MONDAY POSTERS SESSIONS

Blood Flow Regulation

M-01
Upper limb occlusive hyperemia evokes the same response mechanism in the contralateral forearm – data from functional imaging with optoacoustic tomography

Sergio F Andrade MSc, PhD, Tiago Granja PhD, Luis M Rodrigues MSc, PhD
CBIOS – Universidade Lusofona’s Research Center for Biosciences & Health Technologies, Lisboa, Lisboa, Portugal

- To better understand the response to reactive hyperemia and the venoarteriolar reflex.
- These are considered to evoke local adaptive responses used to study vascular physiology.
- Our results demonstrated that any perfusion adaptive response involves a centarlly mediated reflex.

M-02
Maintenance of blood-brain barrier integrity during mammalian hibernation

Maryann Platt PhD1,2, Slav Bagriantsev PhD1, Elena Gracheva PhD1,2
1Yale University, New Haven, CT, USA. 2Yale Kavli Center for Neuroscience, New Haven, CT, USA

- Hibernation provides a naturalistic model of low blood flow in which to study reperfusion.
- Facets of the blood brain barrier are differentially regulated during hibernation.
- Blood flow velocity scales with body temperature during arousal from hibernation.

M-03
Endothelial hemoglobin alpha mediates iron regulation of endothelial nitric oxide

Luke S Dunaway PhD, Shruthi Nyshadham BS, Melissa A Luse MS, Skylar Loeb BS, Adam N Goldfarb MD, Brant E Isakson PhD
University of Virginia, Charlottesville, VA, USA

- Endothelial iron regulates nitric oxide signaling.
- Increased nitric oxide signaling in iron deficiency anemia is not due to reduced blood hemoglobin.
- Endothelial hemoglobin alpha regulates nitric oxide in response to changing iron levels.

M-04
A multiscale mathematical model to examine the role of capillary pericytes in cerebral blood flow control

Niloufar Khakpour PhD Student1, Dabashis Kumar Saha PhD Student1, Nicholas R. Klug PhD, Postdoctoral Associate2, Mark T. Nelson PhD, Distinguished Professor & Chair2, Nikolaos M. Tsoukias PhD, Professor1
1Florida International University, Miami, Florida, USA. 2University of Vermont, Burlington, Vermont, USA

- Present an integrated model to explore pericytes’ role in coordinating local blood flow distribution.
- Pericytes play a key role in maintaining constant brain perfusion through myogenic autoregulation.
- Pericytes promote uniform blood flow distribution preserving supply to vulnerable brain regions.
M-05
Flowing through the kidney microvasculature: glomerular endothelial cells and their role in chronic kidney disease

Nathalie Reinhard, Sabine Leonhard, Annick Werner, Barbara Wilmeringwetter, Matthias Mueller, Ivan Formentini, Philipp Hoppe
Novartis Institutes for Biomedical Research (NIBR), Basel, Switzerland

- Modeling kidney hyperfiltration
- Generating novel glomerular endothelial cell model systems
- Microvasculature drug targets in chronic kidney disease

M-06
Sickle trait and alpha thalassemia increase NOS-dependent vasodilation in human arteries through disruption of endothelial hemoglobin-eNOS interactions

Steven Brooks PhD1, Parker Ruhl MD, MHS1, Xianke Zeng MD, PhD1, Phillip Cruz PhD1, Sergio Hassan PhD1, Olena Kamenyeva PhD1, Juraj Kabat PhD1, Sundar Ganesan PhD1, Rachel Smith BS1, Md Hakim PhD1, Mary Jackson RN1, Jessica de Rivera BS1, Jarrett Jackson MD1, Robert Emeh BS1, Naomi Tesfuzigta BS1, Kyeisha Laurence BS1, Stacy Joyce PA-C1, Christina Yek MD1, Sophana Chea BS1, Brant Isakson PhD1, Jessica Manning MD1, Jeremy Davis MD2, Hans Ackerman MD, DPhil1

1National Institute of Allergy and Infectious Diseases, Bethesda, MD, USA. 2National Cancer Institute, Bethesda, MD, USA. 3University of Virginia, Charlottesville, VA, USA

- Human resistance arteries express hemoglobin that regulates the release of nitric oxide (NO).
- Genetic variation in hemoglobin is associated with NO signaling and blood pressure in humans.
- Disruption of hemoglobin-eNOS interactions increases vascular NO signaling in human arteries.

M-07
Cardiac arrest induces “time sensitive” impairments to cerebrovascular autoregulation

Haishuo Guan, Arvind Pathak, Romergrýko G. Geocadin, Nitish V. Thakor, Janaka Senarathna
Johns Hopkins University, Baltimore, MD, USA

- Cerebral blood flow regulation changes substantially immediately after resuscitation
- Autoregulation is a key indicator of brain health
- Timely clinical intervention is crucial for mitigating brain injury.

Diseases (Vascular and Cardiovascular) I

M-11
Evaluation of IMA2a in a rat model of pulmonary arterial hypertension

Boris Tchernychev PhD, Kevin O’Brien PhD, Lisa Flaman MS, Jennifer Howe BS, Daniel Ortiz PhD, Yves Sabbagh PhD
Inozyme Pharma, Boston, MA, USA

- Enpp1 hydrolyzes extracellular ATP to AMP, a precursor of adenosine.
- In pulmonary arterial hypertension (PAH) adenine nucleotide signaling is dysregulated.
- Effect of IMA2a, an Enpp1-Fc protein, on disease progression was evaluated using rats with PAH.
M-12
TERT transcriptional activation is sufficient to restore NO production in blood vessels following Doxorubicin treatment

Makenna Wells, Andreas Beyer PhD
Medical College of Wisconsin, Wauwatosa, WI, USA
• Cardio-Oncology
• Endothelial function
• Microcirculation

M-13
Epigenetic perturbations in a mouse model of Loeys-Dietz Syndrome: Stress responses and phenotypic plasticity explored through histone modification analysis in Smooth Muscle Cells

Leda Restrepo PhD, Emily Bramel BS, Wendy Espinoza-Carnejo BS, Elena MacFarlane PhD
Johns Hopkins, Baltimore, MD, USA
• LDS showed enrichment of motifs related to phenotypic plasticity, proliferation, and stress response
• Epigenetic changes in LDS SMCs may drive different responses that contribute to aortic disease
• Selectively targeting maladaptive responses could be a promising therapeutic approach in LDS

M-14
Deletion of the acid-sensing ion Channel1A gene leads to hyperaldosteronism-induced hypertension with age

Selina M Garcia PhD, Laura V Gonzalez Bosc PhD, Nikki L Jernigan PhD
University of New Mexico School of Medicine, Albuquerque, NM, USA
• The loss of the acid-sensing ion channel 1a contributes to age-induced hypertension
• The loss of the acid-sensing ion channel 1a contributes to hyperaldosteronism
• The loss of the acid-sensing ion channel 1a does not affect angiotensin II levels

M-15
The role of Neuropilin-1 and p130Cas in pulmonary arterial hypertension

James M Berg1, Jigisha A Patel PhD1, Ameera Milhan1, Elena Rahmani1, Shahin Moledina MBBS2,1, David L Selwood PhD3, Lucie H Clapp PhD3, Paul Frankel PhD1
1Institute of Cardiovascular Science, University College London, London, United Kingdom. 2Pulmonary Hypertension Service for Children, Great Ormond Street Hospital NHS Foundation Trust., London, United Kingdom. 3Wolfson Institute for Biomedical Research, University College London, London, United Kingdom
• Pulmonary Arterial Hypertension (PAH) as a Life-Threatening Disease
• Neuropilin-1 (NRP1) and Growth Factor Signalling
• Unravelling Mechanisms and Pathobiology

Metabolism and Metabolic Diseases

M-17
Identifying how endothelial protease-activated receptors control insulin signaling: implications for diabetes

Rahul Rajala B.A., B.S.1,3, Courtney T Griffin Ph.D.1,2
1Oklahoma Medical Research Foundation, Oklahoma City, OK, USA. 2University of Oklahoma Health Sciences Center, Oklahoma City, OK, USA. 3Harold Hamm Diabetes Center, Oklahoma City, OK, USA
• Identifying novel crosstalk between the Insulin Receptor and Protease-Activated Receptors in ECs.
• Identify how insulin receptor splicing affects endothelial insulin transcytosis.
• Identify synergistic roles for PAR1 and PAR4 in endothelial cells.

M-18
Pharmacological inhibition and endothelial cell deletion of tissue-nonspecific alkaline phosphatase disrupt cellular bioenergetics in brain endothelial cells
Shokofeh Rahimpourkaldeh, Sujung Jun, Divine C Nwafor MD/PhD, Candice M Brown PhD
West Virginia University, Morgantown, West Virginia, USA
• TNAP, Blood-brain barrier, Brain endothelial cell, Glycolysis, Maximal respiration
• Stroke, TNAP, Oxidative phosphorylation, Glycolysis, Brain endothelial cell
• TNAP, BBB, Stroke, Oxidative phosphorylation, Glycolysis, ATP

Pro-Inflammatory Signaling
M-19
Neuropilin-1 interacts with VE-cadherin and TGFB2 to stabilize adherens junctions and prevent activation of endothelium under flow
Anissa Chikh PhD1, Emy Bosseboeuf2, Dhilakshani Vignaraja MRes3, Ridhi Rajendrakumar MRes2, Tom Mitchell PhD2, Thomas D Nightingale PhD2, Rayomand S Khambata2, Anna M Randi PhD3, Justin C Mason PhD2, Amrita Ahluwalia PhD2, Claudio Raimondi PhD2
1St. George’s University of London, London, London, United Kingdom. 2Queen Mary University of London, London, London, United Kingdom. 3Imperial College of London, London, London, United Kingdom
• Under laminar flow, NRP1 stabilizes cell-cell junctions and suppresses endothelial inflammation.
• NRP1 deletion in the endothelium induces vascular inflammation and atherosclerosis
• NRP1 interacts with TGFβ receptor II reducing TGFB2 membrane localization and TGF-β signaling.

M-20
Hresistin initiates early immune responses through macrophages to activate and proliferate pulmonary smooth muscle cells
Udeshika Kariyawasam PhD, John Skinner B.Sc, Qing Lin MD, Ph.D, Roger Johns MD, Ph.D
Department of Anesthesiology and Critical Care Medicine, Johns Hopkins School of Medicine, Baltimore, MD, USA
• Hresistin activates macrophages to induce vascular remodeling over time for PH development.
• Hresistin is a critical regulator of early immune responses in PH development.
• Hresistin is a good therapeutic target to prevent inflammation driven vascular remodeling and PH.

M-21
Loss of endothelial Ripk3 increases vascular permeability following ischemia-reperfusion injury by regulating cytokine and adhesion molecule expression
Charmain F. Johnson Ph.D.1, Kathryn Y. Burge Ph.D.2, Hala Chaaban M.D.2, Courtney T. Griffin Ph.D.1,3
1Oklahoma Medical Research Foundation, Oklahoma City, OK, USA. 2Neonatal-Perinatal Medicine, Department of Pediatrics, University of Oklahoma Health Sciences Center, Oklahoma City, OK, USA. 3Department of Cell Biology, University of Oklahoma Health Sciences Center, Oklahoma City, OK, USA
• Endothelial RIPK3 protects against vascular permeability after ischemia-reperfusion injury (IRI)
• Loss of endothelial Ripk3 increases VCAM-1, ICAM-1 and endothelial IL-6 cytokine expression
• Increased inflammatory markers with loss of Ripk3 contributes to hyperpermeability after IRI
M-22

**Macrophage IL-1β signaling activates Rac1-NF-κB-driven transcription of ACE2**


1Warren Alpert Medical School at Brown University, Providence, RI, USA. 2Ocean State Research Institute Inc. at Providence VA Medical Center, Providence, RI, USA. 3University of Missouri, Columbia, MO, USA

- Macrophage ACE2 expression is dynamic
- Macrophage ACE2 expression is upregulated by inflammatory pathways involving IL-1B-driven NF-kB
- Macrophages may act as a reservoir for ongoing infection through ACE2 upregulation with inflammation

M-23

**Atherosclerotic calcification requires macrophage IL-1β expression**

Cadence Lee Sc.M., Joshua Berus PhD, Rachel Carley PhD, Elizabeth Amelotte, Sheila Sharma, Julia Pierce AB, Chris Mantsounga PhD, Jade Neverson, Celia Butler MPH, Gaurav Choudhary, Alan R. Morrison MD/PhD

1Warren Alpert School of Medicine at Brown University, Providence, RI, USA. 2Ocean State Research Institute Inc. at Providence VA Medical Center, Providence, RI, USA

- IL-1β-deleted animals demonstrated significantly decreased atherosclerotic calcification
- Plasma analysis demonstrated modest changes in lipid metabolism
- Calcified human plaques have CD68+ macrophages expressing IL-1β adjacent to areas of calcification

M-26

**Novel mechanisms controlling NFAT-mediated vascular inflammation**

Ramoji Kosuru PhD, Behshid Ghadrdoost Nakchi PhD, Guru Prasad Sharma PhD, Akiko Mammoto PhD, Tadanori Mammoto PhD, David Zhang PhD, Paul Goldspink PhD, Mohamed Trebak PhD, Magdalena Chrzanowska PhD

1Versiti Blood Research Institute, Milwaukee, WI, USA. 2Medical College of Wisconsin, Milwaukee, WI, USA. 3The University of Illinois Chicago, Chicago, IL, USA. 4University of Pittsburgh, Pittsburgh, PA, USA

- Rap1A restricts Orai1-mediated Ca2+ entry and expression in endothelial cells
- Endothelial Rap1 restricts NFAT-mediated transcription and vascular inflammation.
- A novel role for Rap1A in controlling vascular integrity

M-27

**Interleukin-1β induced neuroinflammation inhibits blood-brain barrier development**

Audrey R Fetsko BS, Dylan J Sebo BS, Michael R Taylor PhD

University of Wisconsin-Madison, Madison, WI, USA

- Induction of IL-1β in the CNS of zebrafish embryos promotes neuroinflammation.
- IL-1β inhibits CNS angiogenesis and the acquisition of blood-brain barrier properties.
- IL-1β blocks Wnt/β-catenin signaling in brain endothelial cells during neurovascular development.

M-28

**Implication of non-histone protein citrullination in smooth muscle cell degeneration and abdominal aortic aneurysm**

Zain Islam, Li Yin, Aaron Marcum, Eric Kent, Nicholas Hoyt, Quang Le, Nina Islam, Campbell Johnson, Mengxue Zhang, Yatao Shi, Lian-wang Guo, Lingjun Li, Bowen Wang
Biomaterials, Bioengineering and Matrix Biology

M-29

Extracellular matrix hydrogel targeting thrombospondin-2 and hypoxia improves diabetic wound healing via angiogenesis

Yaqing Huang1,2, Hao Xing Ph.D.1,3, Themis Kyriakides Ph.D.1,2,3
1Vascular Biology and Therapeutics Program, Yale University, New Haven, CT, USA. 2Department of Pathology, Yale University, New Haven, CT, USA. 3Department of Biomedical Engineering, Yale University, New Haven, CT, USA

- Impaired HIF-1a in diabetic wounds prevents hypoxia-induced repression of TSP2.
- Stabilization of HIF-1a using DMOG reduced TSP2 production in diabetic fibroblasts.
- TSP2KO ECM hydrogel with DMOG enhanced angiogenesis and healing in diabetic wounds.

M-30

Role of Shp2 in angiogenic matrix degradation

Barbara Szynal, Jordan Fauser PhD, Vincent Huyot, Jennifer Klomp PhD, Andrei Karginov PhD
University of Illinois Chicago, Chicago, IL, USA

- Shp2 regulates endothelial matrix degradation in a non-phosphatase dependent manner.
- Shp2 substrate interactions drive initiation and localization of matrix degradation.
- NS-ML Shp2 mutants show altered intracellular localization and matrix degradation.

M-31

Embedding biomimetic vascular networks via coaxial sacrificial writing into functional tissues

Paul P Stankey1,2, Sebastien G.M. Uzel PhD1,2, Katharina K Kroll PhD1,2, Alexander J Ainscough PhD1,2, Daniel S Reynolds PhD1,2, Alexander Elamine1,2, Jennifer A Lewis ScD1,2
1Harvard University, Cambridge, MA, USA. 2Wyss Institute, Boston, MA, USA

- We report a novel technique termed coaxial sacrificial writing into functional tissue (co-SWIFT).
- We have developed a method to 3D print hierarchically branching networks of core-shell vasculature.
- We have embedded biomimetic, multilayer vasculature into thick cardiac tissue via co-SWIFT.

M-32

Exploring endothelial protein c receptor as a therapeutic target: mapping distribution and correlation with nanoparticles targeting

Grace D.M. Eriksen PhD student, Andrew J. Urquhart Associate professor, Martin Bak Senior Officer, David Schultz PhD student, Heidi Arps, Doha Ghanam MSc, Anja Brus PhD, Tsinat Berhane MSc, Thomas L. Andresen Professor Department of Health Technology, Kgs.Lyngby, Denmark

- EPCR targeting in disease dysfunction: Challenges and potential
- Mapping EPCR distribution for enhanced drug delivery
- Nanoparticle strategies for addressing endothelial dysfunction
M-33
A high-fidelity computational study of red blood cell trafficking and capillary hemodynamics in tumor/angiogenic microvascular networks
Abhay Mohan, Prosenjit Bagchi
Rutgers University, Piscataway, NJ, USA
- tumor/angiogenic microvascular networks
- red blood cells and blood flow distribution
- computational model, wall shear stress

M-34
Regulatory role of IncRNA CASC15 in vascular remodeling and its implications for chronic kidney disease
Cristina Espinosa-Diez1,2, Ibrahim Ahmed1, Jianxin Wei1, Mingjun Liu1, Scott Hahn1, Satoshi Okawa1, Thiago Bruder-Nascimento1, Adam C Straub1, Sruti Shiva1, Delphine Gomez1
1University of Pittsburgh, Pittsburgh, PA, USA. 2Wayne State University, Detroit, MI, USA
- Inhibition of Ang-II dependent IncRNA prevents contribution to vascular remodeling
- Loss of this IncRNA could exacerbate CKD progression
- Rescue of Ang-II dependent IncRNA could enhance kidney revascularization and prevents CKD.

M-35
Oncostatin-M signaling drives capillary leak in critically ill children by inducing phenotypical changes in endothelial cells and a novel pathway of transcellular permeability
Giulio Fulgoni PhD, Weiming Ni PhD, Alam Khan PhD, Francesc Lopez PhD, Guilin Wang PhD, John Giuliano MD, Jordan Pober MD, PhD, Richard Pierce MD
Yale University, New Haven, CT, USA
- Oncostatin-M (OSM) signaling is upregulated in ECs isolated from patients who develop capillary leak
- OSM induce transcellular endothelial permeability, possibly through PLVAP1-dependent PM-structures
- OSM signals through a JAK/STAT3 pathway and broadly reprogram EC transcriptome and function

M-36
Do capillary endothelial cell characteristics differ along the length of a capillary: implications for active hyperaemia
Mackenzie E Charter, Barbara M Hyde-Lay, Coral L Murrant
University of Guelph, Guelph, ON, Canada
- Endothelial cells are heterogeneous across a single capillary in skeletal muscle
- Endothelial cell heterogeneity could have implications for how capillaries coordinate blood flow
- Endothelial cell heterogeneity could be an important organizational component of capillary units

M-37
Redox-sensitive mitochondrial acid-sensing ion channel 1a regulates mitochondrial membrane potential in pulmonary arterial smooth muscle cells
Megan N Tuineau, Lindsay M Herbert, Tracylyn R Yellowhair, Thomas C Resta PhD, Nikki L Jernigan PhD
University of New Mexico School of Medicine, Albuquerque, NM, USA
- Acid-sensing ion channel 1 is expressed in mitochondria in pulmonary arterial smooth muscle cells
Mitochondrial acid-sensing ion channel 1 promotes mitochondrial membrane potential depolarization
Oxidants increase localization of acid-sensing ion channel 1 to mitochondria

M-38
The phosphodiesterase-9A inhibitor PF04447943 via a peroxiredoxin mechanism improves coronary microvascular rarefaction in the ZSF1 rat model of HFpEF

Katie Anne Fopiano¹, William M Pearson¹, Vadym Buncha¹, Davis J Hardell¹, Liwei Lang PhD¹, Nazha Hamdani MD, PhD², Zsolt Bagi MD, PhD³
¹Medical College of Georgia at Augusta University, Augusta, USA. ²Ruhr University Bochum, Bochum, Germany
- Coronary microvascular rarefaction contributes to the development of HFpEF.
- Coronary angiogenic endothelial sprouting can be improved via a peroxiredoxin mechanism.
- Treatment with selective PDE9A inhibitors improve coronary microvascular density.

M-39
Stimulating in vivo angiogenesis with sustained local release of an optimized proangiogenic protein cocktail

Stephanie M Roser M.Sc, Alicia J Minor Ph.D., Rajeev J Kant Ph.D., Collin Poluchia B.S., Kareen LK Coulombe Ph.D.
Brown University, Providence, Rhode Island, USA
- Regeneration
- Biomaterials
- Ischemia

Angiogenesis

M-40
Differential endothelial cell cycle status in post-natal retinal vessels revealed using a novel PIP-FUCCI reporter

Ziqing Liu PhD¹,², Natalie T Tanke BS³, Alexandra Neal BS¹, Tianji Yu BS¹, Jean G Cook PhD¹, Victoria L Bautch PhD¹
¹UNC-Chapel Hill, Chapel Hill, NC, USA. ²Medical College of Wisconsin, Milwaukee, WI, USA
- A novel PIP-FUCCI mouse for improved labeling of cell cycle phases with fluorescent reporters
- PIP-FUCCI precisely distinguish the onset of S phase and the S/G2 transition in primary EC and mice
- A novel image analysis pipeline to define EC cell cycle status in distinct vascular zones

M-41
Investigating the role of inflammasome agonism in pathological angiogenesis

Ryan D Makin¹,²,³, Ivana Apicella PhD¹,², Roshni Dholkawala MD¹,², Shinichi Fukuda MD, PhD², Shuichiro Hirahara MD, PhD², Yoshio Hirano MD, PhD², Younghee Kim PhD¹,², Ayami Nagasaka PhD¹,², Yosuke Nagasaka MD, PhD¹,², Siddharth Narendran MD², Felipe Pereira MD², Akhil Varshney PhD⁵, Shao-bin Wang PhD¹,², Jayakrishna Ambati MD¹,²,³,⁴,⁵,⁶, Bradley D Gelfand PhD¹,²,³,⁶,⁷
¹Center for Advanced Vision Science, University of Virginia School of Medicine, Charlottesville, VA, USA.
²Department of Ophthalmology, University of Virginia School of Medicine, Charlottesville, VA, USA. ³Molecular and Cellular Basis of Disease Graduate Program, University of Virginia School of Medicine, Charlottesville, VA, USA.
⁴Department of Ophthalmology, University of Tsukuba, Tsukuba, Ibaraki, Japan. ⁵Department of Ophthalmology and Visual Science, Nagoya City University Graduate School of Medical Sciences, Nagoya, Japan.
⁶Aravind Eye Care System, Madurai, India. ⁷Departamento de Oftalmologia e Ciências Visuais, Escola Paulista de Medicina, Universidade Federal de São Paulo, São Paulo, Brazil. ⁸Dr. Schroff's Charity Eye Hospital, Daryaganj, Delhi, India. ⁹Department of Pathology, University of Virginia School of Medicine, Charlottesville, VA, USA.
Inflammasome agonists exacerbate experimental laser-induced choroidal angiogenesis.
Inflammasome agonist-induced laser CNV exacerbation is inflammasome-dependent.
Inflammasome-dependent macrophage migration drives CNV exacerbation and is mediated by IL-1β.

M-42
The sexual dimorphism of IgG1 anti-angiogenic activity depends on Y chromosome genes
Dionne Argyle MS1, Felipe Pereira MD2, Yosuke Nagasaka PhD3, Shao-bin Wang PhD1, Jayakrishna Ambati MD1, Bradley Gelfand PhD1
1University of Virginia, Charlottesville, VA, USA. 2Universidade Federal de Sao Paulo, Sao Paulo, Brazil. 3University of Virginia, Charlottesville, VA, USA
- The Fc fragment of all human IgG1 antibodies has angiosuppressive properties
- Research suggests that there is a sexual dimorphic response when using IgG1 antibodies in treatment
- A possible tool to explore why there is a sexual dimorphic response with IgG1 is the Y chromosome

M-43
Dab2 regulates VEGFR2 trafficking during angiogenesis in diabetes
Sudarshan Bhattacharjee PhD, Hong Chen PhD
Boston Children’s Hospital, Harvard Medical School, Boston, MA, USA
- Disabled-2 (Dab2) affects VEGFR2 trafficking, impacting angiogenesis in diabetes.
- Dab2 deficiency reduces VEGF signaling, impeding endothelial cell responses.
- Stabilizing Dab2 could enhance angiogenesis for improved wound healing in diabetes.

M-44
Cx40 suppresses sprouting angiogenesis in vitro
Edward K Looker BS1, Femke J Aan BS1, Christopher J Hatch BS2, Christopher CW Hughes PhD2, Michelle L Matter PhD1, Jennifer S Fang PhD1
1Tulane University, New Orleans, LA, USA. 2UC-Irvine, Irvine, CA, USA
- Cx40 knockdown increases sprouting angiogenesis in the in vitro Bead Assay model.
- Cx40 knockdown increases endothelial cell proliferation, but not migration.
- Cx40 regulates Cx37 expression, and Cx37 knockdown alone produces hypersprouting in the Bead Assay.

M-45
Hypoxia regulate developmental coronary angiogenesis potentially through VEGFR2- and SOX17-mediated signaling
Halie E Vitali MS1, Bryce Kuschel MS1, Chhiring Sherpa BS1, Brendan W Jones BS1, Nisha S Jacob BS1, Syeda A Madiha MS1, Samuel Elliott BS1, Eddie Dziennik BS1, Lily Kreun BS1, Cora Conatser BS1, Bhupal P Bhetwal Ph.D.2, Bikram Sharma Ph.D.1
1Ball State University, Muncie, IN, USA. 2Marian University, Indianapolis, IN, USA
- Hypoxia regulate embryonic coronary growth in mice
- Hypoxia stimulate developmental coronary growth through SOX17 mediated pathway
- Hypoxia stimulate developmental coronary growth through VEGF-R2 mediated pathway
Hemogenic Endothelium/Endothelial Diversity

**M-46**
The chromodomain helicase DNA binding protein 8, CHD8, regulates embryonic erythropoiesis through modulating DNA methylation in endothelial cells of the dorsal aorta

*Masahide Sakabe Ph.D., Nong Chen, Mark Verba, Qing Richard Lu Ph.D., Mei Xin Ph.D.*
*Cincinnati Children's Hospital Medical Center, Cincinnati, OH, USA*

- Investigate the role of CHD8 during embryonic erythropoiesis
- The genes that regulate hematopoietic stem cell differentiation was downregulated in CHD8 KO embryos
- CHD8 regulates embryonic erythropoiesis through modulating DNA methylation

**M-47**
ABCA1-dependent cholesterol efflux can regulate membrane cholesterol content in rat mesenteric artery endothelial cells

*Jacob R Anderson B.S., Nancy L Kanagy Ph.D., Laura V Gonzalez Bosc Ph.D., Jay S Naik Ph.D.*
*UNM, ABQ, NM, USA*

- Regional variation in endothelial cell membrane cholesterol exists in the mesenteric circulation
- Membrane cholesterol regulates H2S-signaling
- ABCA1-dependent cholesterol efflux participates in the regulation of membrane cholesterol content

**M-48**
Exploring the impact of sex on endothelial cell functions in vascular diseases – Insights from lineage tracing mice, cultured cells, and RNA sequencing

*Olga A Cherepanova PhD*
*Cleveland Clinic, Cleveland, OH, USA*

- Endothelial cells in atherosclerosis
- Sex differences of the vascular pathologies
- Mitochondrial functions in endothelial cells

**M-49**
Elucidating Adam10a function during the endothelial to hematopoietic transition

*Beatriz C Mercado B.S., Taylor S Turner B.S., Dionna M Kasper PhD*
*Dartmouth Geisel School of Medicine, Hanover, NH, USA*

- Hematopoietic stem cells (HSCs) are made via an endothelial to hematopoietic transition (EHT)
- The mechanisms which regulate EHT remain elusive.
- We identify the metalloprotease Adam10 as a candidate regulator of EHT

**Stem Cells/Organoids/In vitro models**

**M-50**
Modeling the 3D neurovascular unit with human-induced pluripotent stem cells

*Raphael Lis PhD*
*Weill Cornell Medicine, NYC, NY, USA*

- Current model of the BBB are not relying on bona fide endothelial cells
- Overexpression of ETS transcription factor in neuroectodermal epithelial cells -> EC fate
- Reprogrammed ECs are more sensitive to neuro stimuli and adopt a BBB identity
M-51
In vitro modeling of human vasculature using blood vessel organoids provides a robust platform for studying blood vessels in normal and pathological conditions
Ishpreet Dhillon1, Sara Pippard3, Ravenska Wagey3, Reiner Wimmer2, Josef Penninger2,3, Ryan Conder1, Allen C. Eaves1,4, Sharon A. Louis1, Valentina Marchetti1
1STEMCELL Technologies Inc., Vancouver, Canada. 2IMBA, Vienna, Austria. 3Life Sciences Institute, UBC, Vancouver, Canada. 4Terry Fox Laboratory, BC Cancer, Vancouver, Canada
- Human pluripotent stem cell derived blood vessel organoid (BVO) to model blood vasculature
- Exposure of BVO's to "diabetic" conditions induces basement membrane (BM) thickening
- Identified compounds that modulate signaling pathways involved in BM thickening in "diabetic" BVO's

M-52
Generation of functional pluripotent cell-derived brain endothelial cells for in vitro modelling of the neurovascular unit and blood-brain barrier
Tyler M Lu MS1,2, Ugur Akcan PhD3, Alexander Rhee BS1, Aomeng Cuit BS3, Sean Houghton MS1, Fuqiang Geng PhD1, Victor Martinez BS3, Jenny Xiang MD1, Irene Pedersen PhD1, Dritan Agalliu PhD3, Raphael Lis PhD1
1Weill Cornell Medicine, New York, NY, USA. 2SUNY Downstate School of Graduate Studies, Brooklyn, NY, USA. 3Columbia University Medical Center, New York, NY, USA. 4Scintillon Institute, San Diego, CA, USA
- Previous models of in vitro BBB have generated epithelial barrier forming cells rather than ECs.
- Our reprogrammed ECs can adopt a vascular phenotype in 2D and 3D.
- Cellular identity is multifaceted and in vitro models must reflect that.

M-53
Robust differentiation of human pluripotent stem cells into mural progenitor cells via transient activation of NKX3.1
Umji Lee Ph.D1,2, Yonglin Zhu Ph.D1,2, Allen Chilun Luo MS1,2, Liyan Gong Ph.D1,2, Yunhye Kim Ph.D1,2, Ke Yuan Ph.D1,2, Juan Melero-Martin Ph.D1,2
1Boston Children’s Hospital, Boston, MA, USA. 2Harvard Medical School, Boston, MA, USA
- Effective differentiation using transient activation of NKX3.1 to generate mural progenitors
- iMPCs mature into heterogenous mural cell subtypes
- Potential for modeling vascular diseases and therapeutic vascularization in-vivo

M-54
ETV2 induces transdifferentiation of stem cell-derived pancreatic islet-lineage cells toward an endothelial fate
Daniel M Tremmel PhD, Juan Melero-Martin PhD
Boston Children’s Hospital / Harvard Medical School, Boston, MA, USA
- Stem cell-derived islets lack vasculature, limiting their use to model islet function in vitro
- ETV2 rapidly and efficiently drives differentiation of stem cells toward an endothelial fate
- ETV2 rapidly and efficiently drives differentiation of stem cells toward an endothelial fate
M-55
The vascularized islet-VMO platform provides a unique ex vivo tool for studying insulin secretion and the role of leukocyte trafficking in type 1 diabetes pathogenesis
Benjamen T. ODonnell PhD, Hugh Bender PhD, Celeste N. Sanchez B.S., Sima Tahmouresie B.S., Christopher C.W. Hughes PhD
University of California Irvine, Irvine, CA, USA

- We have developed an islet micro-organ model that integrates human islets with vasculature.
- Islets demonstrate superior cell viability and glucose stimulation compared to standard models.
- Perfusable vasculature supports perfusion, extravasation, and killing by activated T-Cells.

M-56
Integrated transcriptomics maps a spatially defined adventitial progenitor niche in human blood vessel
Neelima Thottappillil Ph.D1, Yiyun Wang Ph.D1, Mario Gomez-salazar Ph.D1, Robert Tower Ph.D2, Qizhi Qin Ph.D1, Mingxin Xu Ph.D2, Ray Cheng Ph.D2, Mary Archer B.S.1, Sashank Reddy M.D.1, Kristen Broderick M.D.1, Bruno Peault Ph.D.3, Aaron W James M.D. Ph.D.1
1Johns Hopkins University, Baltimore, MD, USA. 2University of Texas Southwestern Medical Center, Dallas, USA. 3University of California Los Angeles, Los Angeles, USA

- Human vascular adventitial stem cell niche participates in vessel homeostasis and is poorly defined.
- There exists a spatially organized mesenchymal progenitor hierarchy within the adventitial niche.
- Cells with high expression of CD201(PROCR) were primitive progenitors regulated by Wnt signaling

M-57
Modeling early events in prosthetic valve endocarditis using the high-throughput organ-on-chip PREDICT96 platform with physiologically relevant fluid shear stress
Logan R Rubio MS1, Kirsty A McFarland PhD1, Maghnus O'Seaghdha PhD2, Corin Williams PhD1
1The Charles Stark Draper Laboratory, Cambridge, MA, USA. 2Suffolk University, Boston, MA, USA

- Prosthetic valve endocarditis is a rare but serious condition that currently has no good models
- Factors that affect bacterial adhesion involve complex interactions at the valve material surface
- Here we present a microphysiological model that recapitulates early events in PVE

M-58
Macrophages improve the vascularization of iPSC-derived kidney organoids in coculture
Cory P Johnson PhD1, Yulia Kiian PhD2, Ekaterina Chernobrivaia B.S.2, Sergey Tkachuk PhD2, Hannah M Somers BS1, Hermann Haller MD1,2
1MDIBL, Bar Harbor, Maine, USA. 2MHH, Hannover, Lower Saxony, Germany

- Kidney organoids do not recapitulate adult morphology and functionality in vitro
- Kidney organoids lack vascularity
- Macrophages improve kidney organoid development and vascularity in vitro
Vascular Malformations

M-59
A SOX18-mevalonate pathway axis drives vascular growth in infantile hemangioma
Joyce Bischoff Ph.D.1, Annegrete Holm M.D.2, Luke Borgelt B.S.2, Wei Heng Tan B.S.2, Jill Wylie-Sears M.A.2, Matthew S. Graus Ph.D.3, Liang Sun Ph.D.2, Mathias Francois Ph.D.3
1Boston Children’s Hospital and Harvard Medical School, Boston, MA, USA. 2Boston Children’s Hospital, Boston, MA, USA. 3Centenary Institute, Sydney, NSW, Australia
- Repurposing statins for infantile hemangioma
- SOX18-mevalonate pathway axis is central to endothelial differentiation in infantile hemangioma
- Non-beta blocker enantiomer of propranolol blocks vasculogenesis in infantile hemangioma

M-60
Recruited macrophages in GNAQ p.R183Q driven capillary malformations
Sana Nasim PhD1, Colette Bichsel PhD1, Sandra Alexandrescu MD2, Stephen Dayneka BS2, Robert Mannix PhD1, Donald Ingber MD,PhD1,3, Anna Pinto MD,PhD2, Joyce Bischoff PhD1
1Boston Childrens Hospital, Boston, MA, USA. 2Boston Childrens Hospital, Boston, MA, USA. 3Wyss Institute, Boston, MA, USA
- We found activated macrophages expressing MRC1, CD163, LYVE1, and CD68 in SWS brain vascular beds.
- Increased ICAM1 in SWS brain endothelium suggesting a mechanism for macrophage recruitment.
- In vitro showed increased leukocyte transmigration and flow-induced rolling/adhesion in EC-R183Q.

M-61
Cure HHT research network: building the roadmap to cure HHT
Cassi M Friday PhD, Nolie Krock MPH, Nicole Schaefer, Marianne S Clancy MPA
Cure HHT, Monkton, MD, USA
- Hereditary Hemorrhagic Telangiectasia
- Vascular Malformations
- Drug Discovery

M-62
The dual effects of ginsenoside Re in tumor formation and melanogenesis through microphthalmia-associated transcription factor downregulation
Su Jung Hwang Ph.D, Hye Jung Bang Bachelor, Hyo-Jong Lee Ph.D
Sungkyunkwan university, Suwon, Gyeonggi-do, Korea, Republic of
- The ginsenoside Re reduces melanin contents in zebrafish embryos and B16F10 melanomas.
- Re decreases the expression of MITF and its target genes.
- Re showed inhibitory effects on skin melanoma growth and induced tumor vascular normalization.

M-63
Development of a translational rat model of Glenn circulation and pulmonary arteriovenous malformations
Tina Wan PhD, Carol Mattern RDCS, Monica Merbach MS, Andrew D Spearman MD
Medical College of Wisconsin, Milwaukee, Wisconsin, USA
- A surgical rat model of Glenn circulation phenocopies clinical Glenn circulation.
- Pathologic intrapulmonary shunting starts within 4 weeks in Glenn circulation.
- Non-invasive echocardiography can accurately and safely monitor post-surgical blood flow in rats.
M-64
Generation and characterization of iPSC derived TIE2L914F endothelial cells for study of vascular anomalies
Bojana Lazovic1,2, Tuan Nguyen4, Franziska Kohl3, Mohammadhassan Ansarizadeh2, Miguel Carracedo Ortiz5, John Wiseman1, Prateek Singh6, Bilada Bilican3, Pratik Saxena4, Xuechong Hong1, Lauri Eklund2, Ryan Hicks1,6
1BioPharmaceuticals R&D Cell Therapy, Research and Early Development, Cardiovascular, Renal and Metabolism (CVRM), BioPharmaceuticals R&D, AstraZeneca, Gothenburg, Sweden. 2Oulu Center for Cell-Matrix Research, Biocenter Oulu and Faculty of Biochemistry and Molecular Medicine, University of Oulu, Oulu, Finland. 3Translational Genomics, Discovery Sciences, BioPharmaceuticals R&D, AstraZeneca, Gothenburg, Sweden. 4Finnadvance Ltd, Oulu, Finland. 5Research and Early Development Cardiovascular, Renal and Metabolism, BioPharmaceuticals R&D, AstraZeneca, Gothenburg, Sweden. 6School of Cardiovascular and Metabolic Medicine and Sciences, King’s College London, London, United Kingdom
- iPSC-derived locus targeted lines allows for studying mutations at an endogenous expression level.
- iPSC-derived Tie2 LF ECs showed migratory and incomplete tube formation phenotypes.
- iPSC-derived models can be used for studies of EC related diseases and/or drug screening.

M-65
Modeling and pathogenesis of KRAS and RIT1 lymphovenous disease in zebrafish
Scott Paulissen PhD, Dhyanam Shukla BS, Christopher Marshall BS, Benjamin Sempowski BS, Kristina Woodis BS, Sarah Sheppard MD/PhD
NIH, Bethesda, MD, USA
- Modeling human lymphatic anomalies in zebrafish
- Screening for drug effectiveness in patient genetic variants in model organisms
- High-resolution characterizations of disease onset and progression using imaging and NGS techniques

M-66
Therapeutically targeting proteostasis defects downstream of PI3K hyperactivation in venous and lymphatic malformations
Noa Franklin-Shapiro BS1, Ema laconetti MD1, Joseph McCarron BS1, Samantha Kaplan BS1, Meghan Perez BS1, Daniella Rogerson MD1, Averil I Clapp BS1, Hai Li PhD2, Santiago Barbar BS1, Charles Karan PhD3, June K Wu MD1, Carrie J Shawber PhD1
1Columbia University, New York, NY, USA. 2Columbia Univerisity, New York, NY, USA. 3Columbia University University, New York, NY, USA
- PI3K hyperactivation promotes overexpression and mislocalization of CD31 and VE-CADHERIN.
- VMs and LMs have proteostasis defects that can be therapeutically targeted.
- Proteasome inhibitors preferentially induce cell death of ECs with PI3K hyperactivation.

M-67
Recapitulation of the vascular phenotypes of diabetic retinopathy by endothelium-specific activation of H-Ras independent of diabetes
Qingfen Li Ph.D., Kevin Pumiglia Ph.D.
Albany Medical College, Albany, NY, USA
- Explore the role of endothelial-activated HRas in the pathologic development of retinal vasculature.
- HRAS activation leads to retinal vascular pathological development similar to diabetic retinopathy.
- Rapamycin and the ROS scavenger N-acetyl-cysteine treatment markedly reduced retinal pathology.
TUESDAY POSTERS SESSIONS

Cardiovascular Cell Biology I

T-01
Dissecting pericyte-endothelial relationships during brain angiogenesis
Cynthia U Adjeukur1,2, Sarah J Childs1,2
1Department of Biochemistry and Molecular Biology, University of Calgary, Calgary, Alberta, Canada. 2Alberta Children's Hospital Research Institute, Calgary, Alberta, Canada

- Endothelial cell signals (VEGFR, Wnt) are necessary for recruiting the correct number of pericytes.
- Recruited pericytes develop secondary processes to support the matured blood vessel.
- Brain angiogenesis and pericyte recruitment are necessary to support cognitive development.

T-02
Role of the retinoic acid in placental vascularization
Aleksandra Cwiek MS, Karen Hirschi PhD
University of Virginia, Charlottesville, VA, USA

- Placental Vascularization
- Arterial-venous specification
- Placental insufficiency

T-03
Eph-ephrin signaling couples endothelial cell sorting and arterial specification
Jonas Stewen1, Kai Kruse1, Anca T. Godoi-Filip1, Hyun-Woo Jeong1, Susanne Adams1, Frank Berkenfeld1, Martin Stehling1, Kristy Red-Horse2, Ralf H. Adams3, Mara E. Pitulescu1
1Max Planck Institute for Molecular Biomedicine, Muenster, Germany. 2Stanford University, Stanford, CA, USA

- EphB4-ephrin-B2 balance is critical for arterial specification and arteriovenous patterning
- EphB4 and ephrin-B2 regulate arterial specification directly through VEGF and Notch signaling
- EphB4-dependent EC dynamics regulation under arterial flow requires Dach1

T-04
The protective role of endothelial Ripk3 during ischemic retinopathy
Xiang Ma Ph.D, Christopher M Schafer, Courtney T Griffin Ph.D.
Oklahoma Medical Research Foundation, Oklahoma City, OK, USA

- We identified a potent anti-angiogenic protein by studying key cell death pathway components.
- RIPK3 is a multifunctional protein that is tightly regulated in pathophysiological conditions.
- We leveraged a scRNAseq dataset and validated it with endothelial-specific gene modification.

T-05
Macrophage-smooth muscle cell communication in neointima initiation
Mark C Renton PhD1, Meghan W Sedovy BS1,2, Xinyan Leng PhD3, Kailynn Robert1,3, Melissa Leaf DVM1,4, Farwah Iqbal PhD1,4, Scott R Johnstone PhD1,3,4
1. The Fralin Biomedical Research Institute at Virginia Tech Carilion, Center for Vascular and Heart Research, Roanoke, VA, USA. 2. Translational Biology, Medicine, and Health Graduate Program, Virginia Tech, Blacksburg, VA, USA. 3. Department of Biological Sciences, Virginia Tech, Blacksburg, VA, USA. 4. Virginia Tech Carilion School of Medicine, Roanoke, VA, USA

- Macrophage accumulation occurs early in vascular injury and is associated with neointimal formation.
- Macrophages and smooth muscle cells communicate directly through connexin 43 gap junctions.
- Macrophage signaling may initiate Cx43-mediated smooth muscle cell proliferation.

T-06
Hyperhomocysteinemia augments smooth muscle cell phenotypic modulation and atherosclerosis through heat shock factor 1 (HSF1) activation and increased cholesterol biosynthesis
Abhijnan Chattopadhyay PhD, Suravi Majumder PhD, Callie S Kwartler PhD, Dianna M Milewicz MD, PhD
The University of Texas Health Science Center at Houston, Houston, TX, USA
- Homocysteine activates HSF1→HMGCR→PERK signaling and phenotypic modulation in smooth muscle cells.
- Blocking HSF1, HMGCR, or PERK reverses homocysteine-induced phenotypic modulation of SMCs.
- SMC-specific Perk deletion and pravastatin treatment reduce plaques in hyperhomocysteinemic mice.

T-07
SH2 domain protein E (SHE) and ABL signaling regulate blood vessel size
SURENDRA K. ANAND Ph.D.1, Jennifer A. Schumacher Ph.D.2,3,4, Zoë A. Wright2, Diandra S. Florat MS1, Manish Dasyani Ph.D.2,1, Laurita Klimkaite3, Suman Gurung Ph.D.1, Gretchen M. Koller4, Kalia N. Aguera5, George E. Davis Ph.D.5, Saulius Sumanas Ph.D.2,3,1
1University of South Florida, Department of Pathology and Cell Biology, USF Health Heart Institute, Tampa, FL, USA. 2Cincinnati Children’s Hospital Medical Center, Division of Developmental Biology, Cincinnati, OH, USA. 3University of Cincinnati College of Medicine, Department of Pediatrics, Cincinnati, OH, USA. 4Department of Biological Sciences, Miami University, Hamilton, OH, USA. 5University of South Florida, Department of Molecular Pharmacology and Physiology, Tampa, FL, USA
- Loss of She function results in enlarged vascular lumen size and defects in blood flow.
- She acts as a negative regulator of Abl signaling.
- She regulates vascular lumen size by promoting claudin 5a (cldn5a) expression.

T-08
Androgen and mineralocorticoid receptor signaling contribute to pubertal vascular rupture in Vascular Ehlers-Danlos Syndrome (VEDS) mice
Emily Juzwiak BS1,2, Caitlin Bowen MD, Ph.D1,2, Nicole Anderson Ph.D.1,2, Anthony Zeng BS, MS1,2, Harry C Dietz MD2,2
1Department of Genetic Medicine, Johns Hopkins University, Baltimore, Maryland, USA. 2Howard Hughes Medical Institute, Chevy Chase, Maryland, USA. 3Department of Genetic Medicine, Johns Hopkins University, Baltimore, Maryland, USA
- In VEDS, males experience more pubertal arterial rupture than females.
- Blocking the androgen and mineralocorticoid receptors greatly improve survival in VEDS mice.
- Single nucleu RNAseq on VEDS aortas reveal cellular heterogeneity & crosstalk.

Lymphatic Biology and Development

T-09
Engineering biomaterials with stem cells for therapeutic lymphangiogenesis
Donny Hanjaya-Putra Ph.D.
University of Notre Dame, Notre Dame, IN, USA
- Robust differentiation of human pluripotent stem cells into lymphatic endothelial cells.
Hyaluronic acid hydrogels to control lymphatic vascular morphogenesis.
Engineered lymphatic vessels can integrate and function in vivo.

T-10
Interleukin-19 promotes lymphatic permeability and lymphangiogenesis through diverse mechanisms
Amanda M Peluzzo BS, Michael V Autieri PhD, Sheri Kelemen
Temple University LKSOM, Philadelphia, PA, USA
- Cytokine Regulation of Lymphatic Permeability
- Cytokine Regulation of Lymphangiogenesis
- Transcriptional Regulation and Lymphatic Function

T-11
Chemokine signaling regulates early lymphatic development
Meriem Bkhache, Long Do, Xiaolei Liu
Temple University, Philadelphia, USA
- Lymphatic chemokine signaling Cxcr4/cxcl12 regulates early lymphatic development
- Cxcl12/Cxcr4 is required for vegfc mediated lymphangiogenesis
- Cxcr4 activity is required for sustain lymphatic endothelial cell surface VEGFR3 abundance

T-12
Impact of Goreisan on Human Dermal Lymphatic Endothelial protein/mRNA expression
Celene Totry, Laurelis Santiago M.S., Jerome W Breslin PhD
USF, Tampa, FL, USA
- Goreisan treatment alters the HDLEC proteome.
- Mitochondrial function appears to be a target of Goreisan in HDLEC.
- Goreisan stimulates MMP3 mRNA expression in HDLEC.

T-13
β-arrestin1/2 is essential for embryonic lymphatic vessel development
Yanna Tian, Danica D Dy, Bryan Michael Kistner, Elizabeth S Douglas, Kathleen M Caron
University of North Carolina at Chapel Hill, Chapel Hill, NC, USA
- Characterization of the effect of β-arrestin1/2 deficiency in lymphatic development.
- In vivo study to investigate the role of β-arrestin1/2 in the mouse lymphangiogenesis
- Assessment of the consequences of lymphatic specific deletion of β-arrestin1/2

T-14
Elucidation of the role of ACKR3 in the lymphangiogenic response after ischemic heart injury
Laszlo Balint PhD1, Hua Zhang MD1, Amir Aghajanian MD, PhD1, 2, Shubhangi H. Patel1, Kathleen M. Caron PhD1
1Department of Cell Biology and Physiology, University of North Carolina at Chapel Hill, Chapel Hill, NC, USA.
2Department of Medicine, Division of Cardiology, University of North Carolina at Chapel Hill, Chapel Hill, NC, USA
- Characterization of the mechanisms promoting lymphangiogenic response after ischemic cardiac injury.
- Spatial characterization of ACKR3 activation after ischemic heart injury.
• Assessment of the consequences of lymphatic deletion of ACKR3 in under basal and injury conditions.

**T-15**
**Investigating the role of the microRNA miR-223 in lymphatic development**
*Carolyn L Winston B.A., Elizabeth Jones B.S., Dionna M Kasper Ph.D.*
The Geisel School of Medicine at Dartmouth, Hanover, NH, USA

- The function of miR-223 in early vascular progenitors and lymphatic endothelial cells is unknown.
- Zebrafish mutants lacking mature miR-223 show defects in forming early lymphatic structures.
- Etv2 and Prox1a, essential endothelial factors, are predicted to contain miR-223 binding sites.

**T-16**
**Exploring lymphatic system dynamics in edema formation and resolution: Lessons from zebrafish**
*Olamide Olayinka, HyunMin Jung*
University of Illinois, Chicago, IL, USA

- Developed a new zebrafish edema model by inducing osmotic stress
- Osmotic edema-induced lymphangiogenesis
- Lymphatic clearance of edema fluid

**Mechanical Forces and the Vasculature**

**T-18**
**Novel roles for centriolar protein WDR90 in endothelial cells and cardiac tissue**
*Sarah Colijn PhD, Sheng-Chih Jin PhD, Amber Stratman PhD*
Washington University, St. Louis, MO, USA

- De novo mutations in centriolar protein WDR90 have been identified as drivers of CHD
- Adult zebrafish wdr90 mutants display partial lethality, stenosis, and small hearts
- Endothelial WDR90 co-localizes with VE-cadherin and the tips of microtubules

**T-19**
**Pathophysiological hemodynamic changes in choroidal microvasculature**
*Joseph J Olivieri B.S., Bradley Gelfand Ph.D.*
University of Virginia, Charlottesville, VA, USA

- Assess the biomechanical environment of the choroid for its potential to influence AMD pathology
- Shear-responsive TM expression in the choroidal microvasculature may yield mechanistic insights
- Linking microvascular dysfunction and hallmark AMD features inspires hemodynamic-based therapeutics

**T-20**
**Hormone-free medium impairs pulmonary artery endothelial cell response to shear stress**
*Cassandra K. Conway-O'Donnell Ph.D., Christopher J. Hatch B.S., Robel K. Bekele B.S., Christopher C.W. Hughes Ph.D., Naomi C. Chesler Ph.D.*
University of California, Irvine, Irvine, CA, USA

- The absence of steroid hormones disrupts pulmonary artery endothelial cell mechanotransduction.
- 1nM 17β-estradiol does not improve pulmonary artery endothelial cell adaptation to flow.
- The absence of steroid hormones induced stress and upregulated pro-inflammatory cytokines.
Emerging Topics in Microcirculation

T-22
A novel approach to restore endothelial glycocalyx in sepsis using liposomal nanocarriers of preassembled glycocalyx, followed by quantitative analysis of endothelial glycocalyx with sidestream dark field image

Shinya Ishiko M.D., Ph.D, Ghada Ben Rahoma M.D., Ph.D, Sharath Kandhi M.D., Ph.D, An Huang M.D., Ph.D, Dong Sun M.D., Ph.D
New York Medical College, Valhalla, NY, USA

- Novel therapy using liposome to deliver preassembled glycocalyx to endothelial cells was introduced.
- Significantly impaired flow-induced dilation in septic mice was fully restored in response to LNPG.
- Chronic changes of EG and effect of LNPG can be monitored by cranial window approach.

T-23
Oxidation of low-density lipoprotein (oxLDL) by hemoglobin causes lung microvascular endothelial barrier dysfunction via lectin-like oxLDL receptor-1 (LOX-1)

Jamie E Meegan PhD, Lorraine B Ware MD, Julie A Bastarache MD
Vanderbilt University Medical Center, Nashville, TN, USA

- Cell-free hemoglobin (Hb) is a mechanistic driver of microvascular endothelial injury.
- Oxidation of LDL by hemoglobin (Hb-oxLDL) worsens lung microvascular barrier dysfunction.
- Hb-oxLDL disrupts lung microvascular endothelium through lectin-like oxLDL receptor-1 (LOX-1).

T-24
Al-enabled vascularized biological systems: Evaluation of oxygen transport from microscopic images

Jonathan Tronolone1, Abhishek Jain PhD2,3
1Texas A&M University, College Station, TX, USA. 2Texas A&M University, College Station, TX, USA. 3Houston Methodist Research Institute, Houston, TX, USA

- Machine learning enabled vascularized microphysiological systems
- Oxygen or species transport can be predicted
- Standardized analysis of microphysiological systems of the vasculature

T-25
Perivascular adipose tissue as a mediator for anticontractile activity in the coronary microcirculation

Sharanee P Sytha1, Trevor S Self2, Jeff F Bray1, Cristine L Heaps1,2
1Texas A&M University, College Station, TX, USA. 2Michael E. DeBakey Institute for Comparative Cardiovascular Science & Biomedical Devices, College Station, TX, USA

- Coronary arterioles with perivascular adipose tissue demonstrate enhanced anticontractile activity
- K+ channel activation contributes to perivascular adipose tissue-mediated anticontractile activity
- Perivascular adipose tissue mediates anticontractile activity in the swine coronary microcirculation
T-26
Predicting 3D blood flow and hematocrit distributions in microvascular networks using machine learning techniques
Saman Ebrahimi PhD, Prosenjit Bagchi PhD
Rutgers University, Piscataway, NJ, USA
• Machine learning and artificial intelligence (AI) application in microcirculation
• Coupling high-fidelity data and Machine learning for microvascular hemodynamics
• Blood flow, hematocrit and wall-shear stress prediction in microvascular networks using AI

T-27
Heparan sulfate modulation affects breast cancer cell adhesion and transmigration across in vitro blood-brain barrier (BBB)
Yunfei Li MS, Bingmei Fu Ph.D
The City College of New York, New York, NY, USA
• Heparan sulfate (HS) of endothelial cells is disrupted by VEGF which enhances HS of tumor cells
• Disrupted/enhanced HS of endothelial/tumor cells increases tumor cell adhesion/transmigration
• Endothelial HS enhancing agents disrupt HS of tumor cells and reduce their adhesion/transmigration

T-28
Neuropilin 1 (NRP1) acts as a mediator of SEMA3A chemorepulsive cues to guide vessel sprouting
Marco Spreafico¹, Elena Guzzolino¹, Francesca Fanuele¹, Carlotta Tacconi¹, Valeria Catroppa¹, Gaia Gestri², Laura Dent³, Caroline Pellet-Many⁴, Christiana Ruhrberg³, Alessandro Fantin¹
¹University of Milan, Department of Biosciences, Milan, Italy. ²UCL Department of Cell and Developmental Biology, University College London, London, United Kingdom. ³UCL Institute of Ophthalmology, University College London, London, United Kingdom. ⁴Department of Comparative Biomedical Sciences, Royal Veterinary College, London, United Kingdom
• Human endothelial cells are repelled by SEMA3A in a NRP1-dependent manner
• Zebrafish Nrp1a and Nrp1b cooperate to prevent ectopic ISV sprouting and vascular overgrowth
• Nrp1 and Sema3a cooperate to regulate ISV sprouting without affecting endothelial proliferation

T-30
Endothelial ADAM10 utilization defines a molecular pathway of vascular injury in bacterial sepsis
Danielle N Alfano MD, Mark J Miller PhD, Juliane Bubeck Wardenburg MD, PhD
Washington University School of Medicine, St Louis, MO, USA
• Lack of ADAM10 on the endothelium alone is sufficient to improve S. aureus mortality in mice.
• Endothelial injury induced by Hla-ADAM10 complex incites formation of microvascular thrombi.
• ADAM10 differentiates molecular pathways of disease among common human sepsis pathogens.

Atherosclerosis I

T-31
A new role of ATP-binding cassette B8 in preventing endothelial dysfunction and atherosclerosis
ABC8B is a key regulator of vascular health
ABC8B disruption in endothelial cells induces inflammation and plaques formation
ABC8B suppresses TGF-beta pathway activation

Identifying the novel role of IncRNA-SPANXA2-OT1 in macrophage chemotaxis through a systems-based target discovery approach
Prabhash K Jha Ph.D.1, Lucas Yuji Umesaki Itto2, Aatira Vijay Ph.D.2, Adrien Lupieri Ph.D.1, Sarvesh Chelvanambi1, Yuto Nakamura1, Miguel C Barbeiro1, Thanh-Dat Le1, Taku Kasa1, Dakota Becker-Greene1, Abhijeet R Sonawane1, Sasha A Singh1, Elena Aikawa1, Shizuka Uchida1, Masanori Aikawa1
1Brigham and Women's Hospital, Harvard Medical School, Boston, MA, USA. 2Massachusetts General Hospital, Harvard Medical School, Boston, MA, USA. 3Center for RNA Medicine, Department of Clinical Medicine, Aalborg University, Copenhagen, Denmark
- IncRNA SPANXA2-OT1 plays a key role in macrophage chemotaxis in CAD
- Co-expression network analysis for functional characterization of long noncoding RNAs
- IncRNAs can interact with DNA, RNA, miRNA, and proteins, making them potential therapeutic targets.

Endothelial deletion of Nck1 reduces atherosclerosis in ApoE−/− mice
Cyrine Ben Dhau Ph.D, Gerardo Ali Cruz Marquez, Mabruka Alfaidi PhD, Elizabeth Cockerham, Matthew Scott PhD, Brenna Pearson, Arif Yurdagul Jr PhD, Wayne Orr PhD
LSU Health Shreveport, Shreveport, LA, USA
- Endothelial-specific Nck1 deletion significantly reduced atherosclerotic plaque formation.
- Activating Transcription Factor 3 ATF3 was selected as a Nck1-selective binding partner
- Nck1 inhibition did not affect angiogenesis or limb perfusion.

Effects of activity levels on aortic calcification in hyperlipidemic mice as measured by MicroPET/MicroCT
Andy Hon, Jeffery J. Hsu MD, PhD, Angelica Zambrano, Yuxuan Xia, Mimi Lu BS, David Echeverri BS, Sophia Kalanski, Linda L. Demer MD, PhD, Yin Tintut PhD, Soban Umar Md, PhD
UCLA, Los Angeles, CA, USA
- Aortic calcium content increased significantly in both control and treadmill groups.
- Low-speed regimen reduced aortic mineral surface area and vertebral bone density in female mice.
- Low-speed regimen blunted the Western diet-induced LV hypertrophy observed in controls in male mice.

The role of EphA2 in vascular smooth muscle cell proliferation, migration, and mitogenic signaling
Matthew Scott Ph.D., Alexandra Finney Ph.D., Shantel Vital, Brenna Pearson-Gallion, Alika Shum, Wayne Orr
LSU Health Shreveport, Shreveport, LA, USA
- The EphA2 receptor is simultaneously necessary for and antagonistic to VSMC mitogenic signaling.
- EphA2 expression correlates significantly with SMC phenotypic modulation.
• EphA2 expression is sufficient to downregulate expression of SMC contractile genes.

T-36
Knockdown of AdipoR2 compromises adiponectin’s anti-inflammatory actions by mainly promoting a pro-inflammatory chemokine and cytokine secretory profile in THP-1 macrophages
Ioanna Gianopoulos MSc, Stella S. Daskalopoulou MD, MSc, DIC, PhD
Research Institute of the McGill University Health Centre (RI-MUHC), McGill University, Montreal, Quebec, Canada
• Adiponectin increases cell surface levels of AdipoR1/R2 despite AdipoR intracellular knockdown
• Loss of AdipoR2 versus AdipoR1 leads to a pro-inflammatory profile in response to adiponectin
• Selectively targeting AdipoR2 may promote plaque stabilization by reducing inflammatory mediators

Vascular Function in Health Disparities
T-37
Looking further into the significance of perfusion asymmetries in the lower limb of healthy individuals
Margarida Florindo MSc1,2, Joao Gregorio MSc, PhD1, Luis M Rodrigues MSc, PhD1
1CBIOS – Universidade Lusofona’s Research Center for Biosciences & Health Technologies, Lisboa, Lisboa, Portugal. 2ESSCVP—Department of Physiotherapy, The Portuguese Red Cross Health School, Lisboa, Lisboa, Portugal
• perfusion variations between limbs have been commonly found in healthy individuals
• to study potential relations between these limb perfusion variations and hemodynamics
• perfusion asymmetries might accelerate peripheral vascular disease if other determinants are present

T-38
Isolated human mesenteric arteries from organ donors as a model to determine arterial reactivity
Laura Hurtado Osorio Biomedical Engineering, Patricia Zamora Diaz Biomedical Engineering, Charissa Bloom BS, Jenna McQueen, Isabela Zimmermann Rollin, Vishnu Iyer, Chris Katnik, Jerome W Breslin PhD
University of South Florida, Tampa, Florida, USA
• Arteries from donated human organs remain viable and are suitable for studies of function
• Age, sex, Body Mass Index, hypertension, and diabetes did not impact arterial responses to ET-1
• Sex, Body Mass Index, hypertension, and diabetes altered responses to phenylephrine and U46619

T-39
Altered impact of sigma receptor agonists on adrenergic-induced contraction of human mesenteric arteries from donors with a history of heavy alcohol use
Patricia Zamora Diaz, Laura Hurtado Osorio, Jenna F. McQueen, Charissa Bloom, Isabela Zimmermann Rollin, Vishnu V. Iyer, Jerome W. Breslin
University of South Florida, Tampa, FL, USA
• The sigma receptor agonist PRE-084 attenuates phenylephrine-induced contraction of human arteries.
• Sigma receptor antagonists shift the phenylephrine concentration-response upward in human arteries
• Heavy alcohol use reduces the impact of sigma receptor agonists or antagonists on human arteries.
Vascular Anomalies

T-40
Ips-derived endothelial cells facilitate modeling of vascular malformations and enable drug testing
Franziska Kohl¹,², Mikel Roscales¹,², Bojana Lazovic³,², Luc Krimpenfort¹, Xuechong Hong PhD², Ryan Hicks PhD², Lars Jakobsson PhD¹, John W Wiseman PhD¹
¹Karolinska Institutet, Stockholm, Sweden. ²AstraZeneca, Gothenburg, Sweden. ³University of Oulu, Oulu, Finland
- Create cell models by knocking out ENG or ACVLR1 in iPS cells to mimic HHT
- Differentiate iPS cells to endothelial cells and characterize these cell models
- Screen a selection of drugs to identify compounds that affect KO phenotype

T-41
Expression of mutant KRAS in endothelial cells leads to hypertranscription and altered ribosome biogenesis
Negar Khosraviani¹,², Ruolin Wu¹,², Emilie Boudreau², Joshua D. Wythe³, Jason E. Fish¹,²
¹Department of Laboratory Medicine and Pathobiology, University of Toronto, Toronto, ON, Canada. ²Toronto General Hospital Research Institute, University Health Network, Toronto, ON, Canada. ³Cardiovascular Research Institute, Baylor College of Medicine, Houston, TX, USA
- Upregulation of ribosomal RNA synthesis in KRAS-G12V ECs.
- ECs with KRAS-G12V mutation undergo hypertranscription.
- Increased protein synthesis in KRAS-G12V ECs.

T-42
FLVCR2 transports choline at the Blood brain barrier
Dibyanti Mukherjee PhD¹, Rosemary Cater PhD², Ji Hyun Katie Koo B.S¹, Filippo Mancia PhD², Thomas Arnold MD¹
¹University of California, San Francisco, San Francisco, California, USA. ²Columbia University Irving Medical Center, New York, New York, USA
- Identifying Flvcr2 as a novel choline transporter at Blood Brain Barrier
- Loss of Flvcr2 causes vascular defects and hydrocephalus
- Choline transport across the blood brain barrier is required for neuro vascular development

T-43
Collagen Col22a1 functions together with Col5a1 to maintain vascular stability in zebrafish trunk
Vishal Y Mardhekar M. Pharm, Suman Gurung PhD, Kaitlin Ferrari MS, Saulius Sumanas PhD
USF Morsani College of Medicine, Tampa, Florida, USA
- Role of Col22a1 in Zebrafish vascular stability.
- Collagen subtypes associated with vascular aneurysms
- Studying vascular abnormalities

T-44
Ras isoform specificity in endothelial cell function and their role in dysfunctional morphogenesis
Samantha King, Kevin Pumiglia PhD
Albany Medical College, Albany, NY, USA
• Ras isoforms (HRas and KRas) have distinct endothelial signaling and functional outputs.
• KRas protein expression levels are diminished relative to HRas through codon bias.
• Optimizing codons of KRas does not account for differences in AKT signaling or proliferation.

Endothelium in Health and Disease I

T-46
Endothelial PHD2 deficiency promotes angiotensin-II-induced arterial stiffness and coronary microvascular dysfunction

Heng Zeng¹, Jian-Xiong Chen²
¹University of Mississippi Medical Center, Jackson, MS, USA. ²University of Mississippi Medical Center, Jackson, MS, USA

• Prolyl hydroxylase protein-2
• Endothelial specific knockout mouse
• Coronary microvascular dysfunction

T-47
Orchestration of endothelial growth and osteogenesis

Yang Yong, Yan Zhao, Xiuju Wu, Li Zhang, Xinjiang Cai, Kristina Bostrom, Yucheng Yao
UCLA, Los Angeles, California, USA

• Endothelial cells and osteogenic cells alternately showed the potential mutual regulation in femur.
• In skull, endothelial and osteogenic cells followed similar patterns without alternation
• The orchestration of endothelial and bone cells is different between long and flat bones formation.

T-48
Regulating the cell shift of endothelial cell-like myofibroblasts in pulmonary fibrosis

Xiju Wu M.D., Ph.D.¹, Daoqin Zhang Ph.D.², Li Zhang Ph.D.¹, Xinjiang Cai M.D., Ph.D.¹, Yan Zhao Ph.D.¹, John A. Belperio M.D.³, Kristina I. Boström M.D., Ph.D.¹, Yucheng Yao M.D., Ph.D.¹
¹Division of Cardiology, David Geffen School of Medicine, UCLA, Los Angeles, CA, USA. ²Department of Pediatrics, Stanford University, Stanford, CA, USA. ³Division of Pulmonary and Critical Care Medicine, Clinical Immunology, and Allergy, David Geffen School of Medicine, UCLA, Los Angeles, CA, USA

• Identify a previously unknown cell population that contributes myofibroblasts to pulmonary fibrosis.
• A small molecule is discovered to redirect myofibroblasts and reduces pulmonary fibrosis.
• COVID-19 infection drives endothelial-lineage cells towards myofibroblasts in pulmonary fibrosis.

T-49
Endothelial reprogramming and enhanced TGF-β signaling persist after completion of chemotherapy

Kass Sjostrom BS, Shixin Tao PhD, Nataliya Kibiryeva MD, Melissa Cobb MS, Eugene Konorev MD, PhD
Kansas City University, Kansas City, MO, USA

• Doxorubicin treatment causes long-term changes in endothelial cell expression
• Long-term endothelial cell changes are likely mediated by the canonical TGF-β pathway.
• Suppressing the canonical TGF-β pathway may alleviate cardiovascular damage caused by doxorubicin.

T-50
GSK3β inhibition reduced vascular calcification of Ins2Akita/+ and Apoe−/− mice
Yan Zhao PhD, Xinjiang Cai MD, PhD, Xiuju Wu MD, PhD, Yang Yang MD, PhD, Li Zhang PhD, Kristina Boström MD, PhD, Yucheng Yao MD, PhD
David Geffen School of Medicine at UCLA, Los Angeles, California, USA

- GSK3b inhibition reduces vascular calcification in the models of diabetes and atherosclerosis.
- GSK3b inhibition redirects endothelial cell (EC)-derived osteoblast-like cells back to EC lineage.
- GSK3b inhibition induced b-catenin and reduced SMAD1 in diabetic arteries and atherosclerosis.

T-51
The RNA m6A demethylase, FTO, regulates vascular remodeling in the pathogenesis of pulmonary arterial hypertension
Jingbo Dai, Northwestern University, Chicago, IL, USA

- FTO expression level is markedly elevated in the endothelial cells of idiopathic PAH patients.
- The Tie2-Cre-Fto mice exhibited inhibited PH under hypoxia treatment.
- MCT-rat PAH model showed attenuated PAH phenotype with pharmacological inhibition of FTO.

T-52
Infection and pulmonary pathology in a novel model of COVID-19
Cadence Lee M.S.1,2,3, Rachel Carley PharmD1,2,3, Sheila Sharma M.S.1,2,3, Julia Pierce B.S.1,2,3, Crystal Parry B.S.1,2,3, Elizabeth Amelotte1,2,3, Celia Butler MPH2,3, Chris Mantsoura PhD1,2,3, Gaurav Choudhary MD1,2,3, Alan R. Morrison MD/PhD1,2,3
1Brown University, Providence, RI, USA. 2Ocean State Research Institute, Providence, RI, USA. 3VA Providence, Providence, RI, USA

- novel model of COVID-19
- Histopathologic analysis of COVID-19
- ACE2 expression in regards to SARS-CoV-2 infection

Signaling

T-54
Matrix Gla protein directs cell transitions in brown adipose development
Li Zhang, Xinjiang Cai, Yan Zhao, xiuju wu, yucheng yao, Kristina l. Boström
UCLA, Los Angeles, California, USA

- MGP is expressed in endothelial and adipose progenitor cells.
- EC-derived MGP prevented the triggering of white adipogenesis in the perivascular region of BAT.
- MGP plays a complex role in the transitioning of cell populations during brown adipogenesis

T-55
The in vitro and in vivo analysis of the mechanism underlying the accumulation of NOTCH3 CADASIL mutant protein
T-56
Role of neural EGR1 in blood flow recovery after hindlimb ischemia
Gaganpreet Kaur PhD1,2, Amada Caliz MS1,2, Heather Learnard MS3, Hyung-Jin Yoo MS1,2, Soonsang Yoon PhD1,2, Shashi Kant PhD1,2, John F. Keaney MD1,2
1Brigham and Women’s Hospital, Boston, MA, USA. 2Harvard Medical School, Boston, MA, USA. 3University of Massachusetts, Worcester, MA, USA
- Neuronal EGR1 regulates the expression of different growth factors like VEGFα, PGF, and HBEGF.
- Regulation of growth factors secretion by EGR1 can effect endothelial cell function.
- The overexpression of EGR1 increases blood flow recovery after mouse hindlimb ischemia.

T-57
Dissecting the regulatory elements of microRNA-223 in zebrafish vascular and blood development
Elizabeth A Jones BS, Dionna M Kaser PhD
Dartmouth College, Hanover, NH, USA
- microRNA (miR)-223 is a key regulator of endothelial cell fate and hematopoiesis.
- miR-223 is expressed in different tissues at different times in hematovascular development.
- The promoter has binding sites for vascular and hematopoietic regulatory factors.

T-58
Roles of Wnt5a signaling in inflammation and cancer are dependent on tissue context
Ramiro Malgar MD1, Karen Coschigano Ph.D.1, Alyx Hazen BS1, Brakstin Hockley1, Ian Ackers D.O. Ph.D.2
1Biomedical Sciences, Ohio University, Athens, OH, USA. 2Department of Physical Medicine and Rehabilitation, Sparrow Hospital, Michigan State University, Lansing, Michigan, USA
- Wnt5a signaling, a complex pathway activated in inflammation and cancer.
- Could Wnt5a be one of the links responsible for the relationship inflammation/cancer?
- Wnt5a modulates innate immunity in tissue microenvironment.

T-60
Thoracic aortic aneurysms in Marfan Syndrome: from elastic fibers to integrin αvβ8
Soheila Ali Akbari Ghavimi PhD, Ryan Martinez, Mariam Kerolos, Amarri Harrison, Kim de la Cruz, Marie Billaud
Brigham and Women’s Hospital, Boston, MA, USA
- Marfan syndrome caused by mutation of FBN1 and shows overactivity of TGFβ.
Aortic tissue from Marfan Syndrome patients had higher expression of MMP14 and integrin αV and β8.

Targeting integrin αVβ8 offer potential to develop therapeutics to prevent aneurysms in MFS.

Aging, Metabolism and the Vasculature I

T-61
Endothelial Cpt1a reduction causes hyperoxia-induced pulmonary vascular remodeling by upregulating EndoMT in neonates
Katy Hegarty BS1, Jason L Chang BS1, Julie Braza BS2, Gaurav Chaudhary MD2, Phyllis A Dennery MD1, Hongwei Yao PhD1
1Brown University, Providence, RI, USA. 2Providence VA Medical Center, Providence, RI, USA
- Endothelial Cpt1a reduction promotes pulmonary vascular remodeling by upregulating EndoMT.
- Enhancing fatty acid oxidation inhibits neonatal hyperoxia-induced pulmonary vascular remodeling.
- Blocking EndoMT prevents neonatal hyperoxia-induced pulmonary vascular remodeling.

T-62
Sex hormones may be implicated in sex-specific differences in carotid atherosclerotic plaque instability
Karina Gasbarrino PhD1, Edward Daly MSc2, Huaien Zheng MD, PhD1, Ana Carolina Alves dos Santos PhD3, Dajana Vuckovic PhD3, Stella S. Daskalopoulou MD PhD1
1Vascular Health Unit, Research Institute of McGill University Health Centre, Montreal, Quebec, Canada. 2Clinical Proteomics and Mass Spectrometry, Research Institute of McGill University Health Centre, Montreal, Quebec, Canada. 3Department of Chemistry and Biochemistry, Concordia University, Montreal, Quebec, Canada
- Sex differences exist in plaque composition, where men have more unstable plaques than women.
- High testosterone and low estradiol levels are associated with greater plaque instability in men.
- Sex hormones have been detected for the first time in stable and unstable plaques.

T-63
Function of BRI3 in CD16 monocytes in Parkinson’s disease
Kathleen C Paul1, Owen M Wilkins Ph.D., Taylor S Turner, Karl E Biggs, Faith Anderson Ph.D., Stephen L Lee M.D., Ph.D., Fred W Killing Ph.D., Dionna M Kasper Ph.D., Matthew C Havrda Ph.D.
Geisel School of Medicine at Dartmouth, Hanover, NH, USA
- Single cell RNA sequencing of PBMCs from Parkinson’s patients reveals PD-specific transcriptome.
- Brain Protein 13 is differentially expressed in PD CD16 monocytes, but function in PD is unknown.
- Monocyte-like cell line models and zebrafish inflammation models will elucidate BRI3 function.

T-64
The effects of ageing to the cardiac microvasculature
Guillermo Luxán1,2,3, Anita Tamiato1,2,3, Colin Bodemer1,2,3, Timm Winkelmeier1, Mariana Shumliakivska1,2,3, Lukas S Tombor1,2,3, Ariane Fischer1, Marion Muhly-Reinholz1, Büşra N Toğru1, Mariano Ruz Jurado1,2,3, Maximilian Merten1,2,3, Jessica Neitz1, Stefan Günther1,2,3, David John1,2,3, Nina Wettschureck1,2,3, Stefanie Dimmeler1,2,3
1. Institute of Cardiovascular Regeneration, Center of Molecular Medicine, Goethe University Frankfurt, Frankfurt am Main, Germany. 2. German Center for Cardiovascular Research DZHK, Frankfurt am Main, Germany. 3. Cardiopulmonary Institute, Goethe University Frankfurt, Frankfurt am Main, Germany. 4. Max Planck Institute for Heart and Lung Research, Bioinformatics and Deep Sequencing Platform, Bad Nauheim, Germany. 5. Max Planck Institute for Heart and Lung Research, Department of Pharmacology, Bad Nauheim, Germany
Cardiovascular ageing compromises the microvasculature in the heart
Age related deregulation of pericyte genes cause cardiac dysfunction and fibrosis in the heart
Endothelial extracellular matrix is affected in ageing and leads to a pro-inflammatory phenotype

T-66
High levels of O-GlcNAc induce endothelial senescence and vascular dysfunction in a premature vascular aging model
Paula Rodrigues de Barros PhD candidate1,2, Tiago Januario da Costa PhD2, Daniel Rodrigues PhD student1, Carina Amarante Pedersoli Masters student1, Jose Teles de Oliveira Neto PhD student1, Jeimison Duarte dos Santos PhD1, Laena Pernomian PhD2, Camilla Ferreira Wenceslau PhD2, Rita de Cassia Aleixo Tostes Passaglia PhD1
1Universidade de Sao Paulo, Ribeirao Preto, SP, Brazil. 2University of South Carolina, Columbia, SC, USA
• High levels of O-GlcNAcylation in endothelial cells induce senescence and vascular dysfunction
• High levels of O-GlcNAcylation modulate ECs oxidative-stress induced senescence and cell function
• Cycling depend kinase inhibitors activation mediates O-GlcNAcylation induced senescence

T-67
Mechanisms of age-dependent decline in angiogenesis
Yesenia G. Gomez, Tendai Hunyenyiwa, Priscilla Kyi, Tadanori Mammoto, Akiko Mammoto
Medical College of Wisconsin, Wauwatosa, Wisconsin, USA
• Angiogenesis is inhibited in aged human adipose endothelial cells.
• The levels of Y-RNA and RNA binding proteins are altered in aged human endothelial cells.
• Y-RNA in EVs may contribute to the age-dependent inhibition of angiogenesis.

WEDNESDAY POSTERS SESSIONS

Atherosclerosis II

W-01
A Rac1-NF-kB transcriptional axis regulates IL-1β during atherosclerotic calcification
Rachel Carley PharmD1,2, Cadence Lee ScM1,2, Chris Mantsounga PhD1,2, Joshua Berus PhD1,2, Olivia Caballero BS1,2, Crystal Parry BS1,2, Julia Pierce AB1,2, Elizabeth Amelotte1,2, Celia Butler MS1,2, Gaurav Choudhary MD1,2, Alan R Morrison MD/PhD1,2
1Brown University, Providence, RI, USA. 2Ocean State Research Institute Inc at Providence VA Medical Center, Providence, RI, USA
• Rac1 and NF-kB play a role in regulating IL-1B
• Rac1 and NF-kB play a role in regulating atherosclerotic calcification
• Rac1 and NF-kB regulate IL-1B and vascular calcification
W-03
Inflammation targeting contrast agent for MR imaging of atherosclerosis
Joshua Rousseau, Ting Yun Wang, Hui-Chun Huang, Kuei-Chun Wang
Arizona State University, Tempe, Arizona, USA
- Targeted MRI contrast agents enhances precise diagnosis of atherosclerosis
- Cloaking nanoparticles with monocyte membrane enables inflammation targeting
- Biomimetic nanoparticles encapsulating MRI contrast agent allows detection of plaque deposition

W-04
Nanoparticles targeting YAP/TAZ dysregulation in atherosclerosis: impact on vascular cells and macrophages
Ting-Yun Wang, Hui-Chun Huang, Joshua Rousseau, Kuei-Chun Wang
ASU, Tempe, Arizona, USA
- Developing a lesion-targeted, pathway-specific nanodrug for atherosclerosis treatment
- Unraveling pharmacological effects of the targeted nanodrug at single-cell level
- Investigating pharmacological mechanisms in vascular and immune cells

W-05
Novel role of endothelial CD47 in the regulation of pathogenesis of atherosclerosis
Bandana Singh PhD, Kui Cui, Qianman Peng, Kathryn Li, Bo Zhu, Sudarshan Bhattacharjee, Daniel Osorio, Beibei Wang, Yunzhou Dong, Donghai Wang, Yao Wei Lu, Hao Wu, Scott Wong, Douglas B. Cowan Cowan, Hong Chen
Harvard Medical School, Boston, MA, USA
- Deletion of endothelial cell-specific CD47 protects from atherosclerosis progression.
- Deletion of endothelial cell-specific CD47 promotes efferocytosis of dying cells.
- In the athero-CD47iECKO group, the subpopulation of VSMCs decreased, while EC increased.

W-06
The role of endothelial microRNA-33 in atherosclerosis
Kathryn Citrin, Yan Huang, Alberto Canfran-Duque, Xinbo Zhang, Carlos Fernandez-Hernando, Yajaira Suarez
Yale University, New Haven, CT, USA
- Endothelial lipid metabolism regulates pro-atherogenic EC functions
- The EC-specific role of miR-33, a regulator of lipid metabolism, has not been extensively explored
- EC-specific miR-33 knockout does not significantly regulate atherosclerosis progression

Development
W-07
Characterizing the function of klhl4 during vascular development in zebrafish (Danio rerio)
Kaitlin A Ferrari M.S., Suman Gurung PhD, Saulius Sumanas PhD
University of South Florida Morsani College of Medicine, Tampa, FL, USA
- KLHL4 regulates vascular development
- Genetic pathways governing vasculogenesis and angiogenesis
- Vascular defects result from KLHL4 mutation
ETV2/ER71 regulates hematovascular lineage generation and vascularization through an H3K9 demethylase, KDM4A
Dong Hun Lee¹, Min Seong Kim², Raham Lee³, Heesang Song³, Joo Kyung Kim¹, Bum-Yong Kang¹, Karl Agger⁵, Kristian Helin⁶, Changwon Park³,¹
¹Emory University School of Medicine, Atlanta, USA. ²LSU Health Science Center, Shreveport, LA, USA. ³LSU Health Science Center, Shreveport, USA. ⁴Chosun University School of Medicine, Gwangju, Korea, Republic of. ⁵University of Copenhagen, Copenhagen, Denmark. ⁶Memorial Sloan Kettering Cancer Center, New York, USA
- Interaction between ETV2 and KDM4A in differentiating mouse embryonic stem cells
- KDM4A plays an important role in the ETV2-regulated gene expression.
- The novel function of the interaction between ETV2 and KDM4A in neovascularization.

ETS transcription factor flI1b is required for hematopoietic development in zebrafish
Valentina Laverde M.S., Saulius Sumonas PhD
University of South Florida, Tampa, Fl, USA
- Specification of vascular endothelial and hematopoietic cells incompletely understood in zebrafish.
- Zebrafish paralog flI1b functions partially redundantly with ETS transcription factor Etv2/Etsrp.
- Role of FLI1 paralog flI1b in zebrafish embryonic hematopoiesis.

A system-scale approach to uncovering isoforms contributing to endothelial cell differentiation
Madison M. Mehlferber¹, Vasilii Pavelko¹, Jerryd Meade¹, Erin D. Jeffery¹, David Wissel², Elizabeth A. Nelson¹, Karen K. Hirschi¹, Gloria M. Sheynkman¹
¹University of Virginia, Charlottesville, VA, USA. ²University of Zurich, Zurich, CH, Switzerland
- A temporal representation of isoforms associated with endothelial cell development
- System-scale approach integrating long-read sequencing and mass-spectrometry
- Stem cell models used to recapitulate development of endothelial cells and analyze splicing patterns

Plasticity of the Vasculature

PRDM16 suppresses synthetic smooth muscle cell phenotypic modulation in atherosclerosis
Josephine M E Tan PhD, Patrick Seale PhD
University of Pennsylvania, Philadelphia, PA, USA
- PRDM16 is a CAD disease risk gene that regulates vascular plasticity in atherosclerosis
- Enhanced synthetic SMC modulation caused by PRDM16-KO changes lesion composition and structure
- PRDM16 suppresses fibrotic genes associated with synthetic SMC phenotypic switching
Endothelial senescence mediates hypoxia-induced vascular remodeling in the lung

Priscilla Kyi, Tendai Hunyenyiwa, Tadanori Mamamoto, Akiko Mamamoto
Medical College of Wisconsin, Milwaukee, WI, USA
- Yap1 activity and EC senescence are stimulated in the pulmonary hypertension (PH) patient lungs.
- Upregulation of PDGFB in PH patient lungs is inhibited by knockdown of p16INK4A or YAP1.
- EC senescence mediates hypoxia-induced vascular remodeling in the lungs via YAP1-PDGFB signaling.

W-15
Mitochondrial complex i promotes vascular smooth muscle cell proliferation and migration

Alishba Maira PhD Candidate, Dario F Riascos Bernal MD PhD, Nicholas E S Sibinga MD
Albert Einstein College of Medicine, Bronx, NY, USA
- Mitochondrial respiration in smooth muscle cell biology.
- Mitochondrial dynamic in vascular smooth muscle cell plasticity and vascular remodeling
- The roles of mitochondrial Complex I in vascular smooth muscle cell proliferation and migration.

W-16
Coronary microvascular adaptation to left ventricular inflow obstruction in the late gestation fetal lamb

Matthew W Hagen PhD1, Samantha Louey PhD2, Sarah M Alaniz RVT2, Jonathan R Lindner MD1,2, George Giraud MD PhD2,3, Sonnet S Jonker PhD2
1Oregon Health & Science University, Portland, Oregon, USA. 2University of Virginia, Charlottesville, Virginia, USA. 3Portland VA Medical Center, Portland, Oregon, USA
- We developed a novel large-animal model of late gestation left ventricular inflow obstruction.
- Left ventricular inflow obstruction increases coronary microvascular perfusion of both ventricles.
- RV microvascular perfusion increases to a greater degree than LV following LV inflow obstruction.

 Adv. Engineering Models & Imaging Techniques

W-17
Blood science and angiogenesis-enabled Ovarian TME-Chips determine mechanisms and therapeutics against antiangiogenic escape

Lopamudra D. Ghosh PhD1, Abhishek Jain PhD1,2
1Texas A&M University, College Station, TX, USA. 2Houston Methodist Research Institute, Houston, TX, USA
- Blood and angiogenesis-enabled human tumor microenvironment chip
- The spatiotemporal dynamics of angiogenesis in TME is regulated by blood platelets
- Platelets contribute to antiangiogenic escape

W-19
Differential-geometric model of red blood cell equilibrium shapes to investigate skeletal muscle microcirculatory regulation
Red blood cell shape is intimately connected to muscle oxygen regulation. Condensed matter theoretical models explain red blood cell convexity/concavity. Condensed matter theory/geometry models can explain how red blood cells regulate blood flow.

**Inflammation**

W-20

**Vascular inflammation in neuropsychiatric PASC**

*Lindsay S McAlpine M.D.*, *Hailey Reisert*, *Allison Nelson*, *Jennifer Chiarella*, *Shelli Farhadian M.D.*, *Serena Spudich M.D.*

1Yale University, New Haven, CT, USA. 2Mt. Sinai Hospital, New York, NY, USA

- Individuals with neuro-PASC demonstrate persistent systemic inflammation.
- Individuals with neuro-PASC demonstrate persistent vascular inflammation.
- Individuals with neuro-PASC may be at risk of accelerated atherosclerosis.

W-21

**Edn1 is a disease-specific marker in endothelial cells during atherosclerosis**

*Danica M Dy BS*, *Thiel Lehman PhD*, *Jessica Lin*, *Rob Wirka MD*

UNC-Chapel Hill, Chapel Hill, NC, USA

- A specific population of ECs may undergo EndMT, contributing to the progression of atherosclerosis
- A disease-specific population of ECs are characterized by expression of Edn1
- Pathway analysis predicts that EC transition is driven by EndMT-associated transcription regulators

W-22

**VEGF-D improves vascular integrity and resolves inflammation in the lungs**

*Yifan Yuan*, *Lokesh K Sharma PhD*, *Wenwen Tang PhD*, *Micha Sam B Raredon MD, PhD*, *Dianqing (Dan) Wu PhD*, *Laura E Niklason MD, PhD*, *Naftali Kaminski MD*  

1University of Maryland, Baltimore, MD, USA. 2Yale University, New Haven, CT, USA. 3Humacyte Global, Inc, Durham, NC, USA

- scRNAseq data reveals a strong VEGFD-VEGFR2 interaction within the human lung microvascular niche.
- VEGF-D improves vascular integrity and reduces pro-inflammatory response.
- VEGF-D regulates vascular integrity through VEGFR-2- and ROCK-dependent signalling.

**Aging, Metabolism and the Vasculature II**

W-24

**Aging is associated with organ-specific alterations in level and expression pattern of von Willebrand factor**

*Parnian Alavi*, *Douglas Brown*, *John Lewis*, *Stephane Bourque*, *Nadia Jahroudi*

1Department of Medicine, University of Alberta, Edmonton, Alberta, Canada. 2Department of Oncology, University of Alberta, Edmonton, Alberta, Canada. 3Department of Anesthesiology & Pain Medicine, University of Alberta, Edmonton, Alberta, Canada

- VWF mRNA and protein are upregulated in aged mice in an organ-specific manner.
- The increasing proportion of microvessels exhibit VWF expression in organs of aged mice.
- PS3 knockdown abolished the aged- induced VWF upregulation in endothelial cells.
W-25
Age-related increases in DNA methylation impair HuR-mediated VEGF-A mRNA stability and consequent inflammatory arteriogenesis
Christopher Mantsounga PhD1, Olivya Caballero BS1, Sheila Sharma MSc2, Cadence Lee MSc1, Rachel Carley PharmD2, Crystal Parry BS2, Julia Pierce BS2, Celia Butler MPH2, Elizabeth Amello Σ2, Gaurav Choudhary MD2, Alan Morrison MD, PhD2
1Brown University, Providence, Rhode Island, USA. 2Brown University, Providence, RI, USA
- Aging is associated with impaired inflammatory arteriogenesis in response to vascular injury.
- Reduced arteriogenesis was related to decreased muscle tissue VEGF-A and VEGF 165A levels.
- BMDMs from aged mice revealed decreased VEGF-A165a mRNA half-life, and VEGF-A associated with Hur.

W-26
Targeting the microvasculature to treat diabetes-induced heart failure with preserved ejection fraction
Cori Lau1,2, Kai Ellis1,3, Rathnakumar Kumaragurubaran2, Garry Yu2, Lijun Chi2, Paul Delgado-Olguin1, Michael D Wilson2, Jason E Fish2
1University of Toronto, Toronto, ON, Canada. 2Toronto General Hospital Research Institute, Toronto, ON, Canada. 3SickKids Research Institute, Toronto, ON, Canada
- Transcriptome and chromatin landscape of cardiac endothelial cells in T2D mice
- Transcriptome and chromatin landscape of cardiac endothelial cells in SGLT2 inhibitor-treated mice
- Microvascular analysis in the hearts of T2D mice and SGLT2 inhibitor-treated mice

W-27
Direct heterocellular communication between capillary adipose endothelium and adipocytes can regulate lipid handling
Melissa A Luse MS1, Luke S Dunaway PhD1, Meghan Sedovy2, Scott R Johnstone PhD2, Brant E Isakson PhD1
1University of Virginia, Charlottesville, VA, USA. 2Virginia Tech, Roanoke, VA, USA
- Adipose is enriched with capillary endothelial cells which make contact with adipocytes.
- Contact between capillary adipose endothelial cells dictates gene expression in both cell types.
- Connexin 43 facilities contact between adipocytes and capillary adipose endothelial cells.

W-28
Cav1 inhibition of eNOS regulates Cd36 mediated lipid uptake in endothelium
Melissa A Luse MS1, Wyatt Schug BS1, Luke S Dunaway PhD1, Skylar Loeb BS1, Rachel Tessema BS1, Miriam Cortese-Krott PhD2, Brant E Isakson PhD1
1University of Virginia, Charlottesville, VA, USA. 2Heinrich Heine University, Dusseldorf, Germany
- Endothelial Caveolin-1 plays a critical role in lipid uptake.
- Cd36 is located within endothelial caveolae and facilitate lipid uptake in vitro and in vivo.
- Nitrosylation of Cd36 by nitric oxide inhibits the transport of fatty acids into the endothelium.

W-29
Therapeutic reactivation of aging-impaired endothelial regeneration and vascular repair for treatment of acute respiratory distress syndrome in elderly patients via repurposing decitabine or endothelium-targeted nanoparticle delivery of FoxM1

Xianming Zhang PhD1, Narsa Machireddy PhD1, Colin E Evans PhD1, David Wu MD2, Youyang Zhao PhD1
1Lurie Children’s Hospital of Chicago and Northwestern University Feinberg School of Medicine, Chicago, IL, USA.
2University of Chicago, Chicago, IL, USA
• resident endothelial cell proliferation is the intrinsic mechanism of endothelial regeneration
• Aging impairs the intrinsic endothelial regenerative program via hypermethylation of Foxm1 promoter
• Decitabine treatment or nanoparticle delivery of FoxM1 reactivates endothelial regeneration in aging

W-30
Differences in body composition and lipid profile between vegetarians and omnivorous: Relationship with 10-year cardiovascular risk
Cintia Ferreira-Pego MSc, PhD, Tatiana Fontes MSc, Sofia Lopes MSc, Regina Menezes MSc, PhD, Luis Monteiro Rodrigues MSc, PhD
CBIOS – Universidade Lusófona’s Research Center for Biosciences & Health Technologies, Lisboa, Lisboa, Portugal
• To provide evidence on the impact of vegetarian versus omnivore dietary regimens on health
• To identify potential related differences regarding body composition cardiometabolic markers
• To compare cardiovascular risk of participants from both dietary regimens

Neurovascular Unit and Disease

W-31
Studying Erythromelalgia through the laser Doppler flowmetry perfusion components and skin temperature
Joana Caetano MD,MSc1,2,3, Hugo T.D. Ferreira MD,PhD4, Clemente Rocha Msc, PhD1, Sergio F. Andrade MSc, PhD1, Jose D. Alves MD, PhD1,2,3, Luis M Rodrigues MSc, PhD1
1CBIOS - Research Center for Biosciences and Health Technologies, Universidade Lusófona, Lisboa, Lisboa, Portugal. 2Immunomediated Systemic Diseases, Medicina IV Hospital Fernando Fonseca, Amadora, Lisboa, Portugal. 3Nova Medical School, Lisboa, Lisboa, Portugal. 4Institute of Biophysics and Biomedical Engineering, Faculty of Sciences - University of Lisbon, Lisboa, Lisboa, Portugal
• To better characterize Erythromelalgia a still poorly understood disease
• To apply the Wavelet transform to the LDF signals obtained under challenge
• To prove the concept after testing outcomes following a new therapeutical approach

W-32
Development of stem cell-derived microfluidic models of the blood-brain barrier in Alzheimer’s disease
Lily E Takeuchi MSc1,2, Craig A Simmons PhD1,2,3
1Institute of Biomedical Engineering, University of Toronto, Toronto, Ontario, Canada. 2Translational Biology and Engineering Program, Ted Rogers Centre for Heart Research, Toronto, Ontario, Canada. 3Department of Mechanical and Industrial Engineering, University of Toronto, Toronto, Ontario, Canada
• There is an unmet need for physiological models of the blood-brain barrier in Alzheimer’s disease.
• We demonstrate functional impairment to iPSC-derived brain endothelial cells from a patient with AD.
• Shear stress impairs immune activation of the cerebral vasculature in Alzheimer’s disease.
W-33
Circulating extracellular vesicles from mice with chronic cerebral hypoperfusion perturb brain endothelial cell function
Brandon Kirby, Audrey Plumtree, Duaa Dakhilallah PhD, Ekaterina Weil PhD, Candice M Brown PhD
West Virginia University School of Medicine, Morgantown, WV, USA
- Vascular contributions to cognitive impairment and dementia results from extensive vascular disease.
- Mechanisms underlying the pathophysiology of VCID are not well-understood.
- Plasma-derived extracellular vesicles may be a key modulator in the pathophysiology of VCID.

W-34
Activation of Wnt/β-catenin signaling in radial glial cells disrupts BBB development
Dylan J Sebo B.S., Audrey R Fetsko B.S., Michael R Taylor PhD
University of Wisconsin - Madison, Madison, WI, USA
- Wnt signaling works cell non-autonomously within the NVU to inhibit BBB development.
- Wnt and Vgf ligands released from radial glial cells, are sensitive to Wnt signaling.
- Wnt signaling within radial glial cells must be finely controlled for proper CNS and BBB development.

W-35
Reduced endothelial levels of splice factor TDP-43 associated with ALS/FTD- mutations TARDBP348C causes hallmarks of FTD in an animal model
Ashok Cheemala PhD1, Amy Kimble B.A.B.1, Jordan Tyburski B.S.1, Melissa Murphy B.S.1, Evan Jellison PhD2, Bo Reese PhD3, Xiangyou Hu PhD4, Rigiang Yan PhD4, Patrick Murphy PhD1
1University of Connecticut Medical School, Center for Vascular Biology, Farmington, Connecticut, USA. 2University of Connecticut Medical School, Department of Immunology, Farmington, Connecticut, USA. 3University of Connecticut, Center for Genome Innovation, New Britain, Connecticut, USA. 4University of Connecticut Medical School, Department of Neuroscience, Farmington, Connecticut, USA
- ALS/FTD-associated mutations in TDP-43 reduce nuclear levels in brain ECs as well as neurons
- Reduction or endothelial-specific deletion of TDP-43 disrupts RNA splicing, impairs EC-junctions
- Brain EC TDP-43 deletion mimics ALS/FTD: BBB loss, fibrin, glial activation, and cognitive issues

W-36
Wnt-directed specialization of endothelial colony forming cells to generate a perfusible blood brain barrier microphysiological system
Makena L Ewald, Christopher J Hatch, Amber R Keith, Sarah J Hernandez PhD, Duc TT Phan PhD, Leslie M Thompson PhD, Christopher CW Hughes PhD
University of California, Irvine, Irvine, Ca, USA
- Advances in CNS related disease treatments require advanced human-derived in vitro BBB models.
- Endothelial cells specialize into organotypic vasculature based on environmental cues.
- We developed a blood brain barrier model by mimicking natural developmental processes in vitro

W-37
Netrin-1 binding to Unc5B regulates BRB integrity via Norrin/β-catenin
Jessica Furtado BS1, Kevin Boyé PhD2, Anne Eichmann PhD3
1Yale University, New Haven, CT, USA. 2Inserm, Paris, France. 3Yale University, New Haven, USA
- Unc5B regulates blood retina barrier permeability.
- Netrin1 binding to Unc5B regulates the Wnt-related Norrin/β-catenin signaling via pLRP5.
• Endothelial cell and pericyte Unc5B may have opposing roles.

3D Imaging and Organotypic Vasculature

W-38
Creating vascularized gut tube organoid to recapitulate organotypic endothelium and mesenchyme
Yifei Miao PhD1,2, Cheng Tan MD1,2, Nicole Pek BA1, Zhiyun Yu BA1, Kentaro Iwasawa MD1, Daniel Kechele PhD1, Nambirajan Sundaram PhD1, Victor Pastrana-Gomez BA1, Kyle McCracken MD/PhD1, Jason Tchieu PhD1, Jeffrey Whitsett MD1, Michael Helmrath MD1, Aaron Zorn PhD1, James Wells PhD1, Takanori Takebe MD/PhD1, Darrell Kotton MD3, Ya-Wen Chen PhD4, Minzhe Guo PhD1,2
1Cincinnati Children’s Hospital Medical Center, Cincinnati, OH, USA. 2University of Cincinnati College of Medicine, Cincinnati, OH, USA. 3Boston University and Boston Medical Center, Boston, MA, USA. 4Icahn School of Medicine at Mount Sinai, New York, NY, USA
• Gut epithelium, mesenchyme, and endothelium simultaneously co-develop from iPSC-derived organoids.
• Vascularized organoid provides a proper niche to pattern organotypic endothelium and mesenchyme.
• Organotypic endothelium and mesenchyme generated from organoids are similar to human fetal organs.

W-39
Multimolecular ultrasound imaging for vascular regeneration at centimeter depths
G. Edward W. Marti Ph.D., Cheng Liu Ph.D., Kaitlyn Liang M.S., James D. Brooks M.D., Jeremy J. Dahl Ph.D., Steven Chu Ph.D.
Stanford University, Palo Alto, CA, USA
• Molecular ultrasound imaging will enable deep tissue, long-term observations of vascular changes.
• This modality uses ligand- or antibody-labeled microbubbles to localize surface markers.
• We are developing a multi-molecular approach to track several molecular markers simultaneously.

W-40
Investigating the role of endothelial Rac1 towards regulation of vascular and organ homeostasis
Chen Yuan Kam PhD1, Ishani D Singh BS1, Karen K Hirschi PhD2, Valentina Greco PhD1
1Yale University, New Haven, CT, USA. 2University of Virginia, Charlottesville, VA, USA
• Neonatal endothelial Rac1 impairs angiogenesis but not vessel regression in maturing skin
• In adulthood, Rac1 mutant capillaries and venules are perturbed while arterioles are protected
• Endothelial Rac1 mutants display hair growth defects with delayed entry into hair cycle

W-41
Dissecting cutaneous wound healing in zebrafish
Leah Greenspan PhD, Keith Ameyaw BS, Daniel Castranova MS, Van Pham, Gennady Margolin, Caleb Mertus, Brant Weinstein PhD
NICHD, Bethesda, MD, USA
• Delays in vessel regrowth are a major contributor to defects in wound closure.
• Our new rotary tool injury system can be used to study cutaneous wound healing in zebrafish.
• Live imaging reveals insights into immune and endothelial cell behavior during wound healing.

W-43
Discovery of a new external immune surveillance organ in the zebrafish, an exciting model to study immune and endothelial cells
Daniel Castranova MS, Madeleine Kenton MS, Aurora Kraus PhD, Jong Park PhD, Marina Venero Galanternik PhD, Gilseung Park PhD, Daniel Lumbantobing PhD, Miranda Marvel PhD, James Iben PhD, Louis Dye PhD, Lucas Blevins, John K Frazer PhD, Brant Weinstein PhD
1NIH, Bethesda, MD, USA. 2University of Utah, Salt Lake City, UT, USA. 3Smithsonian Institution, Washington, DC, USA. 4University of Oklahoma Health Sciences Center, Oklahoma City, OK, USA
- Zebrafish have an external organ (lobe) behind their operculum that fully regenerates when removed.
- The lobe is teeming with immune cells and believed to be involved in immune surveillance.
- The lobe can be imaged in vivo and ex vivo, making it great to study immune and endothelial cells.

W-44
A zebrafish gill model of mammalian lung endothelium
Jong S Park Ph.D., Celia Martinez-Aceves, Daniel Castranova, Louis Dye, Madeline Kenton, Gennady Margolin, Van Pham, Brant Weinstein
NICHHD, Bethesda, MD, USA
- Many cell types found in lungs are also present in gills.
- scRNAseq revealed aerocyte-like cell population in the zebrafish gill endothelium.
- Gill aerocyte-like cells localize to the lamella of the gill where gas-exchange takes place.

Endothelium in Health and Disease II

W-45
Endothelial protease-activated receptors modulate hepatic vascular permeability during acetaminophen overdose
Rahul Rajala B.A., B.S., Audrey C.A. Cleuren Ph.D., Courtney T Griffin Ph.D.
1Oklahoma Medical Research Foundation, Oklahoma City, OK, USA. 2University of Oklahoma Health Sciences Center, Oklahoma City, OK, USA
- Endothelial PARs promote hepatic vascular permeability following APAP overdose.
- Endothelial PARs can reciprocally regulate their activating proteases.
- Translating Ribosome Affinity Purification can identify novel PAR targets following APAP Overdose.

W-47
Loss of GATA6 in human pulmonary artery endothelial cells shuts down interferon signaling
Vrinda Dambal Graduate Student, Maria Trojanowska Professor
Boston University, Boston, MA, USA
- Endothelium loses response to dsRNA and viral infection with knockdown of GATA6 in HPAECs
- Loss of GATA6 in HPAECs leads to inhibition of interferon signaling.
- Loss of interferon response in GATA6 knockdown HPAECs contributes to PAH development

W-48
Activation of endothelial GCN2 kinase contributes to pulmonary vascular remodeling and pulmonary arterial hypertension
Maggie M Zhu M.S., Mohammad Shafiq Ph.D., Jingbo Dai Ph.D., Yi Peng Ph.D., You Yang Zhao Ph.D., Stanley Manne Children’s Research Institute, Ann & Robert H. Lurie Children’s Hospital of Chicago, Chicago, Illinois, USA. 2Department of
Pediatrics, Division of Critical Care, Northwestern University Feinberg School of Medicine, Chicago, Illinois, USA.
3Departments of Pharmacology, and Medicine, Northwestern University Feinberg School of Medicine, Chicago, Illinois, USA. 4Feinberg Cardiovascular and Renal Research Institute, Northwestern University Feinberg School of Medicine, Chicago, Illinois, USA

- Loss of GCN2 kinase activity in mice inhibited hypoxia-induced PH.
- GCN2 was activated in response to hypoxia to mediate ET1 expression through GCN2/HIF2a/ET1.
- GCN2 activity was highly induced in pulmonary vascular lesions in IPAH patients.

W-49
Gadolinium blocks histone-induced endothelial calcium events and plasma membrane activation
Jade H Cleary, Sophia H Piffard BS, Carlos Lecuier-Garcia, Grant Hennig PhD, Kalev Freeman MD, PhD
University of Vermont, Burlington, VT, USA

- Extracellular histones trigger rapid cytosolic Ca2+ events in endothelial cells.
- Histones activate endothelial membranes causing blebbing and dye internalization within minutes.
- Ca2+ events and membrane effects of histones were completely abrogated by Gd3+.

W-50
The cerebrovasculature in a mouse model of Alzheimer's disease
Lauren R Miller PhD1, Sharon Negri PhD2,3, Debra Saunders1, Michael J Beckstead PhD1, Stefano Tarantini PhD2,3, Audrey C A Cleuren PhD1
1Oklahoma Medical Research Foundation, Oklahoma City, Oklahoma, USA. 2University of Oklahoma Health Sciences Center, Oklahoma City, Oklahoma, USA. 3Semmelweis University, Budapest, Hungary

- Endothelial-specific transcriptomics
- Vascular components of Alzheimer's disease
- Cerebrovascular function

W-51
Single nucleus sequencing reveals vascular microheterogeneity in acute rejection of human cardiac transplants
Christopher Pathoulas1, Amy Kimble1, Omar Omar1, Giulia Protto2, Gregory Fishbein M.D.3, Ryan Lau M.D.3, Matteo Pellegrini Ph.D.2, Patrick Murphy Ph.D.1, Nicole Valenzuela Ph.D.3
1University of Connecticut Health Center, Farmington, CT, USA. 2Department of Molecular, Cell and Developmental Biology, University of California, Los Angeles, CA, USA. 3Department of Pathology and Laboratory Medicine, David Geffen School of Medicine, University of California, Los Angeles, CA, USA

- There is a need for increased understanding of the processes that promote heart transplant rejection
- Endothelial cells become activated and could play an important role in graft rejection
- Longitudinal endothelial gene expression analysis of patient biopsies offers insight into rejection

W-52
Mechanobiology of vascular dysfunction in early age-related macular degeneration
Irene Santiago Tierno1,2, Sathishkumar Chandrakumar PhD1,2, Mahesh Agarwal PhD1,2, Kaustabh Ghosh1,2
1UCLA, Los Angeles, CA, USA. 2Doheny Eye Institute, Pasadena, CA, USA

- Endothelial cells from AMD eyes exhibit significantly impaired neovascularization.
- AMD ECs presented significant increase in FAK activity that was localized to disrupted EC junctions.
- Transcriptomics data revealed 16% of all DEG-enriched pathways were related to mechanotransduction.
W-53
Mechanical regulation of retinal vascular inflammation and degeneration in diabetes
Sathishkumar Chandrakumar PhD¹,², Irene Santiago Tierno MS¹,², Mahesh Agarwal PhD¹,², Timothy S Kern PhD³,⁴, Kaustab Ghosh PhD¹,²
¹UCLA, Los Angeles, CA, USA. ²Doheny Eye Institute, Pasadena, CA, USA. ³UC Irvine, Irvine, CA, USA. ⁴Gavin Herbert Eye Institute, Irvine, CA, USA
- Diabetes leads to lysyl oxidase (LOX)-mediated retinal capillary stiffening
- Stiffer retinal capillaries undergo inflammation-mediated degeneration in diabetes
- Diabetes-induced LOX upregulation is mediated by AGE/RAGE

Diseases (Vascular and Cardiovascular) II

W-54
Cardiovascular safety of nanoparticle vs conventional Doxorubicin
Lukas Brandt, Shelby N. Hader, Laura E. Norwood Toro PhD, Radha Vaddavalli PhD, Venkateswara Gogineni PhD, Sarah B. White MD, Andreas M. Beyer
Medical College of Wisconsin, Milwaukee, Wisconsin, USA
- Endothelial Dysfunction
- Cardio-Oncology
- Cardio-toxicity

W-55
The pH-sensing G protein-coupled receptor GPR68 promotes the contractile phenotype in vascular smooth muscle cells
Madison D Williams MS¹, Joshua S Morgan PhD¹, Michael T Bullock MS², Kristen Carraway¹, Tonya N Zeczycki PhD¹, Kyle D Mansfield PhD¹, David A Tulis PhD¹
¹East Carolina University, Greenville, NC, USA. ²Edward Via College of Osteopathic Medicine, Carolinas Campus, Spartanburg, SC, USA
- Decreased abundance of contractile and cytoskeletal proteins in GPR68 KO versus WT thoracic aortae
- Decreased expression of contractile and cytoskeletal proteins in GPR68 KO versus WT cells in hypoxia
- GPR68 may signal through the small GTPases Rap1A/1B and Rab5a to control VSMC phenotype

W-57
The liver as an inflammatory mediator of pulmonary arterial hypertension
Navneet Singh MD¹,², Amy Principotto², Corey E Ventetuolo MD, MS¹, Elizabeth O Harrington PhD²,¹
¹Brown University, Providence, RI, USA. ²Providence VA Medical Center/CPVB COBRE, Providence, RI, USA
- The liver may mediate inflammation that drives the pathobiology of pulmonary arterial hypertension
- Livers of Sugen-Hypoxia rats may demonstrate an inflammatory infiltrate compared to controls
- Increased GDF2 expression in SuHx livers may be a compensatory response to nonfunctional BMP9

W-58
MicroRNA loss drives tumor development in murine models of angiosarcoma
Annaleigh Benton, Ant Murphy, Jason A Hanna PhD
Purdue University, West Lafayette, IN, USA
- MicroRNA loss results in aggressive tumor development in mouse models of angiosarcoma.
- MiRNA-497 is a key tumor suppressing miRNA in angiosarcoma.
- We have identified several genes regulated by miR-497 that are upregulated in human angiosarcoma.
The slow progression of diabetic retinopathy is associated with transient protection of retinal vessels from death

Yanliang Li MD/PhD, Basma Baccouche PhD, Norma Del-Risco BS, Jason Park PhD, Amy Song BS, J. Jason McAnany PhD, Andrius Kazlauskas PhD
University of Illinois at Chicago, Chicago, IL, USA

- There is compelling evidence for protection from diabetic retinopathy in patients and animal models.
- Diabetes induced transient protection of the retinal vasculature in both T1D and T2D mice.
- As the duration of DM was prolonged, protection waned and vulnerability developed.

Cardiovascular Cell Biology II

STING inhibitors allow plasmid gene expression in primary vascular cells: an easy protocol for gene delivery in hard-to-transfect cells

Shuai Yuan, Adam Straub
University of Pittsburgh, Pittsburgh, PA, USA

- STING inhibitors enhance plasmid transfection in endothelial and smooth muscle cells.
- The inhibitor's effect is transient while protein expression sustains.
- Overexpressed protein retains normal functionality.

The role of COL8A1 in endothelial phenotypic modulation during inflammatory endothelial-to-mesenchymal transition

Qian Li, Sriharsha Talapeneni, Yonghong Meng, Linda L Demer, Yin Tintut, Jeffrey J Hsu
UCLA, Los Angeles, CA, USA

- COL8A1 mediates the endothelial cell transition during inflammatory EndMT.
- COL8A1 maintains normal endothelial function.
- TNF-a decreases COL8A1 expression to facilitate the loss of the endothelial phenotype.

Nuclear functions of smooth muscle α-actin

Callie Kwartler PhD1, Jose Emilianos Esparza Pinelo BS1, Anita Kau MD/PhD2, Dianna Milewicz MD/PhD1
1UTHouston McGovern Medical School, Houston, TX, USA. 2UTHouston McGovern Medical School, Houston, TX, USA

- Smooth muscle (SM) α-actin associates with the INO80 and BAF chromatin remodeling complexes
- SM α-actin p.Arg 179 variants alter histone modifications on critical SM-specific promoters
- Genetic evidence says altered chromatin remodeling may be a common pathway to moyamoya disease

Endothelial cell-targeted nanoparticle delivery of CRISPR/Cas9 plasmid DNA causes robust genome editing of endothelial HIF-2a in adult rats and inhibits experimental pulmonary arterial hypertension: Towards precision gene therapy of vascular diseases

Jingbo Dai PhD, Yi Peng PhD, Youyang Zhao PhD
Lurie Children's Hospital of Chicago and Northwestern University Feinberg School of Medicine, Chicago, IL, USA
• Endothelium-targeted nanoparticle delivery of the genome editing system in adult rats.
• Targeting vascular bed-specific ECs for modulation of gene expression.
• Selective disruption of Hif2a in lung EC inhibits pulmonary hypertension with minimal off-targets.

W-65
Variant-to-function of coronary artery disease: identifying novel regulation of the notch pathway in endothelial cells
Vivian S Lee-Kim Ph.D., Rajat M. Gupta M.D.
Brigham and Women's Hospital, Boston, MA, USA
• Human genetics to identify novel disease pathways
• Notch signaling pathway regulation
• Endothelial dysfunction in coronary artery disease

W-66
How does Dach1 promote vascularization?
Wen-Chuan Hsieh Ph.D., Kristy Red-Horse Ph.D.
Stanford, Stanford, CA, USA
• Assess FDA-approved drugs and small molecules that increase Dach1 level in artery endothelial cells.
• Investigate the molecular mechanism of Dach1 in artery maturation.
• Identify novel regulators and targets of Dach1.

W-67
A chemical biology approach for functional perturbation studies in the vasculature
Joseph Lim Bachelor of Science1,2, Jorge Andrade PhD1,2, Michael Potente MD1,2
1Angiogenesis and Metabolism Laboratory, Berlin Institute of Health at Charité – Universitätsmedizin Berlin, Berlin, Germany. 2Max Delbrück Center for Molecular Medicine in the Helmholtz Association, Berlin, Germany
• A highly efficient, specific, and reversible means to manipulate protein levels in blood vasculature
• Potentially overcoming inherent limitations of conventional genetic tools
• Gaining novel mechanistic insights by leveraging high(er)-temporal resolution of perturbation