

309

Measurement of agonist-induced calcium and mechanical signals in human airway smooth muscle cells

Santina Johnson¹, Naga Annamdevula², Josephine Jalkh², Zara Ijaz², Andrea L. Britain², Dhananjay Tambe², C. Michael Francis², Deepak Deshpande³, Silas J. Leavesley², Thomas C. Rich²

¹University of South Alabama, College of Medicine ²University of South Alabama ³Thomas Jefferson University

OBJECTIVES/GOALS: The goal of this proposal is to develop a technology that combines calcium imaging via confocal microscopy, and force measurement via monolayer stress microscopy to perform simultaneous quantitative measurements of agonist-induced Ca²⁺ and mechanical signals in HASMCs. **METHODS/STUDY POPULATION:** The methods by which second messenger signals and changes in mechanical forces determine specific physiological responses are complex. Recent studies point to the importance of temporal and spatial encoding in determining signal specificity. Hence, approaches that probe both chemical and mechanical signals are needed. We combine hyperspectral imaging for second messenger signal measurements, monolayer stress microscopy for mechanical force measurements, and S8 analysis software for quantifying localized signals. Imaging was performed using an excitation-scanning hyperspectral microscope. Hyperspectral images were unmixed to identify signals from fluorescent labels and microparticles. Images were analyzed to quantify localized force dynamics through monolayer stress microscopy. **RESULTS/ANTICIPATED RESULTS:** Results indicate that localized and transient cellular signals can be quantified and mapped within cell populations. Importantly, these results establish a method for simultaneous interrogation of cellular signals and mechanical forces that may play synergistic roles in regulating downstream cellular physiology in confluent monolayers. **DISCUSSION/SIGNIFICANCE:** We will measure the distribution of chemical and mechanical signals within cells, providing insight into the dynamics of cell signaling. Studies will have implication in the understanding of infections, drug delivery in which non-uniform distributions of drugs are a certainty, and in understanding coordinated responses in cellular systems.

310

Mechanisms of progression of myelodysplastic syndrome to fatal secondary acute myeloid leukemia

Daniel Chang, Klara Noble-Orcutt, Marie Lue Antony, David Largaespada, Chad Myers, Zohar Sachs
University of Minnesota

OBJECTIVES/GOALS: Aim 1: Define the genetic events that cooperate with Trp53 mutations to mediate the transformation of MDS to sAML at the stem cell level. Aim 2: Use Bayesian networks to model the signaling pathway activation states identify aberrant regulators of self-renewal in LSCs. **METHODS/STUDY POPULATION:** To model the transformation of MDS, I will utilize a mouse model of MDS and sAML. The Trp53 mutated mouse mimics the genetics of human MDS and develops MDS. To discover how additional mutations contribute to disease progression, I use the Sleeping Beauty transposon system which induces random mutations. I propose to use this mouse model to define the molecular mechanisms of transformation of MDS to sAML. I will also measure levels of activated signaling intermediates in sAML using CyTOF. CyTOF is a method of flow cytometry that measures up to 40 proteins simultaneously at single-cell resolution. I will also use Bayesian network to model the signaling architecture of

the MDS/sAML stem cells. **RESULTS/ANTICIPATED RESULTS:** I will identify putative mediators of self-renewal in gene expression data and the corresponding Bayesian networks model. I will prioritize genes and signaling intermediates that act in pathways that rank highly in both methods and those that have previously published roles in self-renewal. Based on prior work, I will focus on EZH2, NF- κ B, and mTOR pathways in addition to candidate pathways revealed by these studies. Small molecule inhibitors that target candidate signaling intermediates will be used for experimental validation. I will test the impact of these small molecule inhibitors on LSC self-renewal using serial colony forming assays as well as in vivo mouse reconstitution assays. **DISCUSSION/SIGNIFICANCE:** We focus on the role of stem cells in the transformation of MDS to sAML. Understanding the mutations and signaling relationships that mediate self-renewal in Trp53 mutant stem cells will allow for precise therapeutic target of this disease and improve outcomes for these patients.

311

Mental Illness and the Development of Postoperative Atrial Fibrillation in Transcatheter Aortic Valve Replacement Patients: Trends over Time

Natalie Kolba², Jennifer Morrone², Julia Dokko², Samantha Novotny², Jie Yang³, Vineet Tummala², Sohaib Agha⁴, Ashutosh Yaligar⁴, Puja B. Parikh⁵, Aurora D. Pryor⁴, Henry J. Tannous⁴, A. Laurie Shroyer⁴, Thomas Bilfinger⁴

¹Renaissance School of Medicine at Stony Brook University

²Renaissance School of Medicine, Undergraduate Medical Education, Stony Brook University, Stony Brook, NY ³Renaissance School of Medicine, Office of Dean, Stony Brook University

⁴Department of Surgery, Stony Brook University School of Medicine, Stony Brook, NY ⁵Department of Medicine, Stony Brook University School of Medicine, Stony Brook, NY

OBJECTIVES/GOALS: The purpose of this retrospective cohort study was to evaluate the impact of mental illness on first-time transcatheter aortic valve replacement (TAVR) and repeat TAVR (viv-AVR) outcomes including postoperative atrial fibrillation (POAF/AFL), as well as trends over time. **METHODS/STUDY POPULATION:** Using de-identified data reports from the New York Statewide Planning and Research Cooperative System (SPARCS) database from 2005-2018, multivariate logistics models were used to predict endpoints including POAF, the Society of Cardiothoracic surgeon (STS) endpoint (MM), and 30-day readmission (READMIT) in patients with and without mental illness. The TAVR procedure was approved for high-risk patients after 2012, and intermediate-risk patients after 2016, indicating a need to analyze the two populations separately. Multivariate analysis was only conducted on the first-time TAVR patients because of the small n in the viv-TAVR population. **RESULTS/ANTICIPATED RESULTS:** After 2012, 13.05% (1,810/13,870) of patients undergoing TAVR and 20.83% (15/72) undergoing viv-TAVR were diagnosed with a mental illness before the procedure. After 2016, 15.59% (1,485/9,524) TAVR patients and 20.00% (11/55) viv-TAVR patients had a preoperative diagnosis of mental illness. Multivariate analysis showed that mentally ill patients did not have significant differences in rates of POAF, 30-day readmission, and 30-day composite outcomes when compared to patients without mental illnesses following TAVR procedures after 2012 and 2016. Patients with POAF after both 2012 and 2016 were significantly less likely to be mentally ill, Black, and Hispanic. **DISCUSSION/SIGNIFICANCE:** Of the mentally ill patients who underwent TAVR, there was no significant difference in short-term outcomes after 2012 vs. 2016, compared to patients without mental

illnesses. The small number of mentally ill patients undergoing TAVR may point to provider bias as a contributor to this high selectivity, and further evaluation would be of clinical use.

312

Metabolomics Approach to Cellular Senescence: Characterizing the Secretory Phenotype and Metabolism of Irradiation-Induced Senescent Human Pre- Adipocytes

Zinnarky Kiani Ortiz-Correa, Ian R. Lanza
Mayo Clinic, Rochester, MN

OBJECTIVES/GOALS: The study aims to identify and measure metabolites in irradiation-induced senescent pre-adipocytes induced by the increased secretion of pro-inflammatory cytokines. Through untargeted metabolomics, we seek to understand the alterations to metabolism and mitochondrial activity that occur during irradiation-induced senescence. **METHODS/STUDY POPULATION:** First, commercially available human primary subcutaneous pre-adipocytes were cultured in vitro, and irradiated to induce senescence. To confirm senescence, cells were stained for beta-galactosidase, which was positive in senescent pre-adipocytes. The cells and their conditioned cultured media were then collected and frozen for untargeted metabolomics to profile metabolites. The sample analysis is currently underway and will be conducted using central carbon isotope tracing and chromatograph mass spectrometry. Principal Component Analysis, fold change analysis, and heat maps will be used to detect and report the changes in metabolite signals. Oxygen consumption rate and extracellular acidification rate measurements are in progress at present using the Agilent Seahorse XF Cell Mito Stress Test. **RESULTS/ANTICIPATED RESULTS:** We expect that metabolites of central carbon metabolism and oxidative phosphorylation will be upregulated. The concentrations of metabolites of pathways altered by the pro-inflammatory cytokines that have been identified to be secreted by senescent cells are expected to be altered. The metabolites measured in conditioned culture media will provide insight into the changes to the cellular microenvironment caused by senescence. Finally, we anticipate that mitochondrial function, and both aerobic and anaerobic metabolism will be altered in senescent pre-adipocytes compared to controls. Through the present study, we will achieve a better understanding the metabolic alterations that occur as part of cell senescence in pre-adipocytes. **DISCUSSION/SIGNIFICANCE:** Adipose tissue dysfunction due to the accumulation of senescent cells has been linked to chronic diseases such as diabetes and insulin resistance, and inflammation. Through this study, we expect to provide new insight into the metabolic alterations of senescence and offer a backbone for prospective translational human studies and clinical trials.

313

Microstructural changes in the brainstem regions of the auditory pathway among children with hearing loss

Iram Ahmad, Peter Moon, Kristi Ward, Taseer Din, Sara Saki, Alan Cheng, Kristen Yeom
Stanford University

OBJECTIVES/GOALS: Diffusion MRI can identify microstructural brain changes and can provide insight into neural

development and to potentially prognosticate speech and language outcomes in children with SNHL. The goal of our study was to investigate MRI-based microstructural changes along the brainstem regions of the auditory pathway in pediatric patients with SNHL. **METHODS/STUDY POPULATION:** We reviewed cohort of pediatric patients with SNHL who obtained MRI at 3T between 2011 and 2019. We identified 16 pediatric patients (age **RESULTS/ANTICIPATED RESULTS:** We identified significant differences in FA values of the SON between the SNHL cohort and controls (0.377 ± 0.056 vs 0.422 ± 0.052 ; $p=0.009$). No other FA or MD values were significantly different between the two groups. Among younger children (≤ 5 years), MD was significantly decreased in the SNHL cohort compared to controls in the IC (0.918 ± 0.051 vs 1.120 ± 0.142 ; $p5$ years), there were no significant differences in MD (1.124 ± 0.198 vs 0.997 ± 0.103 ; $p = 0.119$). There were no significant differences in MD or FA in the white matter fibers of the IC-SON tract. **DISCUSSION/SIGNIFICANCE:** This study is the first to assess microstructural changes in brainstem auditory pathway regions among children with SNHL. Longitudinal studies are warranted to assess the predictive value of these MRI-based findings for long-term outcomes and the efficacy of intervention.

315

Non-invasive and quantitative surgical outcome evaluation of patients with sagittal craniosynostosis undergoing sagittal craniectomy*

Connor Elkhill¹, Jiawei Liu², Marius George Linguraru³, Scott LeBeau⁴, David Khechayan⁵, Brooke French⁵, Antonio R. Porras⁶
¹Computational Bioscience Program, University of Colorado Anschutz Medical Campus ²Department of Biostatistics and Informatics, Colorado School of Public Health ³Children's National Health System, Washington, DC ⁴Plastic & Reconstructive Surgery, Children's Hospital Colorado ⁵Plastic & Reconstructive Surgery Children's Hospital Colorado, University of Colorado School of Medicine ⁶Department of Biostatistics and Informatics, Colorado School of Public Health, Children's Hospital Colorado

OBJECTIVES/GOALS: Craniosynostosis is the premature fusion of one or more cranial sutures that produces brain growth constraints and typically requires surgical treatment. We present an age- and sex-specific method to evaluate surgical outcomes using non-invasive 3D photogrammetry that brings objectivity to the current approach for clinical assessment. **METHODS/STUDY POPULATION:** First, we created standardized head anatomy representations for 2,020 patients (1,081 males, 939 females, age 3.14 ± 3.05 years) without cranial pathology from their computed tomography (CT) images based on our previous methods. We used principal component regression stratified by sex to establish age-specific normative ranges of anatomical variability and we designed a new metric called cranial shape abnormality (CSA) index that calculates the number of standard deviations from normality of a given patient's head anatomy. We calculated our CSA index in a group of 56 patients (44 male, 12 female) with sagittal craniosynostosis who underwent sagittal craniectomy from their pre- (22 ± 30 days before surgery) and post-surgical (267 ± 63 days after surgery) 3D photograms to evaluate surgical outcomes. **RESULTS/ANTICIPATED RESULTS:** We observed a reduction in the CSA index from 1.28 ± 0.26 before surgery to 0.87 ± 0.22 after surgery ($p < 0.001$ with a paired Wilcoxon test). The CSA index decreased in 53 of 56 patients (94.6%), who consistently