using a combination of whole-cell patch-clamp electrophysiology, Förster resonance energy transfer (FRET) measurements, and biochemical methods. We report that native I\textsubscript{Kr} current from cardiomyocytes derived from iPSCs was reduced by overexpression of TRIOBP-1-CFP in whole-cell patch-clamp recordings as compared to negative controls, showing a functional consequence of TRIOBP-1 in myocytes. Likewise, in a HEK293 cell line stably expressing hERG, expression of TRIOBP-1-CFP, but not CFP, reduced outward hERG current, suggesting that TRIOBP-1 may regulate hERG surface expression. Biotinylation treatment of HEK293-hERG cells expressing TRIOBP-1 had a marked reduction in staining of hERG surface band intensity, suggesting that TRIOBP-1 reduces the number of mature, fully glycosylated hERG channels at the membrane surface. FRET analysis revealed robust FRET between TRIOBP-1-CFP and the entire cytoplasmic C-terminal portion of hERG (hERG-666-1159-Citrine), suggesting a direct and specific interaction between TRIOBP-1 and hERG channels. Together, these experiments provide evidence that hERG channels are regulated by a direct interaction with TRIOBP-1.

98. Atomic mutagenesis in potassium channels with engineered stoichiometry. JOHN D. LUECK,\textsuperscript{1} ADAM L. MACKEY,\textsuperscript{1} DANIEL INFIELD,\textsuperscript{1} JASON D. GALPIN,\textsuperscript{1} JING LI,\textsuperscript{2} BENOÎT ROUX,\textsuperscript{2} and CHRISTOPHER A. AHERN,\textsuperscript{1}  
\textsuperscript{1}Department of Molecular Physiology and Biophysics, University of Iowa Carver College of Medicine, Iowa City, IA 52242; \textsuperscript{2}Department of Biochemistry and Molecular Biology, University of Chicago, Chicago, IL 60637

C-type inactivation of potassium channels fine-tunes electrical signaling in excitable cells through an internal timing mechanism that is mediated by a hydrogen bond network in the channels’ selectively filter. Previously, we used nonsense suppression to highlight the functional role of the highly conserved Trp434-Asp447 indole hydrogen bond in Shaker potassium channels with a non-
hydrogen bonding homologue of tryptophan, Ind. Here, molecular dynamics data indicate that the Trp434Ind hydrogen bonding partner, Asp447, unexpectedly ‘flips out’ towards the extracellular environment, allowing water to penetrate the space behind the selectivity filter while simultaneously reducing the local negative electrostatic charge. Additionally, a protein engineering approach is presented whereby split intein sequences are incorporated into the N- or C- termini of Shaker monomers that are flanked by endoplasmic reticulum retention/retrieval motifs (ERret). This system enabled stoichiometric control of Shaker monomers and the encoding of multiple Ind amino acids within a channel tetramer. We then applied the protein engineering approach to successfully incorporate two noncanonical amino acids into one peptide, the voltage-gated sodium channel Nav1.4. The developed approach circumvents a significant challenge in the field of genetic code expansion and can be used to study stoichiometry and localization of other proteins in multiple cellular environments.

99. TRP-ing the light fantastic: Activation of TRPC5 and other channels using internal perfusion on an APC platform. GESA RASCHER-EGGSTEIN,1 ANDREA BRÜGGEMANN,1 TOM GOETZE,1 SØREN FRIS,1 MARKUS RAPEDIUS,1 CLAUDIA HAARMAN,1 ILKA RINKE,1 MICHAEL GEORGE,1 JAMES COSTANTIN,2 and NIELS FERTIG,1 1Nanion Technologies Inc., München, Germany 80636; 2Nanion Technologies Inc., Livingston, NJ 07039

Ion channels have been recognized as an important therapeutic target class for treating a number of different pathophysologies. This target class is becoming even more important as the personalized medicine revolution evolves. However, a major limiting factor in the drug discovery process involving ion channels is the technically demanding electrophysiological assay which remains the gold standard since its discovery in the 1970's. Here we present data on Nanion's SyncroPatch 384PE which is a high throughput gigaseal platform, recording up to 768 experiments simultaneously. We show state-dependent effects of tetracaine on Nav1.7 expressed in CHO cells and the effect of temperature on hERG channel pharmacology. Furthermore, we show activation of TRPC5 and TMEM16A by perfusion of intracellular Ca2+ and inhibition by external application of compounds.

100. SCN8A encephalopathy: lessons from a knock-in mouse model. JACY L. WAGNON AND MIRIAM H. MEISLER, Departments of Human Genetics and Neurology, University of Michigan, Ann Arbor, MI 48109

De novo mutations of the voltage-gated sodium channel SCN8A encoding Nav1.6 are responsible for an early-onset epileptic encephalopathy designated EIEE13 (OMIM #614558). More than 140 missense mutations have been identified in individuals with epileptic encephalopathy, which includes intractable seizures, developmental delay and intellectual disability (Meisler et al. 2016. Epilepsia June B Epub ahead of print). Most of the eleven missense mutations evaluated in heterologous systems exhibited gain-of-function changes, including hyperpolarizing shifts in voltage dependence of activation, impaired channel inactivation, and elevated persistent current, that are predicted to lead to neuronal hyperexcitability (Wagnon et al. 2016. Ann.Clin.Transl.Neurol. 3:114-123). In order to evaluate the mechanisms underlying pathogenicity of these gain-of-function mutations in vivo, we used Talen targeting to generate a knock-in mouse model with the patient mutation N1768D (p.Asn1768Asp). The N1768D mutation is located at the distal end of transmembrane segment D4S6. Heterozygous mice exhibited abnormal EEG, spontaneous convulsive seizures and sudden death (Wagnon et al. 2015. Hum Molec. Genet. 24:506- 516). The detection of these patient phenotypes in the knock-in mouse indicates that the Scn8a mutation is sufficient to generate the disorder. In order to investigate the underlying pathophysiology, recordings have been made from isolated cells and brain slices. Elevated persistent current was detected in pyramidal and bipolar hippocampal neu-
rons (Lopez-Santiago...Isom, AES Abstract 3.142, 2015). CA1 and CA3 hippocampal neurons exhibited abnormal AP firing activity and hyperexcitability (Yuan...Isom AES Abstract 2.114, 2015). Hyperexcitability was also observed in layer II neurons of the medial entorhinal cortex (Barker...Patel, AES 2016). Interestingly, hyperexcitability was observed in cardiac ventricular myocytes, which express a low level of Nav1.6 in transverse tubules (Frasier...Isom, AES Abstract 2.152, 2015). Future studies in the SCN8A knock-in mouse will focus on dissection of the abnormal circuits that underlie spontaneous seizures and encephalopathy. (Supported by NIH R01 NS34509)

101. CaBP1/caldendrin regulates signaling by voltage-gated Ca_{1} L-type Ca^{2+} channels and neurite outgrowth in spiral ganglion neurons. TIAN YANG,1,2,3 JEEUN CHOI,1,2,3 DANIEL SOH,1,2,3 STEVEN GREEN,2,4 BERND FRITZSCH,4 MARLAN HANSEN,2,5 AMY LEE,1,2,3 Departments of 1Molecular Physiology and Biophysics, 2Otolaryngology Head-Neck Surgery, 3Neurology, 4Biology, and 5Neurosurgery, University of Iowa, Iowa City, IA 52242

A loss of innervation of cochlear inner hair cells by spiral ganglion neurons (SGNs) can lead to sensorineural hearing loss. Factors that regulate the growth of SGN neurites are poorly understood but could be targeted in hearing restoration strategies. Ca^{2+} signals due to strong depolarization are known to inhibit SGN neurite outgrowth, but the underlying pathways are unknown. Here, we demonstrate that a potent stimulator of Ca_{1} Ca^{2+} channel function, CaBP1/caldendrin, is an inhibitor of SGN neurite outgrowth. Auditory brain stem responses of mice lacking CaBP1/caldendrin (C-KO) show normal hearing threshold at 9 weeks of age but higher wave I amplitudes compared to wild-type (WT) mice, suggesting enhanced function of SGNs in C-KO mice. Consistent with this possibility, the repressive effect of depolarization on SGN neurite growth is significantly weaker in C-KO than WT mice. This effect was blocked by the Ca_{1} channel agonist isradipine. Ca_{1}-mediated phosphorylation of the transcription factor CREB was also reduced in C-KO SGNs. We conclude that CaBP1/caldendrin is a negative regulator of SGN neurite outgrowth, which may be due to its role in augmenting Ca_{1} channel function and coupling to transcriptional activation.

102. Bitter Taste Receptor Agonists as Potent Tocolytics for Preterm Labor. KAIZHI ZHENG,1,4 PING LU,1 TIFFANY A. MOORE SIMAS,2 ELLEN DELPAPA,2 LAWRENCE M. LIFSHITZ,3 FANGXIONG SHI,4 and RONGHUA ZHUGE,1 1Dept. of Microbiology & Physiological Systems, 2Dept. of Obstetrics & Gynecology, 3Program in Molecular Medicine, University of Massachusetts Medical School, Worcester, MA 01655; 4College of Animal Science and Technology, Nanjing Agricultural University, Nanjing, China

Preterm labor (PTL), the leading cause of neonatal mortality and morbidity, occurs in ~11% of pregnancies. Its etiologic mechanisms are complex, multifactorial, and not clearly elucidated. PTL is defined by cervical change in the presence of regular uterine contractions occurring prior to 37 weeks gestation in humans. Given contractions as a central feature, tocolytics are used in its management, yet current tocolytics are not sufficiently effective. In the present study, we report that myometrial cells from mouse and human express bitter taste receptors (TAS2Rs), a class of proteins that were long thought to be solely in the taste buds of tongues, and their canonical signaling components (i.e., G-protein gustducin and PLCβ2). Strikingly, bitter tastants (e.g., chloroquine) relax myometrium pre-contracted by uterotonic (e.g., oxytocin and prostaglandin F2α) more completely than current commonly used tocolytics (i.e., nifedipine, indomethacin and MgSO4) and at a lower dose than MgSO4 and indomethacin. Using isolated single mouse myometrial cells, we also found that bitter tastants reverse the rise in [Ca2+]i and cell shortening induced by uterotonic, and this reversal effect is inhibited by pertussis toxin treatment and α-gustducin deletion. Finally, chloroquine prevents mouse PTB in-
duced by bacterial endotoxin lipopolysaccharide or progesterone receptor antagonist RU486 more often than current commonly used tocolytics, and this prevention is largely lost in α-gustducin knockout mice. In summary, myometrial cells express the canonical bitter taste receptor signaling system, and bitter tastants demonstrate the potential to be more effective tocolytics than those currently used for PTL management.

103. A small molecule inhibitor of TRPC5 suppresses proteinuric kidney disease progression. YIMING ZHOU, PHILIP CASTONGUAY, MORAN LEVITT-DVELA, SOOKYUNG KIM, ERIENE SIDHOM, JONAS SIEBER, JANA REICHARDT, FRANK DUBOIS, SIGRID HOFFMAN, JOHN BASGEN, ASTRID WEINS, PETER MUNDEL, COREY HOPKINS and ANNA GREKA, Department of Medicine and Glom-NExT Center for Glomerular Kidney Disease and Novel Experimental Therapeutics, Brigham and Women’s Hospital and Harvard Medical School, Boston, MA 02115; The Broad Institute of MIT and Harvard, Cambridge, MA 02142

Focal Segmental Glomerulosclerosis (FSGS) is a progressive kidney disease that can be caused by Rac1-activating mutations leading to filter barrier breakdown and podocyte death. Clinical diagnosis depends on the detection of abnormal spilling of essential proteins in the urine called proteinuria. Previous studies have implicated Angiotensin Type 1 Receptor-induced, Rac1-mediated TRPC5 channel activity in podocyte injury. However, the question of whether TRPC5 channel inhibition may confer a therapeutic benefit in acquired, progressive proteinuric kidney disease remains unresolved. Here we demonstrate at the single channel level that TRPC5 activity in podocytes on intact glomeruli escalates in concert with active Rac1 and proteinuric kidney disease progression. A small molecule inhibitor of TRPC5 prevents Rac1-induced ROS generation, and protects podocytes from cell death. We further demonstrate that chronic administration of a TRPC5 inhibitor ameliorates established, heavy proteinuria and prevents podocyte loss in a rat model of FSGS. These data indicate that TRPC5 activity is induced to drive FSGS, and that TRPC5 inhibitors may be valuable for the treatment of progressive proteinuric kidney disease.
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