Summaries of 2019 National Institutes of Health and other Federal Grants Awarded to HFHS

Part I – Department of Internal Medicine

- Allergy and Immunology
- Cardiology/Cardiovascular Research
- Endocrinology and Metabolism
- Gastroenterology
- Hematology/Oncology
- Hypertension and Vascular Research
- Infectious Disease
- Pulmonary
- Sleep Medicine
- General Internal Medicine

Allergy and Immunology

Principal Investigator: Edward Zoratti, M.D.
ICAC 3 - Inner City Asthma Consortium Infrastructure (UM1AI114271) Subcontract

The objectives of the Inner-City Asthma Consortium are to implement a long-range scientific plan to reduce asthma severity and prevent asthma among inner city children and to identify the mechanisms involved in the immunopathogenesis of asthma in these populations. The specific objectives are to: 1) conduct clinical trials to evaluate the safety and efficacy of promising immune based therapies in reducing asthma severity and preventing disease onset in minority children residing in inner cities in the United States; 2) conduct research to delineate the underlying mechanisms of such therapies as an integral part of the clinical trials undertaken by the Consortium; 3) conduct clinical studies on the immunopathogenesis of asthma onset, progression and severity; and 4) develop and validate surrogate/biomarkers to measure disease stage, progression and therapeutic effect.

Cardiology/Cardiovascular Research

Principal Investigator: Dennis Kerrigan, Ph.D.
The Effect of High Intensity Interval Training and Surgical Weight Loss on Distal Symmetric Polyneuropathy Outcomes (R01DK115687-02) Subcontract

Subrecipient will collaborate with the PTE to recruit and train subjects for a bariatric surgery high intensity interval training (HIIT) study. As the Sub recipient PI, Dr. Kerrigan has experience in HIIT studies for both patients with cardiovascular disease and cancer. Additionally, he oversees a weekly exercise class of 10-15 new patients prior to bariatric surgery who could be eligible for study. Dr. Kerrigan would oversee day-to-day operations of the project, be responsible for the protocol adherence, and supervise the research exercise physiologist. The research exercise physiologist would recruit potential subjects, communicate with the PTE project manager, and perform exercise training to those randomized into the HIIT group. Training will be conducted at one of Henry Ford Health System exercise facilities located in Detroit, West Bloomfield and Livonia.

Aims Include:

1. Identify and recruit subjects who will undergo bariatric surgery.
2. Identify and recruit subjects who are candidates for bariatric surgery but do not undergo surgery.
3. Perform the supervised HIIT training protocol.

**Principal Investigator: Steven Keteyian, Ph.D.**
**The Improving ATTENDance to Cardiac Rehabilitation (iATTEND) Trial (R61HL143099)**

The numbers of U.S. adults with either diabetes or pre-diabetes is staggering with nearly 40% of the population affected. In U.S. minority populations, such as African Americans, the diabetes rates are nearly twice as high as those of non-Hispanic white Americans. Skeletal muscle insulin resistance appears to be a nearly universal precursor to overt type 2 diabetes (T2D), and both insulin resistance and T2DM are often accompanied by mitochondrial dysfunction. With 16,569 base pairs and 13 protein-encoding genes, the mitochondrial genome is diminutive when compared with the ~6 billion base pairs in the diploid nuclear genome. To date, genomewide association studies of the nuclear variants have failed to explain a large proportion of the heritability of T2D, but rare mitochondrial mutations have been clearly implicated in T2D syndromes. Mitochondrial genetics has a number of complexities that haven’t been collectively considered in existing studies. First, cells possess hundreds to thousands of mitochondria, so the effect of a given variant may depend on mitochondrial number, as reflected in the DNA copy number. Second, there can be subpopulations of mitochondria within cells such that only portion of mitochondria carry a particular variant – a situation known as heteroplasmy. Moreover, copy number and heteroplasmy can differ between tissue types. Third, a number of genes encoding mitochondrial proteins are located in the nuclear genome. This implies that crosstalk between genomes is required to coordinate gene expression and the efficient production of essential mitochondrial complexes, such as those involved in electron transport. Therefore, epistatic interactions between genes on both genomes (i.e., mitonuclear interactions) may influence risk of T2D. We have two large study populations that will help us delve these complexities and their role in T2D – the Study of Asthma Phenotypes and Pharmacogenomic Interactions by Race-ethnicity (SAPPHIRE) and the Diabetes Multi-omic Investigation of Drug Response (DIAMOND). Through the TOPMed program, we will have whole genome sequence data on 3,596 African American adults in SAPPHIRE. African Americans are a particularly interesting population to study because limited numbers of mitochondrial haplogroups and large-sized chromosomal ancestral blocks minimize the numbers of comparisons needed to identify potentially important mitochondrial variants and mitonuclear interactions associated with T2D (Aim 1). The ongoing DIAMOND study will provide a population of T2D patients and controls in which to replicate the findings from Aim 1 using DNA isolated from skeletal muscle. Given the primacy of skeletal muscle in the pathogenesis of T2D, it is important to replicate mitochondrial findings from Aim 1 in this tissue (Aim 2a), as well as use it to identify unique variants associated with T2D and insulin resistance (Aim 2b). Lastly, given the described effect of exercise training on reversing skeletal muscle insulin resistance, we will investigate how exercise affects the heteroplasmy and copy number of variants identified in the earlier aims and how these changes relate to changes in insulin resistance (Aim 3).

**Principal Investigator: David Lanfear, M.D., and Hani Sabbah, Ph.D.**
**Plasma Metabolomics and Myocardial Energetics in Heart Failure (R01HL132154)**

Heart failure (HF) remains an enormous public health problem despite advances in treatment. Disease progression and response to therapy in HF varies widely between individuals, but breakthrough technologies such as genomics and metabolomics are helping to unravel the disease heterogeneity that confounds patient management. Perturbed energy metabolism may be a key contributor to cardiac dysfunction and the development of clinical HF. Evidence from a variety of sources indicates that impaired structure and function of the energetic apparatus in the myocardium contributes to disease severity, progression, and may influence response to treatment. However, in order to advance these observations toward meaningful interventions for HF patients, several key steps are still needed: 1) confirming the importance of metabolic variation in human HF, 2) developing noninvasive markers of myocardial energetic status, and 3) identifying promising targets for intervention. Our proposed project is a series of interwoven translational investigations in humans and dogs with HF to define the association of plasma metabolite levels with disease severity, myocardial energetics, and disease progression. This project leverages substantial infrastructure already in place, a large existing genetic cohort study, available plasma samples suitable for metabolomic profiling, comprehensive translational laboratory capabilities, and a cohesive multidisciplinary research group focused on HF. Together the planned studies will address the overarching hypothesis that the peripheral metabolomic signature can indicate disease progression/treatment responsiveness in HF patients and that this is driven by altered myocardial energy
metabolism. If true, then these data will help advance personalized therapy and identify novel targets for HF intervention, leading to improved outcomes for HF patients.

**Endocrinology and Metabolism**

**Principal Investigator, Arti Bhan, M.D.**

**Epidemiology of Diabetes Interventions and Complications (U01DK094157) Subcontract**

The Diabetes Control and Complications Trial (DCCT, 1983-1993) compared intensive therapy aimed at near normal glycemia versus conventional therapy with no specific glucose targets in 1441 subjects with type 1 diabetes (T1DM). In 1993, after a mean follow-up of 6.5 yrs., the study showed conclusively that intensive therapy reduced the risks of retinopathy, nephropathy, and neuropathy by 35-76%, and that hyperglycemia was a primary determinant of complications. We also described potential adverse effects of intensive therapy; assessed its effects on cardiovascular disease (CVD) risk factors, neurocognition and quality of life; and projected the lifetime health-economic impact. DCCT intensive therapy was then adopted world-wide as standard-of-care for T1DM. The Epidemiology of Diabetes Interventions and its Complications (EDIC, 1994-present) is the observational follow-up study of the DCCT cohort, with 95% of those surviving actively participating. Most outcomes are evaluated annually. CVD events and deaths are carefully documented and adjudicated. EDIC has notably discovered that the early beneficial effects of intensive treatment on complications have persisted for over 10 years despite the similar HbA1c levels during EDIC in the two groups, termed metabolic memory. Remarkably, former intensive therapy also greatly reduced the risk of CVD events. DCCT/EDIC collaborators have also conducted numerous ancillary studies, with separate funding, most recently including measurement of cardiac function on cardiac MRI and measurement of biomarkers of oxidative stress and inflammation as determinants of complications. The overarching goals for the next 5 years are to follow at least 90% of the surviving cohort; to describe accurately the study-long effects of glycemia (HbA1c) and other established and putative risk factors on diabetes complications and the metabolic memory effects of prior DCCT intensive therapy; and to expand knowledge regarding T1DM and its complications by supporting collaborations for new research funding applications to maximally utilize the cohort, phenotypic data set, and collected biologic and genetic samples. The specific scientific aims are to 1) evaluate effects of risk factors, biomarkers and glycemia on risk of clinical CVD; 2) assess the long-term changes in CVD risk factors; 3) describe effects of DCCT intensive versus conventional therapy on mortality; 4) evaluate risk factors for severe retinopathy/nephropathy; 5) assess effects of diurnal glycemic variation on complications; and 6) conduct eight new research projects involving new measurements and analyses.

**Principal Investigator: D. Sudhaker Rao, M.D.**

**Clinical Assessment of Vertebral Bone Quality Using Direct Biomechanical and Textural Analysis via Digital Tomosynthesis (W81XWH1910374)**

This project relates to the Topic Area “Musculoskeletal Disorders”, and specifically to the encouragement area of research on measures to improve diagnosis, prediction and optimization of health outcomes. This is because the proposed project ultimately aims to improve the accuracy of assessment for spinal bone fragility and fracture risk. The bones of the spine (vertebrae) are the most frequently fractured ones due to osteoporosis. These fractures are economically costly and burden the patients with many downstream problems including back pain. Military personnel are known to be at greater risk for these fractures and complications. An accurate assessment of vertebral fracture risk is essential for appropriate and timely intervention for the prevention of fracture. This research also relates to the Topic Area “Diabetes”, because a diabetic cohort will be included in the study. Current standard techniques for fracture risk assessment rely on radiographic bone density scans. Additional information regarding the patient’s demographic status and medical history is also incorporated in tools predicting fracture risk. However, these techniques are not very sensitive in identifying who will have a fracture and who will not. This is not too surprising, considering the fact that the information used in the assessment is a crude indirect measure of bone strength not based on biomechanics. To address this concern, we developed a new method, in which two images of a patient’s vertebra are taken in the presence and absence of the patient’s body weight by having them stand and lay down for both images respectively. The images are obtained using digital tomosynthesis (DTS), a system that is similar to computed tomography (CT). The advantages of DTS over CT are that DTS allows for standing and lying images to be captured, offers high resolution and exposes patients to less radiation than CT. The
two sets of images are compared using an advanced computational method and deformations in the vertebra caused by standing are measured. From the displacement measurements, vertebral stiffness and overall displacement are calculated as metrics of strength and factor of safety (factor safety is a measure of how strong the bone is relative to the loads it normally experiences). Information on bone microstructure, additional to bone density, is known to increase accuracy in predicting fractures. We can also derive properties related to bone microstructure from DTS images without the biomechanical test. These properties are determined by quantifying the texture in the bone image and called textural properties. We developed these methods in the laboratory in detail using cadaveric vertebrae and laboratory-standard imaging and strength testing. We also performed pilot human studies to establish feasibility of the methods in the clinic. What remains to be determined is how successful the methods will be in the clinical environment for identifying individuals who are at risk. Therefore, this study will be a clinical validation of the new biomechanical and textural DTS methods. In order to determine the ability of DTS methods to correctly identify at-risk patients, the approach will be to compare patients who have conditions or diseases that are known to increase their risk of fracture to normal patients. Therefore, a group of osteoporotic patients with an existing vertebral deformity, a group with primary hyperparathyroidism (pHPT) and a third group with diabetes will be compared to normal patients. These diseases are considered service-related and thus represent a greater risk for military families. Importantly, each of these diseases increase the risk of fracture but alter bone in different ways that are not always detectable by bone density scans. For example, osteoporosis primarily results in loss of bone mass, pHPT alters the organization of bone structure and affects the cortical bone (in the case of a vertebra, the dense bony shell surrounding the vertebra), and diabetes alters the quality of the bone material without reducing bone mass. By studying these groups using DTS and comparing the assessment of bone strength to standard bone density results, we will; i) establish for which types of patients and to what extent the DTS methods might be useful, ii) identify in which way the method must be performed for best results, iii) better understand how differences in bone quality specific to each disease affect the biomechanical outcome, and iv) establish a group of patients that we can follow up for longer term results. In the long term, the method will be useful in other clinically significant issues such as low back pain associated with vertebral fractures, implant stability, degenerative and congenital diseases of the skeletal system resulting in deformities, skeletal response to drug, exercise and disuse.

Gastroenterology

Principal Investigator: Stuart Gordon, M.D.
Chronic Hepatitis Cohort Study II (CHeCS-II) (U18PS005154)

Hepatitis B (HBV) affects over 1.25 million Americans, and hepatitis C (HCV) over 3.2 million Americans. In the decades to come, more than 150,000 Americans are expected to die from these conditions unless steps are taken to increase awareness, diagnosis, and access to necessary care and treatment. Emerging interferon-free, direct-acting all-oral antiviral (DAA) treatments have changed the landscape of HCV treatment and care. These treatments appear to be safer than interferon-based treatments and provide exceptionally high rates of sustained virological response (SVR). Both HBV and HCV treatment guidelines have been updated to reflect evidence regarding initiation of new therapies; however, the evidence for those recommendations is largely based on clinical trials conducted under highly controlled conditions in restricted patient populations with limited data collection. Significant health disparities—across race, sex, age, and co-infection (with HIV or dual hepatitis)—may limit the generalizability of these populations. Data from longitudinal cohorts of “real world” hepatitis patients are needed to assess the population impact of rapidly evolving antiviral therapies, to understand the spectrum of disease and its natural history, and to evaluate the public health impact of chronic viral hepatitis. The Chronic Hepatitis Cohort Study (CHeCS) is the first comprehensive longitudinal cohort study of chronic viral hepatitis in the USC and has served as a model platform for observational data collection in this population. Since 2010, CHeCS has reported valuable information and expanded knowledge on many facets of hepatitis disease and policy. We propose to build upon CHeCS to develop “CHeCS-II,” in order to achieve the long-term goal of applying this rich data and infrastructure resource to inform public health planning, policy decisions, and clinical management of HBV and HCV. To achieve this, we will leverage the established CHeCS infrastructure, which has: (1) a diverse, real-world, non-veteran-based US cohort of >3,000 HBV, >11000 HCV, and >500 HIV co-infected patients receiving care through four U.S. health systems; (2) an experienced multidisciplinary team; (3) an efficient system for patient identification and data collection. We will provide scientific leadership to identify research findings and priorities by: (1) Offering seamless collaboration across study sites and with the Centers for
Disease Control (Aim 1); (2) Expanding our HCV cohort to over 14,000 patients with >2 years’ follow-up; (3) Increasing follow-up of HBV patients to >5 years; (4) Collecting additional data regarding social determinants of health, including access to and uptake of care (Aim 2); (5) Applying rigorous analytical approaches to develop an in-depth understanding of health disparities and comorbidities, as well as investigating how these differences impact access to and uptake of antiviral therapy; (6) Advancing translation of this research to inform hepatitis-related policy and practice (Aim 3).

**Hematology/Oncology**

**Principal Investigator:** Ding Wang, M.D., Ph.D.
**SWOG Network Group Operations Center of the NCTN (U10CA180888) Subcontract**

The impact statement in SWOG's network operations grant application succinctly summarizes our work and our goals: By continuously improving inclusion, engagement, and scientific innovation, SWOG will enhance cancer clinical trial development and conduct, reducing the burden of human neoplasm. The SWOG National Clinical Trials Network Group has established itself as an innovative, collaborative, and cost-effective NCTN constituent. SWOG has 60 years of trial experience, and its work has led to the Food and Drug Administration approval of 14 regimens, changing and informing oncologic practice hundreds of time more. In our 2013 grant application, we promised to make unique contributions to the new NCTN enterprise, and we successfully did so over the last five years. We are strongly committed to furthering our efforts over the next six. SWOG designs and directs high-value, pathway- and immune- driven oncology research, with the goal of achieving practice-changing results that are meaningful to both persons affected by cancer and investigators. The group's current network includes more than 1,000 member sites, with 5,000 physicians who practice across the United States, Canada, South Korea, Mexico, Saudi Arabia, and South America. Twenty-three NCI-designated cancer centers number among our members, as do 22 Specialized Programs in Oncology Research Excellence. From early 2014 through mid-2017, SWOG investigators published more than 188 cancer treatment articles and abstracts and enrolled 12,819 patients into NCTN therapeutic trials. SWOG actively collaborates in NCTN direct research and administrative functions and has developed training and education tools used throughout the network. SWOG's mission is to significantly improve lives through cancer clinical trials and translational research. The following guiding principles, ratified in 9/2017, are the foundation upon which we build to achieve that end: — We make patients our absolute highest priority — We ensure that the best science drives our research — We embrace and encourage diversity in leadership and membership, to effectively solve problems in cancer — We demand integrity, accountability, and ethical behavior in SWOG — We foster and mentor young investigators, to ensure excellent clinical research for future generations. Over the next grant cycle, we will provide an efficient, innovative, and nimble network capable of developing and conducting abroad framework of clinical and translational trials; we will meaningfully contribute to the NCTN; and we will help patients lead longer and meaningful lives. SWOG will remain an innovative force in the design of the next generation of oncologic therapies.

**Hypertension and Vascular Research**

**Principal Investigator:** Emilio Mottillo, Ph.D.
**Direct Analysis of Lipolysis-mediated Signaling Events (5R00DK114471)**

Project Summary/Abstract Obesity has reached epidemic proportions and is tied to the greater prevalence of metabolic disorders such as diabetes and cardiovascular disease. While the precise mechanisms by which obesity causes diabetes are not entirely clear, mounting evidence suggests that the body's normal process of sensing lipids is disrupted. This inability to properly detect lipids can lead to lipotoxicity and cause detrimental effects in key insulin sensitive tissues. Thus, an important scientific goal, and that of this NIH Pathway to Independence Award, is to understand the mechanisms by which cells sense lipids and thereby maintain lipid homeostasis. The training component of this application builds upon the candidate's interest and background in imaging metabolism and metabolic signaling/energy sensing, while providing a unique environment to train in career development activities related to a team science approach of doing research. The research training component will utilize a unique set of tools that will allow the candidate to probe the direct effects of lipolysis independent of transmembrane-protein kinase A (PKA) signaling and image fatty acid metabolism. The candidate will gain experience in global analysis techniques of.
phosphoproteomics and lipidomics, and super-resolution imaging. Utilizing these tools and training, the candidate will determine 1) the signals directly generated by lipolysis, and 2) the dynamics of lipid trafficking and lipolysis-derived signals within a cell. Central to this aim is the hypothesis that signals directly produced by lipolysis function to maintain lipid homeostasis and are highly dynamic. The research component of the award will be accomplished by the following specific aims: Aim 1: Identification of signals that are generated directly by lipolysis. Lipolysis is known to produce signals, but up to this point the direct effects of lipolysis were not distinguishable from transmembrane-PKA signals. Utilizing novel synthetic ligands that activate ABHD5, a lipase co-activator protein, Aim 1 will be accomplished by the following sub-aims: 1a: To identify ABHD5-dependent lipid mediators. Utilizing a lipidomic approach, the candidate will identify the bioactive lipids produced by ABHD5 that regulate downstream metabolism. 1b: To identify ABHD5-dependent kinase activation pathways. Using a phosphoproteomic approach, the candidate will determine the phosphorylation events, kinases and pathways that are a direct consequence of ABHD5 activation. Aim 2: To determine the trafficking dynamics of fatty acids and their metabolites and lipid mediators. This will be accomplished by the use of newly developed genetically encoded fluorescent sensors that allow the monitoring of the temporal and spatial dynamics of fatty acids and fatty acyl-CoAs. The proposed K00/K99 is well aligned with the mission of the NIH and the NIDDK and will train a promising scientist to understand the mechanisms that regulate lipid homeostasis and potentially the pathways that are disrupted during obesity, a significant public health priority.

Principal Investigator: Pablo Ortiz, Ph.D.

Fructose Induced Salt-Sensitive Hypertension: Role of Thick Ascending Limb Transport (R01DK107263)

A high-fructose diet is linked to the epidemic of hypertension, diabetes, and obesity. Up to 25 million Americans consume up to 20% of their calories from added fructose1,2. We found that feeding rats a fructose-enriched diet (20%) for 4 weeks did not increase blood pressure. However, a fructose-enriched diet combined with high salt (4% Na) caused salt-sensitive hypertension within 1 week (Figures 1,11); prior to the development of metabolic abnormalities. The initial phase of salt-sensitive hypertension is in part mediated by a renal defect that prevents NaCl excretion during high salt intake. The thick ascending limb (TAL) reabsorbs 25% of filtered NaCl. Enhanced TAL NaCl absorption is related to salt-sensitive hypertension in humans and rodents3-5. However, the mechanism by which a fructose-enriched diet rapidly (1 week) causes salt-sensitive hypertension is not clear and the role of TAL NaCl absorption in this process is completely unknown. NaCl reabsorption by the TAL depends on the apical Na/K/2Cl cotransporter NKCC2, the target of loop diuretics. Our preliminary data show that a fructose-enriched diet enhanced NKCC2 phosphorylation in TALs. However, the effects of fructose and the signaling induced in the TAL and the distal nephron have not been studied. Our data show that plasma and urine fructose increase rapidly after fructose intake. Thus, fructose reaching the nephron may be transported in by a fructose channel, activating protein kinase signaling. The only kinases known to phosphorylate Thr96,101 of NKCC2 are SPAK (STE20/SPS1-related proline-alanine-rich kinase) and OSR1 (Oxidative Stress Responsive 1) kinases. In the TAL, these kinases specifically phosphorylate NKCC2. In the distal convoluted tubule (DCT), these kinases specifically phosphorylate the thiazide sensitive NaCl transporter NCC. We found that a 20% fructose diet increases SPAK/OSR1 phosphorylation in TALs. In addition, stimulation of β-adrenergic receptors (β-AR) in the TAL activates NKCC213. A fructose-enriched diet may increase sympathetic activity by 2 weeks12, or enhance the sensitivity or signaling of β-AR. Our preliminary data show that β-AR stimulation increases SPAK/OSR1 phosphorylation in TALs. In the Dahl salt sensitive (SS) rat, NKCC2 and SPAK/OSR1 phosphorylation are abnormally enhanced in a normal salt diet. It is not known whether this increases the effect of fructose on blood pressure and NaCl absorption. We hypothesize that a fructose-enriched diet enhances thick ascending limb (TAL) and distal tubule (DCT) NaCl absorption by inducing NKCC2 and NCC phosphorylation via SPAK/OSR1 kinases and enhanced β-AR signaling. These effects occur within 1 week, prior to metabolic alterations, and are maintained chronically (16 weeks), promoting salt-sensitive hypertension in normal rats. In Dahl SS rats, abnormally elevated SPAK/OSR1 in the TAL, enhances the effect of fructose on blood pressure in normal- or high-salt diets.
Diabetes mellitus (DM) afflicts 26 million people in the US. 40-70% of these diabetics die of cardiovascular complications. We and others found that DM increases reactive oxygen species (ROS)-mediated aldehydes like 4-hydroxy-2-nonenal (4HNE) generation. 4HNE forms covalent bonds with macromolecules known as adducts, which lead to cellular damage and decreased cardiac function. Aldehyde dehydrogenase (ALDH2) is a mitochondrial enzyme that detoxifies 4HNE in the heart. We and others have reported that in streptozotocin-induced hyperglycemic models increase in 4HNE protein adducts and decrease in myocardial ALDH2 activity correlate with cardiomyopathy. Although we think this causes cardiac dysfunction, the exact mechanism is unclear. However, most diabetic patients have type-2 DM. Thus, it is imperative to investigate whether hyperglycemia-induced 4HNE and lower ALDH2 activity contribute to cardiac dysfunction in type-2 DM models. We recently demonstrated that high glucose stress or 4HNE administration decreased mitochondrial respiration with increased mitochondrial DNA (mtDNA) damage in cultured cardiomyocytes. In our preliminary study using type-2 diabetic mouse heart, we found an increase in mitochondrial levels of 8-hydroxyguanine (8OHG), an oxidized mtDNA product, which is primarily repaired by 8-oxoguanine glycosylase (OGG)-1. Next, we found increased 4HNE adduct formation on OGG-1 and reduced cardiac OGG-1 levels. These data suggest that 4HNE adduction on OGG-1 reduces its level and activity thereby raising the unmetabolized 8OHG level. Thus, we postulate that 4HNE-mediated mtDNA damage is part of the mechanism by which lower ALDH2 causes mitochondrial respiratory dysfunction and thus cardiac contractile dysfunction. To test our idea, we will use a high-fat diet induced type-2 DM model in wild type (WT) C57BL/6 and ALDH2*2 mutant mice. This mutation mimics East Asians with the E48K variant (ALDH2*2), which exhibits lower ALDH2 activity. We will overexpress ALDH2 gene in the myocardium in situ or treat our diabetic mice with Alda-1, the only specific drug available to improve the catalytic activity of both WT and mutant ALDH2. We propose following three specific aims: **Aim 1-Hyperglycemia in models of type-2 diabetes reduces ALDH2 activity in cardiac myocytes by increasing 4HNE adduction with ALDH2:** Aim 2- Increased 4HNE adduct formation on mtDNA and OGG-1 causes mtDNA damage and poor mitochondrial respiration in type-2 DM: **Aim 3- Augmenting ALDH2 activity reduces 4HNE-mediated mtDNA damage and thereby cardiomyopathy progression in type2-DM.** This study will identify a novel role of ALDH2 in type-2 DM mediated cardiac dysfunction and establish that ALDH2 could be a therapeutic target for restoring cardiac function in type-2 diabetic patients.

**Principal Investigator: Suresh Palaniyandi, Ph.D.**

**4-hydroxy-2-nonenal in Mitochondrial DNA Damage and Contractile Dysfunction in Diabetic Heart: A Role for Aldehyde Dehydrogenase 2 (R01HL139877)**

Hypertension is a major health care burden in the United States, affecting 1 in 3 adults. Hypertension is associated with concomitant coronary artery disease with myocardial infarction (MI) and heart failure (HF). In this study, we will define how N-acetyl-seraryl-aspartyl-lysyl-proline (Ac-SDKP) protects cardiac structure and function in a mouse model of HF that will be induced in two models [angiotensin II (Ang II) hypertension- or permanent left anterior descending coronary ligation (LAD)]. We and others reported that Ang II-induced hypertension or LAD resulted in HF associated with cardiac structural remodeling and impaired function. Ac-SDKP is successively produced from thymosin \(\beta\)-4 (T\(\beta\)4) by two enzymes, meprin \(\beta\) and prolyl oligopeptidase (POP). Circulating and tissue Ac-SDKP depends on the angiotensin converting enzyme (ACE) activity, since Ac-SDKP is mainly degraded by the N-terminal active site of ACE (ACE-N). ACEI are first-line drugs to treat HF. ACEI have strong side effects such as hypotension, cough, rash, angioedema, hyperkalemia, and dysgeusia, whereas Ac-SDKP has none, even at high dosages (up to 48 mg/kg/d). Also, Ac-SDKP is down-regulated in dogs and patients with chronic HF. Whether and how Ac-SDKP therapy could rescue hypertension- or LAD-induced cardiac complications remain to be elucidated. Increasing circulating Ac-SDKP not only inhibited fibrosis and mediators of inflammatory cell infiltration into the injured myocardium, but it also improved cardiac function in mice with LAD or hypertension (preliminary data). We have found that Ac-SDKP inhibits endoplasmic reticulum (ER) stress in cardiac fibroblasts in vitro and in mice with MI and restores phosphor-AKT in hypertensive hearts. Activation of ER stress is detrimental to the endothelium, cardiac fibroblasts, and cardiomyocytes. These findings set the scientific premise of this work, providing foundational work that Ac-SDKP represents a beneficial supplement to the existing cardiac pharmacotherapy. Our central hypothesis is that Ac-SDKP protects and potentiates cardiac protection.
against heart failure via the inhibition of ER stress. We propose to use the mouse model of heart failure induced by hypertension or LAD to address the following 2 aims: (1) we will determine whether Ac-SDKP protects the heart and provides additional cardiac protective effects to ARBs, ACEi, or eplerenone in mice with MI or hypertension, (2) and we will demonstrate that Ac-SDKP improves cardiac function in mice with hypertension or LAD by inhibiting the detrimental ER stress via the PI3K/AKT pathway. A number of conditional and tissue-specific knockout female and male mice will be employed. A team with significant expertise is recruited for this project, which will apply a combination of state-of-the-art in vivo, cell and molecular techniques including measurements of cardiac remodeling and function by echocardiography in non-anesthetized mice and radiotelemetry, which can detect the blood pressure, the electrocardiogram, and the heart rate of conscious mice. These studies will help define the cause-effect relationship between Ac-SDKP and HF and its mechanism towards the protection from HF.

**Infectious Disease**

**Principal Investigator:** Marcus Zervos, M.D.

**Improving antimicrobial use at hospital discharge through a collaborative pharmacist-led transition-of-care intervention (75D30118C02928)** Other Federal Contract

Traditional antimicrobial stewardship interventions in the hospital utilize core strategies such as formulary preauthorization and audit and feedback, while decisions about the choice and duration of discharge antibiotics may be limited to more passive interventions of guidelines and education. Pharmacists practicing in TOC focus specifically on optimizing medication management and patient education at discharge to home, long-term care facilities, or other settings. Using the existing infrastructure of the TOC model to extend the scope of ASP interventions, we can address this gap. The goal of this project will be to implement and evaluate a multi-disciplinary ASP bundle for patients discharged from general medical and/or surgical units on antibiotics. At the time of collaborative discharge rounding, the TOC team will identify patients on, or transitioning to, oral antimicrobials to prepare for hospital discharge and applicable follow-up. Optimal antimicrobial selection, dose, duration, and dispensing location will be determined, and the pharmacist will enter the prescription in the discharge queue, which the provider will send for the patient at the time of discharge. This early entry of discharge orders into the queue also provides opportunity to facilitate medication access for those with financial barriers or insurance plans that do not align with medications. The primary outcome will be appropriate duration of therapy prescribed at hospital discharge.

**Aim 1:** Implement a multi-disciplinary antibiotic discharge assessment for adults with respiratory, urinary, intra-abdominal, and skin infections

**Aim 2:** Evaluate the impact on appropriate durations of therapy, infection-related readmissions, and subsequent antibiotic-associated harms

**Aim 3:** Describe burden of illness and opportunities for improvement with discharge antibiotic prescribing patient access, resistance, and antibiotic-related adverse effects

The long-term objective of this work is to improve patient outcomes such as clinical resolution and reduced development of resistant pathogens adverse drug events (ADE). Inappropriate antibiotic prescribing contributes [.0 medica jon-related adverse events, hospital readmissions, and the development of drug-resistant pathogens. Novel strategies involving all healthcare providers are sorely needed to address these concerns.

**Pulmonary**

**Principal Investigator:** Jayna Gardner-Gray, M.D.

**Clinical Centers (CC) for the NHLBI Prevention and Early Treatment of Acute Lung: Reevaluation of Systemic Early Neuromuscular Blockade (U01HL123031) Subcontract**

**Purpose:** This study is evaluating whether giving a neuromuscular blocker (skeletal muscle relaxant) to a patient with acute respiratory distress syndrome will improve survival. Half of the patients will receive a neuromuscular blocker for two days and in the other half the use of neuromuscular blockers will be discouraged.

**Trial Summary:** Study Design: This is a multi-center, prospective, 2-arm, unblinded, randomized clinical trial of two management strategies of neuromuscular blockade (also called skeletal muscle relaxant and
muscle relaxant). Purpose: To assess the efficacy and safety of early neuromuscular blockade in reducing mortality and morbidity in patients with moderate-severe ARDS in comparison to a control group with no routine early neuromuscular blockade. Sample Size: This trial will enroll approximately 1400 subjects from PETAL network hospital ICUs.

Sleep Medicine

Principal Investigator: Philip Cheng, Ph.D.
Clinical Translation of Phenotypes of Shift Work Disorder (1K23HL138166)

Shift work disorder (SWD) is a significant threat to public health and safety; over 6 million shift workers in the United States experience the debilitating symptoms of excessive sleepiness and insomnia and suffer functional impairments that increases the risk of catastrophic industrial accidents. However, patients with SWD are often inadequately treated because the pathophysiology is not well-characterized, and current diagnostic assessments do not identify specific treatment targets. Consequently, clinicians are unable to deliver precise interventions that target the underlying causes of SWD. The proposed project in this career development award will address these gaps by taking the initial steps of translating two phenotypes of SWD for clinical use. Previous research has indicated that SWD can arise from two independent pathways that can be categorized as pathophysiological phenotypes. The first is the circadian misalignment phenotype, characterized by poor adjustment of the biological clock to the nocturnal work schedule. The second is the sleep reactivity phenotype, characterized by a trait vulnerability to sleep disturbance triggered by environmental stressors. Both phenotypes lead to symptoms of sleepiness and insomnia in SWD and is not currently distinguished in the clinic; however, the requisite treatments for each pathophysiological phenotype are entirely different. As such, the appropriate intervention of SWD requires that these phenotypes be adequately characterized and identified in the clinic. The proposed aims will complete the requisite foundational research to launch the translation of these phenotypes of SWD for clinical use. The first research aim will examine the stability of each phenotype in shift workers to characterize them as either state or trait phenotypes, which will impact both assessments of interventions. The second research aim will identify the specific clinical attributes that can be used to index the phenotypes in a brief, accurate, and cost-effective assessment tool. Finally, the third research aim will identify differences in cognitive and performance deficits between the two phenotypes so that accidents and injuries can be preempted with targeted interventions. To successfully complete the research aims, and to support my long term goal of conducting translational research to improve the health and productivity of shift workers, this career development award will provide further training in the following areas: (1) development of clinical screening tools, (2) advanced methodologies in clinical and translation research, (3) feasibility of real-world behavioral interventions for shift work disorder, and (4) advanced field measurement of circadian phase. In combination, the training activities outlined in this career development award will provide the necessary expertise for a sustained career in translational research and circadian medicine.

Principal Investigator: Christopher Drake, Ph.D.
Sleep to Reduce Incident Depression Effectively (STRIDE) (1R56MH115150)

Abstract Prevention of major depressive disorder (MDD) is a public health priority and is in critical need of innovative strategies that preemptively identify those at-risk in order to enable early intervention. A recent meta-analysis of over 20 longitudinal studies found the risk for incident depression among individuals with insomnia disorder is nearly three times that for normal sleepers, thus making insomnia a potential point of entry for depression prevention. Identification and treatment of insomnia typically occurs in primary care, and is commonly treated with hypnotic medications; however, hypnotics have significant limitations, including increased risk for residual impairment, falls in the elderly, and abuse. Cognitive behavioral treatment of insomnia (CBT-I) has been recommended as a first line approach with demonstrated efficacy that is sustained beyond initial therapeutic intervention. However, effective and widespread implementation of CBT-I is severely limited by the national shortage of trained practitioners in clinical practice. A stepped care approach rooted in primary care holds potential for innovative accessibility and delivery of CBT-I, improving insomnia therapeutics, and reducing rates of MDD by targeting a robust yet modifiable risk factor in insomnia. Our proposed stepped care model uses digital cognitive behavioral therapy (dCBT-I) as an accessible, least-restrictive, first line intervention that reduces specialist time and resources, and adds clinician based face-to-face CBT-I only for refractory patients who need a more personalized, flexible, and durable therapist driven
approach. We propose a large-scale clinical trial in the primary care setting that utilizes a stepped care model (SMART design) to determine the effectiveness of dCBT-I alone and in combination with face-to-face CBT-I for insomnia, and the effects of these sleep interventions on the prevention of MDD. An important innovative component of the trial is the 1 and 2-year follow-up assessments to determine the durability of effectiveness over time and assess the impact on depression incidence and relapse. Early risk-detection and prevention is especially critical in those at elevated risk for depression to reduce health disparities. Thus, individuals with significant vulnerability to MDD, such as high sleep-reactivity, low socioeconomic status, and racial minorities will be included in significant numbers to test for potential moderation of treatment effects stratified by risk. Finally, improving sleep through insomnia treatment may reduce nocturnal rumination, which may mitigate progression toward MDD. As such, we will determine whether changes in nocturnal rumination (i.e., target), a modifiable risk-factor, mediates the effects of CBT-I and dCBT-I on MDD incidence and relapse. This project will test a highly scalable model of sleep care in a large primary care system to determine the potential for wide dissemination to address the high volume of population need for safe and effective insomnia treatment and associated prevention of depression.

Principal Investigator: Timothy Roehrs, Ph.D.

Risks for Transition from Therapeutic Hypnotic Use to Abuse (R01DA038177)

The acknowledged drugs of choice for the pharmacological treatment of insomnia are the benzodiazepine receptor ligand hypnotics (BzRL). Our nighttime studies show that with therapeutic doses used either short-term or chronically, the abuse liability of BzRLs in insomnia is not seen universally and is relatively low. The data from our last grant, a first-ever study, showed the abuse liability of chronic zolpidem use in insomniacs was low. Yet case reports and retrospective studies continue to report BzRL dependence and for the majority of these cases the abuse developed through initial therapeutic use. In our study some subjects showed an increase in dose across time. Understanding the transition from therapeutic use to abuse and identifying risk factors, such as specific patient and drug characteristics, is both mechanistically and clinically important. Our preliminary data have shown that a subset of insomniacs, those insomniacs that have signs of hyperarousal as reflected by elevated Multiple Sleep Latency Test (MSLT) scores, increased their nightly zolpidem dose across time. BzRLs have differential receptor binding affinities and associated anxiolytic or antidepressant properties. Zolpidem has selective alpha 1 BzRL affinity and little mood activity and thus may show less risk for transition from therapeutic use to abuse than another currently frequently prescribed BzRL with less alpha subtype selectivity such as eszopiclone. We propose to study the abuse liability of a selective (zolpidem) vs nonselective (eszopiclone) hypnotic during chronic use (six months) in an at-risk subpopulation (insomniacs with hyperarousal shown by elevated MSLTs). The proposal is highly innovative as it reflects a paradigm shift in understanding the abuse liability of hypnotics. In the end, this proposal will generate a unique set of data addressing a number of previously clinically important unanswered questions regarding hypnotic abuse by insomniacs (i.e., its likelihood as a function of arousal state and specific hypnotic pharmacology, of dose escalation over time and change in mood/drug effect ratings over time). It will provide clinicians with behavioral indicators of abuse risk.

General Internal Medicine

Principal Investigator: David Willens, M.D., MPH, FACP

eAssist A Post-Visit Patient Portal Tool to Promote Colorectal Cancer Screening (R01CA197205-05) Subcontract

The goal of this study is to develop and test a patient portal tool (e-Assist) for engaging and supporting primary care patients to make decisions about and to obtain colorectal cancer screening. Henry Ford Health System will serve as the performance site for this study. As such, the e-Assist tool will be programmed by HFHS programmers and all day-to-day aspects of the study will be coordinated by staff at Henry Ford’s Center for Health Policy and Health Services Research with oversight from the study PI and team. These tasks include identifying eligible participants and coordinating all elements of communication with patients (letter, phone interviews, etc.) In addition, HFHS staff will conduct focus groups and cognitive interviews with patients for preliminary testing of the e-Assist tool.
Part II – All Other Clinical Departments

- Dermatology
- Neurology
- Neurosurgery
- Orthopaedics/Bone & Joint
- Otolaryngology
- Pediatrics
- Radiation Oncology
- Urology
- Women’s Health

Dermatology

Principal Investigator: Qing-Sheng Mi, M.D., Ph.D.

Genetic and Genomic Dissection of Psoriatic Arthritis (R01AR063611)

HFHS will participate in this project by continuing its longitudinal assessment of patients with cutaneous psoriasis (PsC) and psoriatic arthritis (PsA), through our existing collaboration in IPART (the International Psoriatic Arthritis Research Team). It will also continue to collaborate with PI James Elder and Co-PI Dafna Gladman (University Health Network, Toronto, CA) to provide serum samples to identify serum micro-RNA (miRNA) biomarkers of conversion of PsC to PsA. Under the direction of Co-PI Qing-Sheng Mi, HFH will assay serum miRNAs for the purpose of biomarker identification and provide guidance for the assessment of biological functions of identified serum miRNA biomarkers. HFH (through IPART) will contribute additional DNA samples from patients with PsC and PsA as well as normal controls for expanded GWAS genotyping of psoriasis, PsC, and PsA. Finally, HFH (through IPART) will contribute additional blood samples for assessment of blood mRNAs and miRNAs as biomarkers for progression of PsC to PsA.

Applicant Identifier 19-PAF02365 submitted in response to Opportunity ID PA-18-484, is incorporated herein by reference as applicable.

Principal Investigator: Qing-Sheng Mi, M.D., Ph.D.

Uncover the New Subsets of Epidermal Langerhans Cells (1R61AR076803)

Langerhans cells (LCs) are skin-resident dendritic cells (DCs) expressing the C-type lectin Langerin (CD207) that mediate both adaptive immunity and immune tolerance in skin and are involved in various types of skin diseases. Adult LCs are originated from embryonic yolk-sac-derived macrophages and fetal liver monocytes in the steady state. Interestingly, LCs could also be derived from the bone marrow or peripheral monocytes and repopulate the skin under inflammatory conditions. However, due to the lack of molecular profiles at individual LC level, a significant gap remains in our understanding on how a single CD207+ epidermal LC population can induce both immunity and tolerance. Fortunately, new technologies such as the single-cell RNA-sequencing (scRNA-seq) can evaluate cell-to-cell transcriptomic variation, while the single-cell assay for transposase-accessible chromatin using sequencing (scATAC-seq) can assess the epigenomic heterogeneity at single-cell resolution in an unbiased manner. Recently, we identified two major LC subsets in mice, ATF3+Bal2a1b- (mLC1) and ATF3-Bal2a1b+(mLC2) subsets, and three major LC subsets in human including ATF3+ (hLC1) subset using scRNA-seq. We also found in ATF3 knockout mice that lack of ATF3 enhances LC maturation and promotes LCs-induced Th1 and Th17 cell differentiation suggesting immune suppressive function induced by ATF3+LC1. Hence, these preliminary data support our hypothesis that LCs are heterogeneous consisting of distinct subset with different immune functions. Our objective is to use single-cell analysis platforms plus the LC fate-mapping and mutation mouse models to further validate this. We will pursue two Specific Aims in the R61 phase: Aim 1) Characterize the gene signatures and regulatory elements of mLC1 and mLC2 by profiling LCs during embryonic, young, and aging development at steady-state and repopulated LCs at inflamed state using scRNA-seq and scATAC-seq; Aim 2) Generate ATF3negEGFP reporter mice to fate-map ATF3+LC1 embryonic development and the dynamic change of ATF3+LC1 and ATF3-LC2 subset at steady state.
during adult and aging development and at inflammatory state and functionally characterize LC subsets in vitro by sorting ATF3EGFP+ LC1 and ATF3- LC2 cells and rederviving ATF3.loxp mice, which will be crossed with hLangerin-Cre mice to generate LCspecific/time induced ATF3KO for in vivo functional study. In the R33 phase, we will pursue the following Specific Aim: Aim 3) Functionally characterize ATF3+LC1 subset in vivo using LC-specific ATF3 deletion hLCcre.ATF3KO mice to evaluate the potential immune regulation function of ATF3+LC1 subset in the different disease models, including autoimmune vitiligo, melanoma, and fungi infection models. Our work will uncover the mystery of LC subsets with their specific functions, which will provide new insights into the biology of LCs and lead to the development of LC-based intervention strategies for skin diseases.

Principal Investigator: Qing-Sheng Mi, M.D., Ph.D.
microRNAs and NKT Cell Development and Function (R01AI119041)

Natural killer T (NKT) cells are an evolutionarily conserved subset of T cells that are developmentally and functionally distinct from conventional T cells. The ability to quickly secrete large quantities of a variety of cytokines upon activation enables NKT cells to be potent regulators of diverse immune responses. The deficiencies in NKT cell number and function have been linked to the development of many diseases. However, a significant gap remains in our understanding of how the development and function of NKT cells are precisely regulated. MicroRNAs (miRNAs), a recently discovered class of evolutionarily conserved small non-coding RNAs, negatively regulate the expression of protein-coding genes and thereby control essential biological functions and contribute to the development of many diseases. We were the first to report that the deletion of Dicer (a key enzyme for miRNA biogenesis) during hematopoiesis results in a significantly reduced NKT cell number and impaired NKT cell maturation and function, without alternating conventional T cell development in the thymus, suggesting that miRNAs are required for NKT cells. Our long-term goal is to understand how miRNAs regulate NKT cell development and function. While more than 1000 experimentally reported miRNAs, very few specific miRNAs are linked to NKT cells so far. Our objective here is to define specific miRNAs and their targets that regulate NKT cell development and function. Using miRNA arrays, we recently identified dynamic expression of miRNAs, including miR-155, and miR-17-92 cluster, during NKT cell development and activation. These findings plus our recent other report lead to our central hypothesis that these dynamically expressed miRNAs serve as critical regulators controlling NKT cell development and function through fine-tuning of specific target genes. Here we will further test this hypothesis. We will investigate how dynamic and miR-155 and miR-17-92 expression regulates NKT cell development and function using specific miRNA mutant mice with the gain or loss of miRNA gene. The results from proposed studies may not only illuminate the new immunological and molecular mechanisms underlying NKT cell development but may also facilitate the development of new and more efficient intervention strategies for autoimmune diseases, infection, and cancer based on the NKT cell therapy.

Principal Investigator: Qing-Sheng Mi, M.D., Ph.D.
Roles of HDAC3 in Epidermal Langerhans Cell Ontogeny and Function (R01AR069681)

Langerhans cells (LCs), the skin residing dendritic cells (DCs), form a contiguous immune network in skin and are involved in allergy, infection, cancer, and autoimmune disease development. However, the regulatory mechanisms involved in the development and functions of LCs have not been completely elucidated. Histone deacetylases (HDACs) are enzymes that regulate gene expression by modifying chromatin structure through removal of acetyl groups from target histones or directly deacetylating nonhistone proteins and represent a key epigenetic regulatory mechanism. HDAC inhibitors (HDI) are shown to have anti-tumor and anti-inflammatory effects in a variety of diseases, in which LCs play an important role. However, the mechanisms underlying the clinical effectiveness of HDI remain largely unknown. We recently reported that the inhibition of Class I/II HDACs by Trichostatin A (TSA) regulates the homeostasis and function of LCs in vitro and in vivo and modulates the non-coding miRNA expressions in LCs, while miRNAs also control LC development and function. Our preliminary data indicate that LCs express all Class I/II HDACs. To evaluate the role of individual HDACs in LC development and function, we generated knockout (KO) mice with selective deletion of HDAC3 (Class I) or HDAC4 (Class II) in epidermal LCs. Interestingly, LC number was significantly reduced in LC-HDAC3KO mice, but unaffected in LC-HDAC4KO mice. Furthermore, LC maturation and function were altered in LC-HDAC3KO mice. Thus, we hypothesize that HDAC3 is a key epigenetic component that controls LC development and function. In Aim 1, we will investigate the roles of HDAC3 in LC development and homeostasis, using LC-HDAC3KO mice for homeostasis after birth and using constitutive Csfr1r-specific HDAC3-deletion mice (Csfr1r-HDAC3) and
inducible Csf1r-specific HDAC3-deletion (Csfr1.Mer-HDAC3) mice for early embryonic LC development; Aim 2, we will investigate the roles of HDAC3 in LC function, using inducible LCER. HDAC3KO mice. In Aim 3, we will elucidate the molecular mechanisms and signaling pathways by which HDAC3 regulates LC development and function, by combining cDNA array, miRNA array and ChiP-Seq techniques. The proposed studies will uncover the epigenetic regulatory mechanisms of HDAC3 in LC development and function and may also elucidate new mechanisms for HDI therapy.

Principal Investigator: Qing-Sheng Mi, M.D., Ph.D.
Serum MicroRNA Biomarkers of Islet Autoimmunity (R01AI123258) Subcontract

Under Dr. Mi’s leadership, the team at Henry Ford Health System will perform miRseq profiles and quantitative miRNA analysis on serum samples using the Exiqon RT-PCR platform. Based on preliminary data, a custom panel of 188 microRNAs will be used. This strategy will allow greatly reducing the cost of measuring microRNAs by almost 50% and yet allow to study serum microRNA extensively; making it possible to measure a larger number of samples for increased statistical power. Over the course of the four-year program, we anticipate measuring microRNA levels in 600 serum samples from the DPT1 cohort, as described in the experimental plan. In addition to this, the team at Henry Ford Health System will perform miRseq to define potential candidates that may be missed by the Exiqon platform.

Principal Investigator: Li Zhou, M.D.
miRNAs Regulate Skin Langerhans Cell Ontogeny and Function (R01AR072046)

Langerhans cells (LCs), the skin residing dendritic cells (DCs), control both the induction of adaptive immunity, and immune tolerance in skin and are involved in variety of skin disease development. However, the regulatory mechanisms involved in the development and functions of LCs have not been completely elucidated. MicroRNAs (miRNAs), a class of non-coding small RNAs, are recognized as important regulators of protein-coding genes through the inhibition of mRNA translation. Using Cre-loxP Dicer deletion mouse models, our laboratory and others have reported that deletion of miRNAs by CD11c-Cre or hLangerin-Cre significantly reduced the number and interrupted the function of LCs, indicating that miRNAs are required for LC homeostasis and function after birth. While there are more than 1000 experimentally reported miRNAs, very few individual miRNAs are linked to LCs so far. We were the first to report that miR-150 and miR-223 differentially regulated LC-induced T cell proliferation and cytokine production. Most recently, our embryonic lineage-tracing studies showed that miRNAs, including miR-17-92cluster, regulate LC embryonic development. Furthermore, using miRNA arrays, we identified that mature LCs have a unique miRNA gene expression profile compared to immature LCs, and that miRNA expression is dynamically changed during LC embryonic ontogeny. These findings led to our central hypothesis that the dynamically changed miRNAs may serve as critical regulators controlling LC ontogeny, homeostasis and function through fine-tuning specific target genes. In Aim 1, we will investigate the roles of miRNAs in LC ontology and homeostasis. Constitutive or inducible Csf1r-specific individual miRNA mutant mice will be used for studying embryonic LC ontology and LC repopulation after inflammation, while LC-specific Dicer or individual miRNA mutant mice will be used for LC homeostasis after birth. In Aim 2, we will investigate the roles of miRNAs in LC function, inducible LC-specific Dicer or individual miRNA mutation mouse models will be used. In Aim 3, the direct target gene(s) of miRNAs and related signaling pathways involved in LC development and function will be investigated by the combination of RNA-seq, miRNA bioinformatics and related target functional validation strategies. The proposed studies will uncover the dynamic miRNA-mRNA regulation and related molecular mechanisms and signaling pathways that control LC development and function, which will not only provide new insight into the biology of LCs, but may also facilitate the development of LC-based intervention strategies for diseases.
Neurology

Principal Investigator: Jieli Chen, Ph.D.
Diabetic Stroke Cardiac Dysfunction; Treatment with CD133 + Exosomes (R01HL143432)

Cardiovascular complications are primarily responsible for the high morbidity and mortality in people with stroke and diabetes mellitus (DM). Cardiovascular diseases are roughly three times higher in patients with neurological deficits than in patients without neurological diseases. DM is a prominent risk factor for cardiovascular diseases and cerebral ischemic stroke. Our preliminary data show that ischemic stroke and type two DM (T2DM) each induces cardiac dysfunction, while T2DM animals subjected to ischemic stroke exhibit profound cardiac dysfunction compared to non-stroke T2DM mice or non-T2DM stroke mice. Therefore, there is a compelling need to develop therapeutic approaches specifically designed not only to reduce neurological deficits, but also to decrease cardiac dysfunction after stroke with diabetes. Our preliminary data indicate that treatment of stroke in T2DM mice with exosomes derived from human umbilical cord blood isolated CD133+/KDR+ cells (CD133+Exo) 3 days after stroke not only improves neurological and cognitive outcome, but also significantly improves cardiac function and increases heart microRNA (miR)126 and miR29b expression. In a novel and clinically relevant approach, based on our robust preliminary data, we propose to investigate the underlying cardioprotective therapeutic mechanisms of CD133+Exo treatment of stroke in T2DM mice, and we will test the hypothesis that miR126 and miR29b mediate CD133+Exo-induced cardiac protective effects in male and female mice in vitro and in vivo. Two Aims are proposed. Aim 1: To investigate the effect of cerebral ischemic stroke and stroke-related factors (age, sex and T2DM) on cardiac and neurological function in mice. To test the therapeutic effects of CD133+Exo treatment of T2DM-stroke in male, female and aged mice, time window, dose response, multiple doses and combination with anti-diabetic drug (Metformin) studies will be performed. Aim 2: To investigate the mechanism of CD133+Exo induced cardiac protective effects in male and female T2DM-stroke mice in vitro and in vivo. We will focus on miR126 and miR29b, and will test: 1) whether CD133+Exo treatment of T2DM-stroke increases heart and serum miR126 or miR29b levels; 2) whether increasing miR126 or/and miR29b expression in heart or/and serum mediates the CD133+Exo induced cardiac beneficial effects in male and female T2DM-stroke mice; 3) whether the miR126/Spred-1 and/or the miR29b/DPP4 signaling pathways mediate CD133+Exo treatment induced myocardocyte protection of cultured cardiomyocytes. A major significance of our investigations is that it opens up important and novel ways to understand how exogenously administered CD133+Exo communicate with and alter heart cells by means of miR delivery to thereby activate endogenous cardiac protective events. This proposal is highly clinically relevant and if successful, it will significantly impact the treatment of stroke, diabetes, and cardiac dysfunction. Importantly, this proposal will elucidate novel mechanisms of action and generate therapeutic targets for CD133+Exo treatment of cardiac dysfunction after stroke with T2DM in male, female and aged mice.

Principal Investigator: Michael Chopp, Ph.D.
miR-17-92 Exosome Treatment of Stroke (R01NS088656)

Exosomes, small lipid microvesicles (30-150 nm), are active biological containers, which transport regulatory genes and proteins between cells and form a major biological communication conduit, facilitating a plethora of biological responses. The regulatory molecules contained in the exosomes include microRNAs (miRNAs), short (22-25 nt) non-coding RNAs which regulate gene translation and play primary roles in mediating a vast range of biological functions. In this proposal, based on strong preliminary data, we propose to manufacture a distinct exosome population which contains increased levels of the miR-17-92 cluster as a proof-of-principle and a mechanistic demonstration of a new method of treating stroke and possibly other neurological diseases and injury. We test the premise, that by modulating their miRNA content, exosomes can be designed to enhance plasticity of axons and thereby further promote neurological recovery post stroke. Success of this novel approach may lead to a new designer-based paradigm for the treatment of stroke and neurological disease. The following Specific Aims and associated Hypotheses are proposed: Specific Aim 1: To employ exosomes derived from multipotent mesenchymal stromal cells (MSCs) to treat stroke in order to enhance neurovascular remodeling and thereby, functional recovery post stroke. Hypothesis: Exosomes, derived from MSCs when administered to rats after stroke promote neurovascular remodeling which improves functional outcome. Specific Aim 2: To alter specific miRNAs contained...
within exosomes generated by MSCs as a means to enhance axonal plasticity and neurological recovery post stroke. Hypothesis: Administration of exosomes with increased miR-17-92 cluster to rat post stroke promotes axonal remodeling and enhances functional outcome. There are multiple layers of innovation in our application: we generate biological exosome carriers tailored for specific miRNAs; we use these exosomes to treat stroke, without the administration of exogenous cells; we employ electrophysiological methods, laser capture, fiber track tracing, a battery of neurological tests, and an array of novel approaches, e.g. microfluidic chambers, and ex vivo slice cultures, to mechanistically determine the molecular pathways of the target exosomes which mediate axonal outgrowth. Development of this designer exosome-based therapy also serves as a prototype for capitalizing on the characteristics of exosomes to transport specific miRNAs and for the manufacture of designer exosomes. Developing a therapy for stroke that is exosome-based opens up a wide variety of means to deliver targeted regulatory genes to enhance multifaceted aspects of central nervous system (CNS) plasticity and to amplify neurological recovery for neural injury and neurodegenerative diseases.

Principal Investigator: Xu Cui, Ph.D.
ABCA1 Regulates White Matter Remodeling and Oligodendrogenesis after Stroke (R01NS092917)

Stroke is a major cause of white matter (WM) damage which induces long-term disability. There is limited WM remodeling in the adult brain. Many neuroprotective treatments of stroke have failed in clinical trials because they cannot protect WM. Therefore, there is a compelling need to investigate the mechanism underlying WM remodeling and oligodendrogenesis of the adult brain and to develop effective long-term stroke therapy. Cellular cholesterol modulates axonal and dendritic outgrowth and is required for myelination. The level of HDL-cholesterol is related to the progression and recovery of stroke patients. ATP-binding cassette transporter A1 (ABCA1) is a major cholesterol transporter and plays critical roles in regulation of HDL-cholesterol and ApoE synthesis and metabolism in the central nervous system. Brain specific- ABCA1 deficient (ABCA1-B/-mice have very low brain HDL-cholesterol/ApoE level and exhibit neuronal ultrastructure changes and functional deficits. Both HDL-cholesterol and ApoE increase neurite outgrowth in culture conditions. Our preliminary study shows that ABCA1-B/-mice have increased WM damage and reduced oligodendrogenesis and exacerbated neurological functional deficits after stroke. Primary cultured neurons derived from ABCA1- B/-mice show decreased neurite outgrowth, which can be attenuated by HDL treatment. ABCA1-B/-astrocyte-conditioned media also decreased wild type neurite outgrowth after hypoxic ischemia. Therefore, we propose the following three specific aims: Aim1 to investigate whether brain-deficient in ABCA1 exhibits decreases in WM-remodeling and axonal growth after stroke. ABCA1-B/-B and floxed-control mice will be subjected to stroke, WM-changes and oligodendrogenesis will be measured. Aim2 To investigate molecular mechanism underlying ABCA1 in regulation of WM-remodeling and oligodendrogenesis after stroke, we will examine whether ABCA1 regulates brain HDL and ApoE level, and whether brain HDL and ApoE levels mediate ABCA1-induced WM-remodeling and oligodendrogenesis after stroke. Aim3 to investigate cellular mechanisms of ABCA1 in regulation of WM-remodeling and oligodendrogenesis, we will examine neurons and oligodendrocytes and the cross talk of astrocytes with neurons and oligodendrocytes on ABCA1-induced WMremodeling and oligodendrogenesis in vitro and in vivo. We expect that ABCA1 deficient brain will exhibit significant decreases in HDL and ApoE level and decreases WM-remodeling and oligodendrogenesis as well as reduced functional outcome after stroke. The level of HDL/ApoE in brain or cerebrospinal fluid will, at least partially, mediate ABCA1-induced WM-remodeling and oligodendrogenesis in the ischemic brain after stroke. To our knowledge, no one has investigated the functional effect of ABCA1 on oligodendrogenesis and WMremodeling post- stroke recovery, especially by using ABCA1- B/-mice. The new insights gleaned from this study will contribute to our understanding of the beneficial role of ABCA1/HDL-C/ApoE in brain plasticity which will impact development of rational restorative approaches to improve neurological outcome for stroke patients.

Principal Investigators: James Ewing, Ph.D., Neurology and Stephen Brown, Ph.D., Radiation Oncology
MRI Signatures of Response to High-Dose Radiotherapy in Rat Models of Cerebral Tumor (R01CA218596)

In some cases, e.g. small brain tumor metastases, responses to single or multiple fraction high-dose radiation therapy (HD-RT) have been remarkable, suggesting that HD-RT tumor control is at least as effective as biologically equivalent doses of conventional fractionated radiation therapy
(CF-RT), even in radioresistant tumors. Although the mechanism for its effectiveness is not well understood, HD-RT is becoming accepted practice for a variety of tumors, including brain tumors. Our recent preclinical study using MRI measures of short-term changes in tumor physiology after HD-RT in a small-animal model of cerebral tumor suggests a physiological response that includes vascular effects but is multifactorial and temporally variable. Hypothesizing that these short-term changes may both explain the increased effectiveness of HD-RT, and serve as a predictor of long-term response, we propose to investigate the relationship between short-term physiological changes after HD-RT and long-term outcome as a result of that therapy. In counterpoint, we will also study physiological changes during and after CF-RT. Detailed poroelastic modeling is proposed to generate a map of local solid and fluid parameters (stress, flow) that will help explain short-term changes in physiology. Aim 1 studies short-term changes in measures of tumor physiology as predictors of response. Aim 2 describes the behavior of these same measures over the course of CF-RT. Our long-range goals are to develop noninvasive biomarkers of response that predict tumor control after HD-RT and CF-RT, and to describe physiological changes and related biomarkers that might be used to optimize the order and timing of RT and adjuvant chemotherapies.

**Principal Investigator: Shailendra Giri, Ph.D.**

**Novel Regulation and Targeting of Macrophages Metabolism in Neuroinflammatory Disorders (R01AI144004)**

Myeloid cells play a critical role in CNS demyelination and axonal destruction of multiple sclerosis (MS) and experimental autoimmune encephalomyelitis (EAE). The early phase of the disease is characterized by the presence of pathogenic activated macrophages (M1 type), while the recovery phase is associated with alternatively activated macrophages (M2 type) which release anti-inflammatory cytokines that resolve the pathogenic inflammation. Activated M1 macrophages depend on glycolysis to boost biosynthetic pathways to produce inflammatory mediators. However, anti-inflammatory M2 macrophages rely primarily on mitochondrial respiration. Adenosine monophosphate-activated protein kinase (AMPK) regulates energy metabolism, and thus controls the balance between glycolysis and mitochondrial respiration. We previously reported that AMPKa1 knockout (KO) mice develop severe EAE indicating AMPK activation is protective, yet the molecular mechanism by which AMPK regulates EAE disease progression is not known. AMPKa1-KO macrophages exhibit a hyperinflammatory phenotype and have a lower rate of metabolism. AMPKa1-KO macrophages also show glycolysis-tricarboxylic acid (TCA) cycle remodeling, which results in an imbalance in the levels of the endogenous metabolites, succinate and itaconate, which regulate pro-and anti-inflammatory macrophage functions, respectively. Their levels are tightly controlled by succinate dehydrogenase (SDH) and immune responsive gene 1 (IRG1), respectively. We hypothesize that the loss of AMPKa1 remodels the glycolytic-TCA pathway causing an imbalance in the levels of succinate and itaconate, which promotes an M1 phenotype over an M2 phenotype. This, in turn, promotes Th17 cells and suppresses regulatory cells leading to a hyperinflammatory CNS immune response and CNS tissue damage. To test our hypothesis, we have generated monocyte-specific AMPKa1 KO and macrophage-specific, constitutively active AMPKa1T172D transgenic mice. In **Aim 1**, we will examine how the loss or gain of function of AMPKa1 in macrophages regulates M1 versus M2 macrophage polarization and consequently, Th17 and Tregs differentiation and disease outcomes. Studies under **Aim 2** will elucidate the mechanism by which the loss of AMPKa1 reprograms glycolysis-TCA metabolism leading to an imbalance of succinate and itaconate metabolites in macrophages, which in turn, determine the macrophage phenotype. The proposed study is expected to have a positive impact by elucidating the metabolic regulatory mechanism responsible for macrophage plasticity during disease and investigating AMPKa1 as a potential therapeutic target for MS. Our innovative genetic mouse models and precise metabolomics approach will allow us to identify the apparent rewiring of cellular metabolic pathways specific to AMPKa1 in hyperinflammatory cells. Ultimately, this process could be exploited to tailor novel therapeutic strategies to resolve or limit autoimmune inflammation in the CNS.

**Principal Investigator: Shailendra Giri, Ph.D.**

**Endogenous Metabolite Restricts GM-CSF Signaling Pathway in Pathogenic Macrophages to Ameliorate CNS Autoimmunity (1R01NS112727)**

Identifying a therapeutic option that can modulate the innate immune response without generally suppressing the immune system as a whole has been a key barrier to improving treatment for patients with MS. Using metabolic profiling, we have reported that resolvin D1 (RvD1), a pro-resolving lipid metabolite of
omega-3 polyunsaturated fatty acids, is significantly decreased in the plasma of patients with MS. Consistent with this finding, MS patients have lower levels of omega-3 metabolites, which are precursors of resolvis, compared to healthy controls, a finding that has been replicated in animal models of the disease, experimental autoimmune encephalomyelitis (EAE). Notably, we found that daily supplementation with RvD1 significantly attenuated clinical symptoms in both chronic and relapsing remitting EAE. These data are provocative for their translational potential, particularly because the immune system is not depressed by RvD1 treatment as it is with steroids and most other MS therapies. The immunomodulatory effect of RvD1 is mediated through its receptor, formyl peptide receptor 2 (FPR2), leading to modulation of AMP-activated protein kinase (AMPK), an important regulator of cell metabolism. In other human disease models, the RvD1-FPR2 signaling cascade protects by inducing an anti-inflammatory phenotype in macrophages. However, the mechanism affording this protection remains elusive. Granulocyte-macrophage colony-stimulating factor (GM-CSF) is a key player both in the pathology of both MS and EAE, promotes an inflammatory environment and neuronal damage. There is a growing interest in inhibiting the pro-inflammatory effects of GM-CSF signaling as a therapeutic target in MS. In our preliminary work, we found that RvD1 treatment inhibited GM-CSF signaling in macrophages and that this inhibition was AMPK-dependent. However, how RvD1 affects AMPK activity and GM-CSF signaling to attenuate EAE is unclear. Our long-term goal is to identify natural endogenous signaling mechanisms that can be harnessed to treat autoimmune diseases, particularly MS. Our overall objective here is to determine the mechanism of action of RvD1 in resolving inflammation and disability in mouse models of MS. Our central hypothesis is that RvD1 attenuates EAE disease progression by abrogating GM-CSF signaling resulting in the polarizing of pro-inflammatory macrophages into an anti-inflammatory phenotype. And the underlying mechanisms of macrophage phenotype switch are through FPR2-AMPK-dependent metabolic reprogramming. To test this hypothesis our specific aims are: 1) to identify the effects of RvD1 on the cellular phenotype and function of CNS-infiltrating macrophages in EAE; and 2) to determine the effects of RvD1 on metabolic reprogramming in macrophages. We will address these aims with a combination of immunological, biochemical and innovative metabolomic approaches that are already well in hand. The proposed studies will form the foundation for the development of innovative therapeutic strategies to resolve inflammation during MS with no side effects and will likely apply to other diseases involving pathogenic activation of the immune system.

Principal Investigator: Quan Jiang, Ph.D., and Jieli Chen, Ph.D.
Investigation of D4-F Effects on Neurovascular Remodeling after Diabetic Stroke (R01NS097747)

Ischemic stroke patients with Diabetes mellitus (DM) exhibit a distinct risk-factor and etiologic profile and a worse neurovascular prognosis than non-DM patients. Therefore, there is a compelling need to investigate neurovascular changes after stroke in the DM and non-DM population and to develop therapeutic approaches specifically designed to reduce neurological deficits after stroke. Type 2 diabetes (T2DM) constitutes 90% of diabetic patients and is associated with low high-density lipoprotein cholesterol (HDLC), impairment of the anti-oxidative capacity of HDLC, low phosphorylation of endothelial nitric oxide synthase (p-eNOS), and with reduced ATP-binding cassette transporter A1 (ABCA1) gene expression. D-4F is an economical apolipoprotein A-I (ApoA-I) mimetic peptide, presently employed in clinical trials to reduce coronary atherosclerosis in patients with acute coronary syndrome. However, the therapeutic effects of D-4F in post-ischemic stroke have not been investigated. Our preliminary data show that D-4F treatment of stroke starting 2h or 24h after ischemic stroke improves recovery of neurological function in both T2DM and non-DM mice and also increases p-eNOS and ABCA1 in the ischemic brain. In a novel and clinically relevant approach, based on our robust preliminary data, we propose to use D-4F in the treatment of stroke in the non-DM and T2DM population in mice. We seek to develop D-4F as a novel neurorestorative therapy to reduce white matter (WM) dysfunction and vascular damage, in T2DM and non-DM mice when treatment is initiated at 24h after onset of ischemic stroke. In addition, most development of stroke treatments has focused on young adult animals, but not on old animals, the prevalent population with stroke. Increased age also increases neurological impairment after stroke. We have also developed and implemented multimodality MRI imaging which can dynamically monitor neurovascular remodeling in both the animal and the patient. In the current study, we will measure WM and vascular changes and elucidate the mechanisms of action of D-4F in young adult and aged animals with and without T2DM after stroke. Our hypothesis is that D-4F increases ABCA1 and p-eNOS signaling activity which mediates vascular and WM remodeling and in concert improve functional outcome after stroke. We, therefore, propose two highly integrated and longitudinally designed Specific Aims. Aim 1 will investigate the delayed (24h after stroke) therapeutic effects of D-4F in non-DM and T2DM in young adult and aged mice after stroke. The differences in cerebral WM and vascular changes, and neurological
functional outcome after stroke between non-DM and T2DM mice treated with or without D-4F will be analyzed. MRI will be employed to measure the dynamics of neurovascular reorganization underlying therapeutic response and recovery. In Aim 2, using eNOS knockout mice and specific loss of brain ABCA1 mice, we will investigate the mechanisms by which D-4F promotes neurovascular remodeling and hence, neurological recovery. The long-term objective of this RO1 is to develop a neurorestorative treatment for stroke in patients with or without diabetes.

Principal Investigator: Quan Jiang, Ph.D.
Glymphatic and Cognitive Impairment of Aging and Diabetes (RF1AG057494)

The objective of this application is to investigate glymphatic impairment and cognitive deficits during progression of aging with and without diabetes. Emerging data indicate that the glymphatic system in the brain mediates the cerebrospinal fluid (CSF)-interstitial (ISF) exchange and solute clearance from the brain parenchyma. However, despite the well-described dysfunction of the glymphatic system in the development of neurodegenerative conditions, there is still no reported study that focuses on the role of the glymphatic system in the development of cognitive impairment during aging and with type-2 diabetes (DM). Using noninvasive MRI methodologies to investigate cerebral solute waste clearance in middle-age control and type-2 diabetic (DM) rats, we have found increased impairment of the glymphatic system, as indicated by reduced clearance of interstitial Gd-DTPA in brain parenchyma, primarily in the hippocampus and hypothalamus in DM rats (Fig.2&3). In parallel, 3D confocal microscopic analysis of the brain-wide distribution of fluorescent tracers revealed increased delayed clearance of ISF in the hippocampus and hypothalamus from DM rats (Fig.2&3). Impairment of the glymphatic system in DM rats was shown to be highly correlated with cognitive deficits as measured by an array of cognitive tests including the Morris Water Maze (MWM) for hippocampal related learning and memory. Importantly, histopathological analysis shows that delayed clearance of interstitial solutes is associated with sporadic cerebral microvascular thrombosis in the hippocampus 2 months after hyperglycemia (15 months from birth), while extensive microvascular thrombosis and para-vascular accumulation of beta-amyloid (Aβ) are detected at 4 months after induction of hyperglycemia (17 months from birth), suggesting that the impairment of the glymphatic system leads to Aβ accumulation. Collectively, our preliminary data, for the first time, demonstrate that non-invasive MRI methodologies can detect DM-induced early impairment of the glymphatic system which is highly correlated with hippocampal related dysfunction of learning and memory. Based on our novel preliminary data, we will employ MRI and 3D confocal microscopy to evaluate and quantitatively measure kinetic clearance parameters of the glymphatic system during progression of aging with and without DM (Aim 1). We will then investigate whether impairment of the glymphatic system predicts cognitive dysfunction, the sensitivity and association between impairment of the glymphatic system, the onset of brain vascular dysfunction, and cognitive deficits during aging with and without DM (Aim 2). Data generated from this application will provide new insights into aging and age-matched DM associated impairment of the glymphatic system and the relationship of the glymphatic system with vascular and cognitive dysfunction.

Principal Investigator: Quan Jiang, Ph.D.
Interaction Between Glymphatic and Vascular Systems for Waste Clearance in Brain (R01NS108463)

The objective of this application is to first develop and validate micro vessel measurement for the entire brain to enhance detection sensitivity of micro vessels by ten-fold using superparamagnetic iron oxide (SPIO) enhanced susceptibility weighted imaging (SWI, SPIO-SWI) and then to investigate the interaction between glymphatic and vascular systems for waste clearance in the diabetic brain. Emerging data indicate that the glymphatic system in the brain mediates the cerebrospinal fluid (CSF)-interstitial (ISF) exchange and solute clearance from the brain parenchyma and plays an important role in neurological diseases. Despite many milestone achievements, conclusive findings on the solute efflux pathways are relatively limited. Consequently, the interaction between vascular and glymphatic systems on waste clearance, especially with neurological diseases, is unclear. The paucity of research into the efflux pathway may be attributed in part to technical difficulties, such as the challenging need to perform minimally invasive in-vivo, ultra-high detection sensitivity for tube-shaped influx and efflux pathways, and whole brain imaging. Although MRI can overcome the weak points of two-photon confocal microscopy to provide non-invasive whole brain in-vivo imaging of the glymphatic system, conventional MRI sensitivity is insufficient for the required spatial resolution for investigating micro vessels of glymphatic and vascular systems. We have developed highly sensitive MRI methods (Fig. 1) which significantly improve the detection sensitivity of small vessels by using the combination of high susceptibility...
of MRI agents with blooming effects7-9. The new methods provide excellent tools for investigating the efflux pathways of waste clearance under normal and pathophysiological conditions. Three efflux routes have been recently proposed and solutes in the brain could reach the lymphatic network by the olfactory bulb across the ethmoid plate10, 11 or by functioning conventional lymphatic vasculature in the meninges12. We found that tracer concentration in the venous system significantly increased with diabetes (Fig. 9), thus adding a new route for brain waste clearance. Based on our novel preliminary data and published studies by others, we hypothesize that, the newly developed SPIO-SWI technique significantly increases detecting sensitivity of micro vessels in both vascular and glymphatic systems, and the efflux pathways of waste clearance with and without diabetes can be identified and investigated using this optimized SPIO-SWI method. To test these hypotheses, we will first (Aim 1) further develop, optimize and validate SPIO-SWI techniques to enhance the detection sensitivity for both vascular and glymphatic micro vessels. We will perform computer simulation, optimize SWI technique and experimental conditions in animal studies and then validate USPIO-SWI technique by LSCM measurements. We will then (Aim 2) investigate the interaction between vascular and glymphatic systems for waste clearance in diabetic brain using the optimized USPIO-SWI technique. Data generated from this application will provide new insights into the efflux pathways between glymphatic and vascular systems in diabetic brain.

Principal Investigator: Jaspreet Singh, Ph.D.
Plasma Biomarkers of Cerebral Disease in X-Linked Adrenoleukodystrophy (R21NS104560)

There are no biomarkers to predict the onset of fatal demyelinating phenotypes in males with inherited X-linked adrenoleukodystrophy (X-ALD) disease. 60% of male X-ALD patients develop demyelination in childhood (5-12 years; ALD) or in adulthood (25-35 years; cerebral adrenomyeloneuropathy, cAMN). On average, cAMN and ALD are fatal in 2 to 5 years of onset. The primary genetic defect in X-ALD (mutation/deletion in ABCD1 gene) and the biochemical defect (accumulation of very long chain fatty acid; C>22:0 in plasma and tissues) cannot predict the onset of ALD or cAMN. Our long-term goal is to dissect microRNA (miRNA) and metabolic pathways underlying neurodegeneration in X-ALD. The objective of this application is to identify plasma miRNA and metabolite biomarkers predictive of progression to fatal cAMN and ALD in X-ALD males. Plasma exosome miRNAs and circulating metabolites have been used as diagnostic and prognostic biomarkers for neurodegenerative diseases. No plasma (or other biofluid) miRNA and metabolite biomarkers have been explored for the fatal X-ALD phenotypes. Our preliminary proof-of-concept data, with next generation sequencing and untargeted metabolomics, identified differential miRNA and metabolites between healthy-control and ALD- phenotype postmortem brain. Within the ALD brain white matter, unique miRNA and metabolite changes were recorded between distant normal looking areas and areas adjacent to the plaque suggesting an association with disease progression. Our central hypothesis is that metabolomic and miRNA analysis in retrospective plasma samples, collected both before and at the time of detection of demyelination in the patients, will provide biomarker(s) predictive of fatal cAMN and ALD progression in X-ALD males. To test our hypothesis we propose two specific aims: 1) Identify a plasma metabolome and miRNA signature for the cAMN phenotype. 2) Define a plasma metabolome and miRNA signature for the ALD phenotype. We will take advantage of a large cohort of control, non-converting AMN, cAMN (pre and post) and ALD (pre and post) plasma samples already available in the biorepository of the Moser Center for Leukodystrophies, Kennedy Krieger Institute, Baltimore, for discovery and validation. This proposal is innovative, because it departs from the status quo by identifying novel plasma regulatory (miRNA) and active (metabolite) biomarkers predictive of cAMN and ALD. The proposed research is significant because our pre-post design will identify plasma biomarkers able to predict disease course before the onset of fatal cAMN or ALD. X-ALD was added to the Recommended Uniform Screening Panel in February 2016, a federal list of all genetic diseases recommended for state newborn screening programs. This study will nominate and validate plasma biomarkers that have the potential to provide an effective monitoring tool for identified X-ALD infants. This study will lay the foundation for our future, large-scale, prospective clinical trial studies using novel plasma miRNA and metabolite biomarkers to predict fatal cAMN and ALD phenotypes. In collaboration with Kennedy Krieger Institute we will apply for funding for these future clinical trials from NINDS.
Vascular dementia (VaD) is common in patients after a stroke or after a series of mini-strokes and results from several mechanisms, one of which involves injury to blood vessels supplying deep white matter (WM) of the brain resulting in silent, multifocal, brain microinfarcts, vascular dysfunction, decrease in cerebral blood flow, and cerebral parenchymal cell damage. Extensive WM damage such as vacuolization, rarefaction, and demyelination in the periventricular region have been reported in patients with VaD. There is a critical need to develop therapeutic strategies for VaD that identify and target key pathophysiological events driving axonal/WM damage and cognitive deficits. The therapeutic effects of Vasculotide, an Angiopoietin-1 mimetic peptide, in VaD have not been investigated. Our preliminary in-vitro studies show that Vasculotide treatment can dose dependently increase axonal outgrowth in primary cortical neurons (PCN). In male retired breeder rats subjected to a multiple microinfarction (MMI) model of VaD, Vasculotide treatment initiated at 24 hours after MMI, significantly decreases axonal/WM injury and improves long term cognitive outcome. In a novel and clinically relevant approach, based on our robust preliminary data, we propose to use Vasculotide for the treatment of MMI induced VaD in male and female middle-aged rats (10-12 months old). We seek to develop Vasculotide as a therapeutic agent to decrease vascular dysfunction and axonal/WM injury, decrease inflammatory responses, attenuate glymphatic dysfunction and improve cognitive outcome. By affecting gene regulation, microRNAs (miRs) are involved in most biological processes and act as molecular rheostats that fine-tune and switch regulatory circuits governing tissue repair, inflammation, hypoxia-response, and angiogenesis. Elucidation of the role of miRs in VaD pathogenesis, and identification of key miRs that can potentially serve as therapeutic targets in VaD are lacking. We hypothesize that Vasculotide treatment induced vascular and axonal/WM remodeling; anti-inflammatory responses and cognitive recovery are mediated via modulation of key miRs and their target gene expression. Therefore, we propose three highly integrated and longitudinally designed Specific Aims. In Aim 1, we will perform dose-response studies and investigate the safety and long-term cognitive outcome of Vasculotide treatment in middle-aged, male and female rats subject to MMI model of VaD. In Aim 2, we will investigate the therapeutic effects of Vasculotide on vascular remodeling, axonal/WM remodeling, synaptic plasticity, inflammatory responses and glymphatic waste clearance pathway in middle-aged rats subject to MMI. In Aim 3, using “gain or loss” of brain miR-145 and miR-124, we will test whether Vasculotide treatment induced therapeutic effects after MMI in rats are mediated via the miR-124/Interleukin-6 and miR-145/Aquaporin-4/ATP-binding cassette transporter A1 (ABCA1) signaling pathways. The long-term objective of this R01 application is to develop a novel treatment for VaD.

Principal Investigator: Poornima Venkat, Ph.D.
Vasculotide Promotes Cognitive Improvement in Rats with Vascular Dementia (1R01AG063750)

Stroke is one of leading causes of death and disability worldwide, mainly affecting elderly. Tissue plasminogen activator (tPA), the only Food and Drug Administration (FDA) approved treatment, is limited in its use to < 8.5% of stroke patients. Therefore, there is a compelling need to develop new and broader utility therapies for acute ischemic stroke. Vepoloxamer is a well characterized proprietary amphipathic copolymer with rheological properties, which is currently under investigation in a global phase III clinical trial for patients with sickle cell disease. Our preliminary studies demonstrate that administration of Vepoloxamer in combination with tPA 4h after embolic stroke facilitates recanalization and thrombolysis reduces ischemic neuronal damage and improves neurological outcome but does not increase cerebral hemorrhage in young adult rats. We also found that platelet-derived exosomes contribute to the therapeutic effect of Vepoloxamer on enhanced tPA-thrombolysis. In this application, we propose to investigate effect of Vepoloxamer in combination with tPA on acute stroke and molecular mechanisms underlying the combination therapy on the thrombolysis and neurovascular function in the aged male and female rats. Data generated from this application may provide a novel and potentially useful treatment strategy for patients with acute stroke.

Principal Investigator: Li Zhang, M.D.
Combination Treatment with Vepoloxamer and tPA for Acute Stroke (R01NS102744)

Large cerebral vessel occlusion is the most disabling and life-threatening form of ischemic stroke. Human stroke primarily occurs in late middle age and beyond. Approximately two thirds eligible patients treated with tPA experience incomplete reperfusion. Thrombectomy is now also a standard of care for treatment of acute stroke with large vessel occlusion. However, recanalization of the occluded large vessels by thrombectomy
only leads to ~71% of patients achieving improved tissue reperfusion, often incomplete. In addition, due to unfavorably large ischemic cores, many patients with large artery occlusion are not eligible to receive tPA or thrombectomy. Patients with reperfusion of the ischemic tissue are closely associated with good clinical outcome. Thus, there is a compelling need to develop therapies in combination with tPA and thrombectomy to enhance cerebral perfusion and thereby augment the therapeutic efficacy of tPA and thrombectomy monotherapies. Also, therapies to block ischemic core expansion will increase numbers of patients who would be eligible to receive tPA and thrombectomy. Using rat models of embolic middle cerebral artery occlusion (eMCAO) and transient MCAO (tMCAO, ischemia/reperfusion), we found that exosomes derived from cerebral endothelial cells (CEC-exos) in combination with tPA after eMCAO or CEC-exos given upon reperfusion after tMCAO substantially increased recanalization and downstream cerebral blood flow (CBF), and reduced blood brain barrier (BBB) leakage and infarction compared to tPA or tMCAO alone. Exosomes are nano-vesicles that contain lipids, proteins, and RNAs including microRNAs (miRs). Our preliminary data suggest that exosomal cargo miRs likely contribute to the therapeutic effect of CEC-exos in combination with tPA on acute stroke by acting on cerebral endothelial cells to suppress proteins that promote thrombosis and BBB disruption. We thus propose to develop CEC-exo therapy as an adjunctive treatment to enhance tPA and thrombectomy treatments of acute ischemic stroke. Aim 1 is to investigate whether the CEC-exo therapy as an adjunctive treatment enhances tPA and thrombectomy treatments in aged rats after large artery occlusion. Aim 2 is to investigate whether CEC exosomal cargo miRs contribute to CEC-exos-amplified thrombolysis leading to reduction of neurovascular damage. Aim 3 investigates whether a special set of CEC-exo cargo miRs contribute to the therapeutic effect CEC-exos on stroke-induced neurovascular damage by suppressing a network of pro-BBB leakage and thrombotic genes. Accomplishing these aims will potentially lead to development of a mechanistically based exosome therapy as an adjunctive treatment to enhance tPA and thrombectomy treatments of acute ischemic stroke, leading to improvement in the neurological outcome.

Principal Investigator: Zhenggang Zhang, M.D., Ph.D.

Exosomes and Platinum-Induced Peripheral Neuropathy (R01NS111801)

Platinum-based drugs are commonly used to treat cancers. Platinum drugs are the first line therapy for ovarian and colorectal cancers. However, chemotherapy-induced peripheral neuropathy (CIPN) is one of the most common complications. More than 70% of the patients receiving oxaliplatin are affected by neuropathy. Oxaliplatin induces two symptoms of peripheral sensory neuropathy; an acute and transient cold-aggravated, and a chronic form that has onset after multiple exposures to the drug and does not disappear with drug cessation. The neurotoxicity often leads to platinum drug dose reductions, compromising efficiency of platinum drugs to suppress tumor progression. On an average of 6 years after chemotherapy, 47% of women still reported symptoms of CIPN. Studies to develop a neuroprotective agent have, to date, been unsuccessful to reduce CIPN. There is an imperative need to develop new therapies to CIPN. Challenges to develop such therapies include that a therapy needs not to impede antitumor efficacy, but to effectively inhibit CIPN. Our preliminary data demonstrated cerebral endothelial cell derived exosomes (CEC-exos) abolish oxaliplatin-induced peripheral neuropathy in tumor bearing mice and sensitize oxaliplatin on cancer cell killing. Exosomes are nanovesicles and mediate intercellular communication by transferring cargo proteins, lipids, and genomic materials including mRNAs and microRNAs (miRNAs) between source and target cells. We found that treatment of the tumor bearing mice with CEC-exos along with oxaliplatin induces a network of miRNAs/miRNAs in sciatic nerves that exerts neuroprotection in sciatic nerves and DRG neurons but triggers a distinct miRNAs/miRNAs network in tumor to promote cancer cell death. We, thus, hypothesized that CEC-exos mitigate peripheral neurotoxicity induced by platinum drugs and that CEC-exos enhance the anti-cancer efficacy of platinum drugs on tumor cells. Three specific aims are proposed to test this overall hypothesis. Aim 1 is to investigate the efficacy of the CEC-exos on ameliorating platinum drug-induced peripheral neurotoxicity and on improving the treatment of tumor. Aim 2 is to investigate molecular mechanisms underlying the therapeutic effect of CEC-exos on platinum drug-induced peripheral neuropathy with a focus on the interaction between CEC exosomal miRNAs and their target proteins in axons and DRG neurons. Aim 3 is to investigate molecular mechanisms underlying the effect of CEC-exos on sensitizing tumors to platinum drugs with a focus on the interaction between CEC exosomal miRNAs and their target proteins in tumor cells. Accomplishing these aims will potentially lead to development of a new CEC-exo based therapy for CIPN, leading to improvement in the quality of life and possibly cure of cancers.
Neurosurgery

Principal Investigator: Meser Ali, Ph.D.
Treatment of Glioma with Nanocombretastatin with MRI Monitoring (R01CA206190)

Glioblastoma (GBM) is a highly aggressive hypervascularized brain tumor characterized by high recurrence rates and poor prognosis despite advanced treatment. The vasculature of GBM is fundamentally different from that of normal vasculature and offers a unique target for anti-cancer therapy. Therefore, direct targeting of tumor vasculature with vascular disrupting agents (VDAs) is distinctly different from antiangiogenic strategies and offers a complementary approach to standard therapies. Combretastatin A4 (CA4) is a potent vascular disrupting drug. CA4 induces rapid shutdown of tumor blood supply, typically promoting a necrosis at the core of the tumor but leaves a rim of viable tumor cells at the periphery which can then rapidly re-grow. However, CA4 is not effective in inducing necrosis at the core of GBM tumor. The ineffectiveness of small molecule chemotherapy drugs in treating malignant brain tumors has been attributed to the blood-brain barrier (BBB) being a significant impediment to the transvascular extravasation of drug fraction across the barrier into the extravascular compartment of tumor tissue and the high tumor interstitial fluid pressure also presents an additional delivery barrier. Nanotechnology is already benefiting to deliver drugs across the BBB and into brain tumors. We have engineered a nano sized polymeric CA4 conjugate which demonstrates high water solubility. Preliminary intravenous (i.v.) delivery of G3-CA4 in an orthotopic glioma model demonstrated necrosis at the core of the tumor leaving a rim of viable tissue. By applying the designed nano-prodrug strategy and tumor-specific prodrug activation mechanism, we observed the true success of inducing necrosis at the core of the tumor in an orthotopic U-251 glioma animal model first time. Tumor-VDAs have significant potential when combined with cytotoxic chemotherapy and radiotherapy in treating other tumor models. Combined treatment with radiation is attractive, as radiation therapy (RT) represents a standard of care and RT should effectively kill the well-oxygenated cancer cells in the well-perfused tumor rim. We have shown that GBM cancer stem cells are sensitive to radiation exposure in culture and a single dose of 50Gy irradiation yielded necrosis in primary GBM rat model. Therefore, this study is extended to include SRS and standard cytotoxic temozolomide (TMZ) therapies with G3-CA4. We hypothesize that the combination of G3-CA4 with SRS and TMZ will show synergistic cytotoxic effect in clinical relevant primary GBM model. Our objectives of the proposed research are A) To incorporate CA4 molecules with dendrimer-based nanoparticles (G3-CA4) that demonstrates full solubility in aqueous media, B) To determine the efficacy and safety of small molecule CA4, CA4-P and G3-CA4 nano-prodrug in U251 glioma tumor model, C) To determine the efficacy and safety of G3-CA4 alone or in combination with SRS in primary GBM, D) To determine the efficacy and safety of a combined G3-CA4 and standard TMZ therapy in primary GBM model. The overall therapeutic effect from G3-CA4 alone or in combination with SRS/TMZ will be evaluated by image-guided MRI monitoring of long-term survival rats.

Principal Investigator: Meser Ali, Ph.D.
Extracellular pH Mapping as Therapeutic Readout of Nanoparticle-based Drug Delivery in Glioblastoma (R01EB023366) Subcontract

Extracellular acidosis (i.e., low pH) is a tumor microenvironment hallmark, caused by atypical metabolism and perfusion. Acidic pH enhances cancer growth, proliferation, and builds therapy resistance. The prognosis remains dismal for most brain tumor patients. Malignant gliomas, including glioblastoma multiforme (GBM), fail treatments because gliomas invade outside tumor boundaries conventionally demarked by MRI contrast and the blood-brain barrier (BBB) blocks most drugs. Furthermore, conventional MRI methods are insensitive to physicochemical parameters like pH and mainly track intratumoral volume. Among the primary MRI methods are paramagnetic agents for longitudinal (T1) contrast, where assessment of treatment response involves 2D or 3D measurement with Gd3+ enhanced MRI contrast. However, such methods are not reliable in distinguishing pseudoprogression and pseudoresponse from actual changes in tumor status. Thus there is an urgent need for alternative MR techniques sensitive to metabolic changes, which can aid in effective monitoring of therapeutic response in addition to measuring the tumor size. Because acidic pH milieu is conducive to tumor growth and builds resistance to therapies, simultaneous mapping of pH inside and outside the tumor (i.e., intratumoral-peritumoral pH gradient) is an important cancer imaging need. A novel way to map intratumoral-peritumoral pH gradient is using lanthanide III (Ln3+) agents with BIRDS methods, where physicochemical factors like pH contribute to shifts of non-exchangeable protons. To meet the need for MR readouts of the tumor physicochemical state, we
developed BIRDS to map the intratumoral-peritumoral pHe gradient and found that it is a sensitive readout of cancer growth and treatment. Based on preliminary data obtained from GBM models (e.g., U251), including patient-derived xenograft (PDX) models, we will validate high-resolution pHe mapping with BIRDS as therapeutic readout of chemotherapy drugs delivered into human GBM models. Although we detected 1-2 mm diameter tumors with BIRDS using non-methylated agents, higher resolution mapping of intratumoral-peritumoral pHe gradients will be reached with a novel methylated multiplexed agent in Aim 1.

In Aim 2 we will validate intratumoral-peritumoral pHe gradient mapping by BIRDS with fluorescent pHe probes. We will use BIRDS to examine how intratumoral-peritumoral pHe gradients change with tumor aggression (Aim 3). We will also test compatibility of pHe mapping with BIRDS for tracking response to chemotherapy drugs (e.g., Temozolomide and Sorafenib) being used to treat GBMs (Aim 4). Both of these drugs are known to cross the BBB and are used in GMB therapy. Temozolomide activates apoptosis by alkylating DNA to stall cell replication and Sorafenib is a multiple kinase inhibitor targeting several oncogenic pathways and enhances glycolysis. If successful, pHe mapping by BIRDS will enable monitoring of therapeutic response of various chemotherapy drugs for preclinical PDX models to potentially be translated clinically.

Principal Investigator: Ana deCarvalho, Ph.D.
Targeting Oncogene Amplification in Glioblastoma (W81XWH-19-1-0693)

Malignant gliomas originate in astrocytes or neural progenitor cells within the brain. These tumors rarely metastasize outside of the brain and exert their devastating health effects by damaging this vital organ. Glioblastoma is the most aggressive type of glioma, and unfortunately the most frequent in adults, with an incidence of about 12,000 new cases per year in the US. Surgery followed by radiation and chemotherapy with a DNA targeting agent temozolomide are able to control the disease temporarily. We have learned from the many failed clinical trials that these tumors have been unexpectedly resistant to therapy directed at what seemed rational targets. In over 70% of the cases, proteins that drive glioblastoma growth are overexpressed due to high level gain in copy numbers in regions of the genome that become amplified. In this project we are focusing on two oncogenes, CDK4 (cyclin-dependent kinase 4) and MDM2 (mouse double minute 2 homolog) that overrule the two most important pathways restricting cell proliferation. Novel potent inhibitors targeting these oncogenes are now in clinical trials enrolling glioblastoma patients. This project involves bringing the clinical challenges to the research laboratory in a tangible way, by testing these novel therapies in patient-derived models (neurosphere cells and mouse xenografts) that are as hard to treat as the original tumors they came from, because we have shown they indeed preserve the genomic complexity and oncogene amplifications of glioblastomas. Similar to clinical studies, this project uses genomic information to assign glioblastoma patient-derived models to drugs that in theory should be effective. However, unlike the clinical setting, the same “patient” will be assigned to multi-arm treatments simultaneously. In this project we will investigate the pharmacological properties, specificity and efficacy of the most promising pharmacological inhibitors of MDM2 and CDK4 in treating glioblastoma models with a diverse set of genomic abnormalities. Not only MDM2 and CDK4 genes are amplified, but they are amplified in a circular segment of DNA, that like the chromosomes is packed in chromatin, but does not have centromeres, so they can increase dramatically in number. Furthermore, the tumor cells have different amounts of these circular DNA elements, thus variable levels of the drug target among cells within the same tumors. This likely poses a challenge for these new drug treatments. For that reason, we are adding another therapeutic strategy to our study, we will target an enzyme that is a central regulator of DNA repair, named DNA-PK, also using novel more selective and potent drugs in currently in clinical trials. This has a broader application for glioblastoma treatment, because radiation and temozolomide inflict DNA damage, and so does the rapid cell proliferation rate, and if DNA repair is inhibited the damage to the DNA becomes toxic and eventually lethal to the cell. However, here we will also test if this strategy affects the propagation of the circular DNA elements carrying oncogenes, as a novel application for DNA-PK inhibitors. First, we will test each of the CDK4 and MDM2 inhibitors separately in the glioblastoma cells and xenografts obtained from various patients. We will quantify their efficacy to selectively inhibit their targets, and to what extent they can control glioblastoma growth, and what are the effects of combining them with radiation and temozolomide. If resistance to therapy is observed, we will analyze these tumors in novel and important ways. We will quantify the shifts in the number of the circular DNA elements that carry the oncogenes and also other molecular changes that will be analyzed by Dr. Poisson, a co-investigator in this project and bioinformatics expert. We will then test the efficacy of the DNA damage inhibitor. It is not feasible to conduct these analyses to identify possible mechanisms of resistance to therapy in the clinic. Our results will greatly help the optimization of these therapies. This proposed research happens in the context of a truly multi-
disciplinary team, with the relevance for patient treatment provided by Dr. Snyder, a neuro-oncologist and collaborator in this project, as with any pre-clinical study, the final validation for our findings will take place in the clinic.

Glioblastoma affects men and women of all ages. Military service members, veterans, and their family members are among those who have suffered from the significant symptoms associated with this tumor. Next, they have to deal with craniotomy, radiation therapy and chemotherapy. In most cases, what follows are just months of decline before patients succumbs to this devastating disease. It is becoming common for the tumors that were removed to undergo genomic profile, and the presence of “targetable genomic abnormalities” is used in molecular tumor boards to recommend experimental treatments for patients that have progressed after the standard of care, even when evidence for clinical benefit is not present, due to the lack of options. Our work takes place in the interface between the research laboratory and clinic, and by tackling the complexity of drug resistance in the laboratory we can contribute in a meaningful way to the long road of bringing glioblastoma into the category of treatable diseases.

**Principal Investigator: Houtan Noushmehr, Ph.D.**

**Epigenomic Master Regulators that Define IDH1/2 Mutant Glioma Tumor Progression**

(W81XWH1810540) Department of Defense

Diffuse gliomas are a heterogeneous group of primary brain tumors that develop from glial cells, such as glioblastoma, astrocytoma, and oligodendroglioma. Although environmental risk factors for glioma and glioblastoma remain poorly defined, with the exception of exposure to ionizing radiation, evidence has shown that traumatic brain injury may predispose service members to gliomagenesis via inflammation and stem cell transformation. Despite advances in surgical techniques and clinical regimens, treatment of gliomas remains challenging and the tumors usually progress or recur. Therefore, understanding the mechanisms involved in glioma progression and recurrence is essential and will have clinical implications. The hypothesis tested here is based on our recent reports that defined and characterized glioma subgroups as related to specific genetic alterations associated with \( IDH1/2 \) mutations and chromosome 1p and 19q deletions (codel) and distinct epigenetic alterations. Within the \( IDH - \)mutation subgroup, two novel subtypes were defined by intact codel status and DNA methylation levels, namely Glioma-CpG Island Methylator Phenotype (G-CIMP)-low and -high. G-CIMP-low patients were younger at diagnosis (<39 yo) and exhibited a shorter overall survival compared to G-CIMP-high. Longitudinal analyses of matched glioma samples revealed that 12% of G-CIMP-high cases progressed to G-CIMP-low and the molecular changes occur at candidate non-coding functional elements, suggesting the existence of a potential master regulator affecting brain tumor progression in young adults.

**Objective/Hypothesis:** The hypothesis that G-CIMP-high tumors may relapse as G-CIMP-low gliomas with no NA methylation and other epigenomic events could be a key determinant of the mechanisms that drive glioma progression. Using advances in next generation sequencing and insights of the relationship between transcription factor (TF), histone modifications and DNA methylation, the goal of this project is to investigate the functional genomic elements (e.g. enhancers or silencers) that define brain cancer progression between G-CIMP-high and G-CIMP-low.

**Specific Aims:** In aim 1, we will screen our nationally recognized brain tumor bank for G-CIMP-high to G-CIMP-low transformation using our published signatures. We will obtain matched recurrent pairs, isolate the molecular DNA/RNA and chromatin and generate high-throughput next generation sequencing of distinct epigenomic profiles such as whole-genome bisulfite sequencing, chromatin immunoprecipitation of H3K27ac, CTCF followed by sequencing and RNA-sequencing. In aim 2, we will define candidate master regulators in non-coding functional genomic elements by integrative bioinformatics approach and integrate the most significant findings with publicly available data from ENCODE/Roadmap to define candidate functional genomic elements that will discriminate subgroups during tumor progression. In aim 3, we will validate the identified genomic functional elements in appropriate \( IDH - \)mutant cell culture using targeted CRISPR/Cas-9 to delete the genomic elements, and shRNA to do s of the targeted alterations on gene expression will be assessed by RNA-seq.
Principal Investigator: Ye Xiong, M.D., Ph.D.
Exosome-based Therapeutics in TBI (R01NS100710)

Traumatic brain injury (TBI) is a major cause of death and disability worldwide. There are no effective therapies available for TBI patients. Thus, there is a compelling need to develop novel therapeutics in order to improve neurological recovery after TBI. Mesenchymal stem cells (MSCs) are adult multipotent cells that give rise to various mesodermal cell types. The use of MSCs for tissue repair is of great interest because of their ability to home to damaged and inflammatory tissues. However, previous studies from us and others show that only a small proportion of transplanted MSCs actually survive and few MSCs differentiate into neural cells in injured brain tissues. The predominant mechanisms by which MSCs participate in brain remodeling and functional recovery are related to their secretion-based paracrine effect rather than a cell replacement effect. Our recent data suggest that posttraumatic treatment with cell-free exosomes isolated from rat and human MSCs improves functional recovery in male rats after TBI. Exosomes play an important role in intercellular communication. Exosomes transfer not only proteins and lipids but also genetic materials including mRNAs and microRNAs (miRNAs) to recipient cells, thereby mediating a variety of biological responses. Our preliminary data further demonstrate that the labeled exosomes administered intravenously after TBI reach the brain and are incorporated into brain cells as well as in macrophages in peripheral organs. Our encouraging findings indicate that MSC-derived exosomes have equivalent restorative effects as their cellular counterparts on brain remodeling and functional recovery after TBI. Thus, MSC-generated exosomes are novel candidates as a cell- free therapy that can overcome the obstacles and risks associated with the use of naive or engineered stem cells or MSCs. While our results are promising, the precise therapeutic mechanisms underlying exosome therapy for TBI recovery warrant further elucidation. In this proposal, we will first determine therapeutic efficacy of naïve MSC-exosomes for improvement in functional recovery in male and female rats after TBI. We will then evaluate the effect of MSC-exosomes on brain neuroplasticity, and growth factor expression as well as on the brain and peripheral immune response, effects that likely underlie and contribute to functional recovery (Aim 1). We will then evaluate the role of the miRNA content of the MSC-derived exosomes on brain angiogenesis, neurogenesis, synaptogenesis, cell death, growth factors and immune responses underlying functional recovery (Aim 2). Finally, we propose to enhance the therapeutic effects of exosome treatment of TBI by generating and employing tailored MSC-derived exosomes enriched with the miR-17-92 cluster as a treatment for TBI. In addition, we will investigate the molecular mechanisms underlying cellular exosome uptake (Aim 3). This proposal is innovative, and highly translational. This study will provide novel insights into mechanisms underlying the MSC-derived exosome-promotion of functional recovery after TBI, develop a means to amplify the therapeutic effects of exosome therapy for TBI, and form the foundation for clinical translation of exosome therapy for TBI.

Principal Investigator: Yanlu Zhang, M.S, M.D.
Treatment of Traumatic Brain Injury with Velpoloxamer (R01NS109477)

Traumatic brain injury (TBI) is a major cause of death and disability worldwide. There are no effective therapies available for TBI patients. Thus, there is a compelling need to develop novel therapeutics in order to improve neurological recovery after TBI. Among many secondary injury events that occur after TBI, cerebral microthrombosis is an under-recognized, yet important contributor to the secondary brain ischemia and damage that occurs after TBI, and would therefore seem to be one of the central secondary events after brain trauma to bear in mind when designing treatment strategies. Cerebral microthrombi not only lead to ischemia and cell death but also prevent therapeutic drugs from entering into the affected brain and therefore constrain the efficacy of therapeutic drugs, which may be one of important factors ignored during preclinical and clinical trials. Our recent study indicates that early (2 hours post injury) intravenous administration of Velpoloxamer promotes sensorimotor function and cognitive functional recovery after TBI induced by controlled cortical impact (CCI-TBI), which is associated with its robust effect on reducing cerebral microthrombosis formation and neuroinflammation. Velpoloxamer is a purified form of Poloxamer 188 where impurities associated with renal dysfunction have been removed, which is an amphiphilic polyethylene-polypropylene-polyethylene tri-block copolymer that is reported to seal membranes and restore plasma membrane integrity in damaged cells. However, to date, there is a paucity of information about Velpoloxamer for treatment of TBI and the mechanisms underlying its therapeutic effects. von Willebrand factor (vWF) released into blood from injured endothelial cells inversely correlates with clinical outcome of severe TBI. vWF can induce microthrombosis formation. Our previous study demonstrated that the level of vWF released into plasma increases at 1-4 hours, peaks at 1-3 days, declines at 8 days, and returns to
normal at 15 days in rats after CCI-TBI. We hypothesize that TBI induces the blood-brain barrier (BBB) damage and release of endothelial-derived vWF, which leads to platelet aggregate and subsequent cerebral microthrombosis-induced secondary injury. In Aim 1, we will first conduct a dose-finding study to identify Vepoloxamer dose and therapeutic window effect on functional recovery without toxicity in young rats (male and female) with TBI. In Aim 2, we will then investigate the mechanisms by which IV administration of Vepoloxamer enhances cerebral microvascular perfusion and promotes functional recovery after TBI. Microvascular integrity, cerebral blood flow, and BBB leakage will be measured dynamically using either laser scanning confocal microscopy or magnetic resonance imaging (MRI). This work will address a previously understudied important issue and is highly translational. Successful completion of this proposed research will elucidate mechanisms underlying IV Vepoloxamer-mediated promotion of TBI recovery and facilitate development of Vepoloxamer as a novel therapeutic approach targeting endothelial cells/microthrombi to improve neurological outcome for TBI patients.

Orthopaedics/Bone & Joint

Principal Investigator: Michael Bey, Ph.D.
Shoulder Function After Rotator Cuff Repair (R01AR051912)

Rotator cuff tears affect about 40% of the population over age 60 and are a common cause of pain and disability. Approximately 250,000 rotator cuff repairs are performed in the United States each year, but healing following surgery is a significant challenge (e.g., 20-70% of surgical repairs fail) and postoperative shoulder function is unpredictable. There is also often a disconnect between repair tissue healing and shoulder function where patients have poor shoulder function (e.g., limited strength and pain) despite an intact repair or, conversely, excellent shoulder function despite a failed repair. Conventional clinical data (e.g., patient age, tear size) are not strong predictors of clinical outcome, and Therefore, this disconnect between healing and function remains difficult to explain. Recent research suggests that repair tension and repair tissue elongation may provide insight into post-operative healing and shoulder function that is not adequately provided by clinical data. However, the relationships between repair tension, repair tissue deformation, healing, and shoulder function are not well understood. The objectives of this application are to determine how rotator cuff repair affects shoulder motion, strength, and patient-reported outcomes, and to assess the influence of repair tension and repair tissue deformation on these outcomes. The rationale for this project is based on several important findings from our on-going work regarding the progression and treatment of rotator cuff tears: 1) rotator cuff pathology, even in the absence of symptoms, has a significant impact on shoulder function, 2) physical therapy improves clinical outcomes despite only minor changes in joint motion, 3) surgical repair appears to alter glenohumeral joint (GHJ) motion in a way that suggests excessive repair tension, and 4) shoulder motion, strength, and patient-reported pain/function scores are interrelated after surgery. Based on these findings and the purported roles of repair tension and repair tissue elongation, our central hypothesis is that repair tissue elongation (up to and including failure) is due, at least in part, to repair tension approaching or exceeding the mechanical capacity of the healing repair tissue. We also hypothesize that repair tissue deformation affects joint motion in ways that have a significant impact on strength and patient-reported outcomes. Our approach will be to conduct a longitudinal study that measures repair tension, repair tissue deformation, joint motion, strength, and patient-reported outcomes before and after surgical repair. The proposed research is innovative because it will use a state-of-the-art imaging technique to provide an accurate assessment of the mechanical progression of healing rotator cuff repair tissues. The contribution of this research will be significant because it will advance our understanding of how surgical repair influences shoulder function and clinical outcomes, ultimately leading to improved patient care.

Principal Investigator: Michael Bey, Ph.D.
Shear Wave Elastography to Predict Repair Tissue Healing and Shoulder Function After Rotator Cuff Repair (R21AR072785)

Rotator cuff tears are common, affecting up to 40% of individuals over age 60 and accounting for an economic burden of $3-5 billion per year. Surgical repair is a satisfactory solution for many patients, but clinical outcomes and healing of the repair tissue after rotator cuff surgery can be unpredictable. Tear chronicity (i.e., the extent to which the muscle/tendon unit has degenerated over time) is a critical
factor in determining healing and clinical outcomes. However, conventional approaches for assessing
tear chronicity uses only qualitative descriptions (mild, moderate, severe) or grades (0 to 4) without any
explicit assessment of the quality of the muscle and tendon tissues. Consequently, it is perhaps not
entirely surprising that these conventional assessments are only weak predictors of healing and clinical
outcome after surgery. This is important, because without a reliable measure of tear chronicity it is
difficult for surgeons to know prior to surgery how challenging the repair may be what alternatives may
need to be considered during surgery, what post-operative rehabilitation activities should be prescribed,
and how best to counsel patients on expected outcomes. Ultrasound shear wave elastography has
emerged as a promising technique for non-invasively assessing the in-vivo stiffness of soft tissues.
Given that the pathologic processes associated with rotator cuff tears are characterized by changes in
tissue stiffness, shear wave elastography may have clinical utility in assessing the chronicity of rotator
cuff disease. However, even though this advanced technique has been used extensively for breast and
liver imaging, it has seen only limited use in musculoskeletal tissues. Consequently, the objective of
this study is to determine the extent to which rotator cuff shear wave speed (SWS) predicts healing and
clinical outcomes after rotator cuff repair. Our approach will be to use shear wave elastography to
measure SWS in patients who are having surgical rotator cuff repair. These data will be acquired prior
to surgery and then related to conventional tear characteristics (tear size, tear retraction, muscle
atrophy, fatty degeneration), healing, and conventional clinical outcomes (strength, ROM, patient-reported
outcomes) collected at 12 months post-surgery. Our central hypothesis is that SWS will be a significant
predictor of healing and clinical outcomes and superior to conventional predictors of healing and clinical
outcome. The proposed research is innovative because it will use an emerging technology to assess the
quality of the rotator cuff tissues, which cannot currently be obtained in any other way. This contribution of
the proposed research will be significant because we believe it will establish the clinical utility of shear wave
elastography by identifying SWS as a superior predictor of clinical outcome and repair tissue healing. In
turn, clinical use of shear wave elastography will provide physicians with the information necessary to
improve care for patients suffering with rotator cuff tears

Principal Investigator: Jamie Fitzgerald, Ph.D.
The Role of SHIP2 in Mineralization (R21AR072297)

Skeletal mineralization is fundamentally important to all vertebrate species. Too little mineralization results
in structurally compromised bone that is prone to failure. On the other hand, pain and disability occur when
there is inappropriate or ectopic mineralization and calcification of soft tissues. The major mechanisms
controlling mineralization are poorly understood resulting in a major gap in knowledge. Our ongoing studies
on the genetic basis of opsismodysplasia (OPS), a rare chondrodysplasia that is characterized by a marked
delay in endochondral ossification, identified a new potential regulator of matrix mineralization: SH2
Domain-containing Inositol 5-phosphatase 2 (SHIP2). SHIP2 functions as a phosphatase that
deposphorylates phosphatidylinositol (3,4,5) P3 (PIP3) to generate hosphatidylinositol (3,4) P2 (PIP2).
Data from our in vitro SHIP2 inhibitor and SHIP2-deletion studies confirmed that SHIP2 deficiency leads to
a mineralization defect. Furthermore, experiments on matrix vesicles (MVs) isolated from chondrocytes and
osteoblasts demonstrated that the loss of SHIP2 leads to a failure of MVs to support mineral deposition.
Together, our data support the overall hypothesis that SHIP2 regulates MV function.

SPECIFIC AIM 1: To investigate the regulation of cell surface phosphoinositides (PIs) by SHIP2 in
mineralizing cells. Hypothesis 1 states that SHIP2 controls MV formation by regulating PI cell surface
levels. We have generated new tools to address this hypothesis including ATDC5 chondrocyte and SaOs-2
osteoblast cell lines engineered using CRISPR to eliminate SHIP2 protein, and several highly specific,
fluorescently labeled PI-binding proteins. These reagents will be used to track the sub-cellular and cell
surface PIP3, and metabolites of PIP3 in the presence or absence of SHIP2.

SPECIFIC AIM 2: Investigate the control of MV composition by SHIP2. Hypothesis 2 states that SHIP2
regulates mineralization by controlling MV composition. This hypothesis will be addressed in two Sub-Aims:

Sub-Aim 2.1: Define the proteins recruited to the cell surface by PIs: PI pull-down experiments
will be conducted separately on cell lysate, MV, and membrane preparations using reagents against
biosensor proteins. We will focus on PIP3 initially and then other PIPs that have disrupted distribution in the
absence of SHIP2. Immunoprecipitated proteins will be identified by mass spectroscopy and confirmed by
immunoblotting.
Sub-Aim 2.2: Define how SHIP2 controls MV composition: A comprehensive proteomic analyses will be conducted on MV fractions from both skeletal cell types. Mass spectroscopic findings will be confirmed in immunohistochemical and immunoblot experiments.

The proposed studies have the potential for new insight into the mechanisms of the fundamental, but poorly understood, process of matrix mineralization. Significantly, these studies may identify new, rational targets for the clinical treatment of bone mineralization defects and other mineralization-associated disorders.

Principal Investigator: Yener Yeni, M.D.
Clinical Assessment of Vertebral Bone Quality Direct Biomechanical and Textural Analysis via Digital Tomosynthesis (W81XWH1910373)

This project relates to the Topic Area “Musculoskeletal Disorders”, and specifically to the encouragement area of research on measures to improve diagnosis, prediction and optimization of health outcomes. This is because the proposed project ultimately aims to improve the accuracy of assessment for spinal bone fragility and fracture risk. The bones of the spine (vertebrae) are the most frequently fractured ones due to osteoporosis. These fractures are economically costly and burden the patients with many downstream problems including back pain. Military personnel are known to be at greater risk for these fractures and complications. An accurate assessment of vertebral fracture risk is essential for appropriate and timely intervention for the prevention of fracture. This research also relates to the Topic Area “Diabetes”, because a diabetic cohort will be included in the study. Current standard techniques for fracture risk assessment rely on radiographic bone density scans. Additional information regarding the patient’s demographic status and medical history is also incorporated in tools predicting fracture risk. However, these techniques are not very sensitive in identifying who will have a fracture and who will not. This is not too surprising, considering the fact that the information used in the assessment is a crude indirect measure of bone strength not based on biomechanics. To address this concern, we developed a new method, in which two images of a patient’s vertebra are taken in the presence and absence of the patient’s body weight by having them stand and lay down for both images respectively. The images are obtained using digital tomosynthesis (DTS), a system that is similar to computed tomography (CT). The advantages of DTS over CT are that DTS allows for standing and lying images to be captured, offers high resolution and exposes patients to less radiation than CT. The two sets of images are compared using an advanced computational method and deformations in the vertebra caused by standing are measured. From the displacement measurements, vertebral stiffness and overall displacement are calculated as metrics of strength and factor of safety (factor safety is a measure of how strong the bone is relative to the loads it normally experiences). Information on bone microstructure, additional to bone density, is known to increase accuracy in predicting fractures. We can also derive properties related to bone microstructure from DTS images without the biomechanical test. These properties are determined by quantifying the texture in the bone image and called textural properties. We developed these methods in the laboratory in detail using cadaveric vertebrae and laboratory-standard imaging and strength testing. We also performed pilot human studies to establish feasibility of the methods in the clinic. What remains to be determined is how successful the methods will be in the clinical environment for identifying individuals who are at risk. Therefore, this study will be a clinical validation of the new biomechanical and textural DTS methods. In order to determine the ability of DTS methods to correctly identify at-risk patients, the approach will be to compare patients who have conditions or diseases that are known to increase their risk of fracture to normal patients. Therefore, a group of osteoporotic patients with an existing vertebral deformity, a group with primary hyperparathyroidism (pHPT) and a third group with diabetes will be compared to normal patients. These diseases are considered service-related and thus represent a greater risk for military families. Importantly, each of these diseases increase the risk of fracture but alter bone in different ways that are not always detectable by bone density scans. For example, osteoporosis primarily results in loss of bone mass, pHPT alters the organization of bone structure and affects the cortical bone (in the case of a vertebra, the dense bony shell surrounding the vertebra), and diabetes alters the quality of the bone material without reducing bone mass. By studying these groups using DTS and comparing the assessment of bone strength to standard bone density results, we will: i) establish for which types of patients and to what extent the DTS methods might be useful, ii) identify in which way the method must be performed for best results, iii) better understand how differences in bone quality specific to each disease affect the biomechanical outcome, and iv) establish a group of patients that we can follow up for longer term results. In the long term, the method will be useful in other clinically significant issues such as low back pain associated with vertebral fractures, implant stability, degenerative and congenital diseases of the skeletal system resulting in deformities, skeletal response to drug, exercise and disuse.
Otolaryngology

Prinicipal Investigator: Lamont Jones, M.D., M.B.A.
Characterization of Keloid Specific Exosomes and Determination of Exosomal Critical Signaling Pathways in the Keloid Microenvironment (1K08GM128156)

There are more than 11 million people in the world with keloids and more than 425,000 associated clinic visits, yearly, in the United States. Keloids are benign fibroproliferative tumors which cause pain, pruritus, emotional distress and loss of function. Current therapies are unsatisfactory with unacceptably high recurrence rates, mainly because of an incomplete understanding of keloid pathogenesis. Fibroblasts are a key player in keloid pathogenesis, but the drivers are unknown. Keloid disease is influenced by aberrant signaling pathways. However, no clear signaling pathway has been identified. Exosomes mediate cell-cell communication, exercising primary physiological and pathophysiological function. Exosomal cargo, such as microRNAs (miRNAs), regulate cellular function.

Our group identified RAB27, important for exosome secretion, as being differentially hypomethylated in keloid compared to normal skin. Our group has isolated keloid-specific exosomes. To date, there are no published studies on keloid exosomes or the contribution of RAB27 methylation on exosome function. We propose to test the central hypothesis that exosomes communicate critical signaling events in the keloid microenvironment mediated by RAB27 gene methylation. Aim 1: To determine the effect of RAB27 methylation on keloid exosome production and miRNA cargo profiles. Hypotheses: (1) Keloid exosomal production correlates with RAB27 gene methylation. (2) Keloid exosomal cargo miRNA expression profiles correlate with RAB27 gene methylation. (3) Keloid exosome miRNA’s putative target genes lie within pathways essential for wound healing and/or fibrosis. Aim 2: To determine the effects of keloid exosomes on the keloid microenvironment. Hypothesis: Keloid fibroblast exosomes compared to normal fibroblast exosomes will cause pro-fibrotic phenotype changes in normal fibroblasts. Aim 3: To determine the effect of keloid exosomes on scar formation in vivo. Hypothesis: Exosomes generated in aim 1 and tested in aim 2 will increase scar formation in a rabbit ear scar model.

Significance: This project will lead to an enhanced understanding of keloid pathogenesis and the potential for exosome-based therapy. Innovation: (i) rational progression from preliminary data supporting the novel role of exosomes in keloid pathogenesis; (ii) investigating the influence of RAB27 methylation on the function and production of keloid exosomes would suggest a mechanistic basis for novel epigenetic biomarkers; (iii) using unique resources which includes fibroblast cell lines from primary untreated keloid (25) and matched normal skin (25) from a multi-ethnic group of patients and an in vivo animal model allow for the pragmatic translational application of results; (iv) entirely new field of keloid investigation. In summary, this project, mentoring and career development plan will position, Lamont R Jones, MD, MBA, to become an independent clinician scientist and leader in keloid pathogenesis.

Pediatrics

Principal Investigator: Maureen Connolly, M.D.
Using Evidence-Informed Interventions to Improve Health Outcomes among People Living with HIV: Transgender Women Engagement and Entry to Care Project (TWEET) (U69HA310670100)

There is a pronounced need for implementation of evidence-informed interventions to reduce HIV-related health disparities and improve health outcomes, including improving retention in care, treatment adherence, and viral suppression for people living with HIV (PLWH). In 2016, 81.7% of PLWH in the U.S. were retained in care, and approximately 85% were virally suppressed. The need for these efforts is felt most deeply among racial/ethnic minority men who have sex with men (MSM) and among transgender women. Retention in care for young Black MSM (YBMSM) was lower (75%) than the national RWHP average. 79% of transgender women has achieved viral suppression. Transgender Black/African American had lower percentages of viral suppression across demographic subgroups compared to transgender Hispanic/Latinos and whites. PLWH often have complex behavioral health comorbidities that complicate their ability to maintain treatment adherence and continuous care. A 2010 survey of 246 Ryan White Part C medical providers found that 30% of PLWH had a substance use disorder and 35% had a serious mental illness. Other studies have
found between 35-64% of PLWH suffer from PTSD. 3,4 Although research has defined best practices for addressing steps along the HIV care continuum, the implementation of such interventions lags behind. This is especially true for the implementation of interventions that: 1) are tailored for Black MSM and transgender women; 2) address the co-occurring behavioral health needs of PLWH; 3) tackle the social, structural, and environmental barriers— including experiences of trauma—that hinder attainment of positive health outcomes. This initiative will focus on supporting the implementation of interventions to improve HIV-related health outcomes in the above focus areas. The implementation of the interventions will be evaluated using an implementation science approach. The evaluation will systematically collect and analyze project data in order to measure and monitor progress towards meeting the goals and objectives of the project, while also evaluating the ability of specific interventions to improve the HIV care continuum outcomes of linkage, retention, re-engagement, and viral suppression among client participants. Lessons learned and best practices will be identified throughout the course of the initiative and will be shared rapidly with the larger field.

Radiation Oncology

Principal Investigator: Carri Glide-Hurst, Ph.D.
Development of Anatomical Patient Models to Facilitate MR-only Treatment Planning (1R01CA204189) Subcontract

Accurate delineation of targets and organs at risk for radiation therapy planning (RTP) remains a challenge due to the lack of soft tissue contrast in computed tomography (CT), the standard of care imaging for RTP. Radiation Oncology has addressed this limitation by registering magnetic resonance images (MRI) to CT datasets to take advantage of the superior soft tissue contrast afforded by MRI. MRI brings considerable value to RTP by improving delineation accuracy which, in turn, has enabled dose escalation to improve local control while maintaining or reducing normal tissue toxicities. However, the current integration of MRI as an adjunct to CT has significant drawbacks as it requires image registration and contour transfer between datasets. This process introduces systematic geometric uncertainties that persist throughout treatment and may compromise tumor control. Thus, we propose to translate MR-only RTP into clinical use, with the ultimate goal of improving patient outcomes accomplished via improved treatment plan design. MR-only RTP will eliminate redundant CT scans (reducing dose, patient time, and costs), streamline clinical efficiency, entirely circumvent registration uncertainties, and fully exploit the benefits of MRI for high-precision RTP. Yet, MRI is not routinely used alone for RTP, largely due to its known spatial distortions, lack of electron density, and inability to segment the bone needed for online image guidance and electron density mapping for dose calculation. The central hypothesis is that the innovative technologies that our multi-disciplinary academic/industrial (Henry Ford Health System/Philips Healthcare) collaboration develop will yield geometrically accurate patient models built from MRI data across several platforms/field strengths with CT-equivalent densities that can be used in confidence throughout the entire RTP workflow. In Aim 1, we will perform geometric distortion corrections, determine distortion variability with changing anatomy, benchmark the results in a novel modular phantom, and develop an image processing toolkit. In Aim 2, we will fully automate MR image segmentation in the brain and male/female pelvis to yield accurate synthetic CT patient models derived from novel MRI sequences, including provisions for metal implants, and benchmark the results in phantom. In Aim 3, we will conduct end-to-end testing to characterize the uncertainties in the MR-only RTP workflow. We will perform a virtual clinical trial of MR-only RTP for brain and male/female pelvis and compare to the standard of care. Final translation will include developing physician-physicist practice guidelines, end-user validation of all translational steps, and dissemination of image processing tools into the Radiation Oncology community. This research will systematically address the major challenges limiting MR-only RTP and lay the groundwork for multi-institutional clinical trials across MRI platforms. It will support future work related to MR-guided RT, functional MRI for biologically adaptive RT, and focal RT to areas of high tumor burden.
The androgen receptor (AR) plays a critical role at all stages of prostate cancer (PCa) (1-4). Therefore, for over seven decades, treatments that target AR action have been a mainstay for the treatment of advanced PCa. However, these therapies do not provide a lasting remission and the disease usually recurs as castration-resistant prostate cancer (CRPC), which, remarkably, still relies on AR. New drugs that target AR action (e.g., enzalutamide and abiraterone) provide incremental benefit but no cure. We have discovered a new role of AR, and we have found a way to exploit this role and create a more effective way to kill PCa cells. We discovered that AR plays a critical role in maintaining telomere stability, even in CRPC cells (5-7). This is important because telomere stability is essential for genome stability and cell survival (8,9).

Telomeres are the DNA-protein structures that cap the ends of linear chromosomes, which are double stranded DNA with a single-stranded overhang (12), and that protect them from fusing to each other (8,9). When such protection is lost (an event referred to as telomere dysfunction), chromosome ends are recognized as lesions and this results in the activation of a DNA damage response (DDR) at the telomeres (8,13). The DDR includes recruitment of specific proteins to the telomeres and activation of ATM. ATM activation leads to the activation of Chk2, a cell cycle checkpoint protein that activates a checkpoint control. Checkpoint activation causes cells to arrest, allowing for repair so that cells may resume cell cycle progression to mitosis. Unprotected telomere ends may undergo fusion or recombination (processes referred to as ‘repair’), leading to the formation of telomere end to end fusions and telomere sister chromatid exchanges. Such telomere aberrations push cells into breakage fusion-bridge cycles, resulting in unequal distribution of genetic material to daughter cells and the development of genome instability (14,15).

Notably, when cells with telomere dysfunction are treated with an ATM inhibitor, checkpoint activation is abrogated, allowing cells to enter mitosis with damaged telomeres; the more damage, the more likely the cell will activate a cell death pathway (7,16,17). We discovered that a subset of AR in human PCa cells is associated with telomeres, and that AR antagonists, including bicalutamide (Casodex), enzalutamide (MDV3100), or AR-siRNA, induce telomere dysfunction and activate a DDR at the telomeres, in both androgen-sensitive (e.g., LNCaP) and castration resistant (e.g., C42B and 22Rv1) PCa cells (5-7). Most notably, we found that treating CRPC cells with AR antagonist plus ATM inhibitor abrogated checkpoint activation and killed cells that were resistant to growth inhibition by AR antagonist alone (7). These effects on cells in vitro lead us to test the hypothesis that co-treatment with AR antagonist and ATM inhibitor will suppress the growth and recurrence of CRPC in vivo (Aim 1). In addition, we have observed that AR antagonist treatment induces telomere aberrations in LNCaP, C4-2B and 22Rv1 cells (6,7). Therefore, we hypothesize that PCa cells that survive treatment with AR antagonist may develop genome instability, which may promote tumor progression (15,18,19) and help to explain the recurrence of PCa treated with AR antagonist alone (Aim 2). Two interrelated aims (illustrated in Fig.5) will be pursued as follows:

Aim 1: To evaluate the in vivo efficacy of ATM inhibitor to potentiate the anti-tumor activity of AR antagonists: We will test the effect of an ATM inhibitor (KU59403) in combination with an AR antagonist (enzalutamide) on the growth and recurrence of patient derived CRPC tumor xenograft (PDX) models in mice. These studies will validate observations made using cultured cells and establish proof-of-concept that combined treatment with AR antagonist and ATM inhibitor is more effective than either treatment alone in suppressing the growth and recurrence of CRPC.

Aim 2: To determine the effect of AR antagonists on genome stability: We will use a PCR-based approach to analyze telomere fusion (Aim 2-1) and array Comparative Genomic Hybridization (aCGH) to analyze copy number alterations (Aim 2-2) in castration resistant 22Rv1 and C4-2B cells treated with AR-antagonist in vitro, and in recurring PDX tumors from mice treated with enzalutamide alone in Aim 1. A higher level of telomere fusions and copy number alterations in enzalutamide-treated cells and recurrent tumors than in untreated cells and tumors, respectively, may explain the progression of PCa in the face of AR antagonist.

Impact: Combined treatment with AR antagonist plus ATM inhibitor may represent a new way to effectively treat patients with CRPC.
Prostate cancer is a complex disease with multiple tumors originating independently at different stages of growth. Although morphological differences have been well recognized, the underlying molecular complexity in each tumor foci has not been well studied. Tumor growth in each foci can be determined by independent driver molecular aberration(s). Understanding the molecular level of differences in each tumor foci would help to differentiate the patients who may undergo indolent or aggressive disease course. Further, morphological differences mostly help to understand the stage of the disease, but it is not possible to select appropriate targeted therapy. If different tumor foci carry different driver molecular aberrations, targeted therapy for single molecular aberrations may not yield the curative benefit to the patients. Conventionally, systematic sampling of large tumor foci or high Gleason grade tumor foci have been considered for various genetic and molecular studies. In this approach smaller tumor foci with important driver molecular aberration and high metastatic potential can be easily missed. Therefore, using our novel approach, we propose to screen the entire prostate tissue to assess molecular differences in each tumor foci using well characterized prostate cancer specific molecular markers.

Given the distinct ancestral background of the African American genome, there may be fundamental molecular differences that may drive the prostate cancer differently compared with other ethnic groups. Such differences are not studied in detail to understand the racial differences to offer better and more effective treatment options. Recent studies on in-depth molecular characterization of prostate cancer using next generation sequencing approach revealed extensive genetic aberrations in different regions of prostate cancer. From these studies, driver or targetable aberrations within the spectrum of complex molecular aberrations are not identified other than revealing the genomic complexity of the disease. Therefore, in this study, we for the first time attempt to characterize the entire prostate cancer landscape for molecular differences in each tumor foci in African American and compare with Caucasian prostate cancer to understand the racial disparity. We selected a panel of well characterized prostate cancer driver molecular aberrations, including ERG, SPINK1, ETV1, ETV4, ETV5 and PTEN to study the expression pattern on whole mount prostatectomy tissues (entire prostate tissue- not sampling of specific tumor foci). In this approach we will be able to study individual tumor foci, irrespective of the size, to assess the differences in the molecular aberrations and assess tumor molecular heterogeneity and correlate with clinical and biochemical parameters to understand theassociation with disease progression and clinical outcome. Our preliminary analysis of whole mount prostate tissues from 368 patients revealed hitherto unidentified molecular subsets of prostate cancer with more than one driver molecular aberration in different foci in a vast majority of patients. We observed cases with expression of ERG, SPINK1 and ETV1 in three distinct tumor foci in the same patient and associated PTEN deletion. We also observed many cases with two driver aberrations in two independent tumor foci and additional tumor foci negative for the selected markers. Such observations revealed the extent of genomic complexity of prostate cancer and confirms the independent origin of each tumor foci. Until now analyses based on systematic sampling of dominant/index tumor nodule alone eluded the observation of these distinct subsets of patients. We are in the process of analyzing the clinical implications of such observations in each patient by considering the ethnic origin, and other clinical and biochemical parameters. Based on our encouraging preliminary studies, we would like to extend the analysis to at least 500 African American and 500 Caucasian American whole mount prostate tissue to assess the significance of the molecular differences to understand the racial disparity. Since we have already made significant progress in this direction we will be able to complete the analysis of the proposed number of cases within the three years of the study period. The outcome of this study will have greater benefit to the patients by understanding the extent of genomic complexity and also help clinician select appropriate options for better management of the disease. Based on our preliminary screening of 368 cases, we classified patients into the following four major groups a) patients negative for all the selected molecular markers, b) positive for only one marker in all tumor foci (rare), c) positive for two markers in two independent foci or in the same foci with additional foci negative for any of the markers, d) positive for three markers in three independent foci and yet additional foci negative for any of the markers. These observations are not based on random cases. We have identified a significant number of cases in each group (preliminary data- project narrative) and we are in the process of correlating with clinical and biochemical parameters to understand the clinical impact of the results. We will select the African American cases that are negative for all the selected markers for next generation sequencing to discover new molecular markers.
We and others have demonstrated the re-purposing of the anti-diabetic drug metformin in EOC. Retrospective studies also support the beneficial survival effects of metformin in diabetic EOC patients. The mechanism of metformin’s anti-cancer effects has been largely attributed to activation of its target enzyme, AMPK (Adenosine monophosphate-activated protein kinase). AMPK is a highly conserved cellular energy sensor that plays a critical role in the regulation of cell growth by regulating protein synthesis via mTOR; cell cycle by cyclins and p21; insulin and IGF-1 signaling and lipid metabolism. From a previously funded DOD Pilot grant, we recently showed that AMPK is a key player in metformin’s ability to limit high fat-diet and adipocyte mediated promotion of EOC. We also reported that AMPK activation occurred both in the host and the tumor cells, indicating its modulation on both host factors and tumor cells.

To further gain insight into the role of host AMPK in restraining EOC growth, we generated syngeneic ID8 ovarian tumors in AMPK alpha1 knockout (KO). Our preliminary data showed: 1) AMPK KO mice exhibited an accelerated EOC growth and significantly decreased survival compared to wild type (Wt) mice. 2) Absence of AMPK α1 resulted in a dysregulated host immune environment characterized by increased myeloid derived suppressor cells (MDSCs) with amplified immunosuppressive ability. 4) Absence of AMPK altered the energy metabolism of MDSCs by shifting their metabolic state to a highly active state, independent of fatty acid oxidation, which correlated with its increased arginase dependent immunosuppressive function. MDSCs have emerged as critical elements of cancer-induced immune dysfunction by creating immunosuppressive conditions, allowing unchecked tumor growth via their ability to suppress T-cell proliferation and function. EOC has been shown to be associated with infiltrated MDSCs that enhance incidence, metastasis and stemness. Recently, the understanding of energy metabolic pathways used by immune cells to convey their effector functions has seen major advances. AMPK has emerged as one of the central molecules regulating the metabolic shift in the energy pathways of various immune cells vital to their functioning, but its role in regulating MDSC is unknown.
Part III – Population and Health Sciences

- Center for Health Policy and Health Services Research
- Center for Individualized and Genomic Medicine Research
- Department of Public Health Sciences

Center for Health Policy and Health Services Research

Principal Investigator: Brian Ahmedani, Ph.D.
Effects of Medical Products on Suicidal Ideation and Behavior in Serious Mental Illness (HHSF223201810201C) Subcontract

This study will provide a comprehensive program of infrastructure development, methods development, and innovative research to generate real-world evidence addressing critical gaps in the regulation of medical products. We will explore four related issues: Potential for existing medical products to precipitate suicidal ideation and behavior; potential for existing medical products to reduce risk of suicidal behavior; potential for anticipated new medical products to reduce risk of suicidal behavior; and individual variation in liability to both adverse and therapeutic effects.

Principal Investigator: Brian Ahmedani, Ph.D.
An Evaluation of the National Zero Suicide Model Across Learning Healthcare Systems (U01MH114087)
Developing Tools to Evaluate the Impact of Safety Planning and Lethal Means Assessment on Suicide Outcomes (U01MH114087S1) Subcontract

Suicide is a major public health concern – it is the 10th leading cause of death and number one cause of injury related death in the United States (US). Due to national concern about this problem, the National Action Alliance for Suicide Prevention and the US Surgeon General published the joint 2012 National Strategy for Suicide Prevention (NSSP). The NSSP outlines a series of Aspirational Goals (AG) with the specific objective to reduce the national suicide rate by 20%. AG 8 and 9 promote healthcare settings as primary targets for suicide prevention. Consistent with this message, Henry Ford Health System’s (HFHS) Perfect Depression Care (PDC) Zero Suicide Initiative was the first US program linked with a substantial decrease in the suicide rate among behavioral health patients after implementation. These findings have motivated national promotion of this model for suicide prevention in health systems. As such, the National ZS Model (NZSM) was developed, based on the HFHS PDC program, but with flexibility to allow adaptation to diverse settings and patient populations. Overall, the NZSM is founded on the realization that suicidal individuals often fall through multiple cracks in a fragmented and sometimes distracted healthcare system, and on the premise that a systematic, comprehensive approach to care is necessary for suicide prevention. The comprehensive approach of the NZSM includes implementation of a series of clinical and quality strategies within the following components: 1) Identification of those at-risk, 2) Engagement and care management; 3) Effective treatment, and 4) Care transition. Despite being a model program promoted internationally for healthcare system quality improvement in suicide prevention, the NZSM has very limited evidence outside of the findings from the HFHS PDC program. The proposed study seeks to conduct a comprehensive process and outcome evaluation of NZSM implementation in real-world clinical settings across 6 large, diverse Mental Health Research Network Affiliated Learning Healthcare Systems providing healthcare for over 9 million individuals each year. The project aims are to: 1) Collaborate with health system leaders to develop EHR metrics to measure specific quality improvement targets and care processes tailored to local NZSM implementation, 2) Examine the fidelity of the specific NZSM care processes implemented in each system, and 3) Investigate suicide attempt and mortality outcomes within and across NZSM system models. Study data are captured using electronic health records and insurance claims. Given strong national support for NZSM, if it is found to be effective to reduce suicide behavior, this model will have nationwide implications for suicide prevention in healthcare settings.
Suicide is a major public health concern – it is the 10th leading cause of death and number one cause of injury related death in the United States (US). Suicide rates have risen over 25% in the last 15 years. In parallel, the nation is struggling with an opioid epidemic. Opioid prescribing, heroin use, and opioid related overdose deaths have risen substantially. Approximately 15% of all suicide deaths are due to drug overdose, and prescription opioids specifically, are commonly used among people who attempt suicide. Health systems across the country have made decisions to tackle both of these public health crises – implementing policies to dramatically reduce opioid prescribing as well as clinical processes within the Zero Suicide model to improve suicide prevention for their patients. The parent award for this supplement is focused on evaluation of Zero Suicide implementation, including fidelity to each of these clinical processes and suicide outcomes, across 6 large, diverse Mental Health Research Network-affiliated Learning Healthcare Systems providing healthcare for over 9 million individuals each year. Given the overlap, significant reductions in opioid prescribing as part of newly implemented policies should lead to a reduction in the availability of opioids. These reductions may result in a public-health level means reduction approach to reduce suicide. Means reduction is among the interventions recommended within Zero Suicide. The concurrent implementation of these new opioid prescribing policies in the context of implementation of Zero Suicide allows the opportunity to evaluate how changes in opioid prescribing impacts suicide outcomes in health care. This supplement project seeks to accomplish three specific aims: 1) Evaluate changes in opioid prescribing patterns during the period of NZSM implementation across health systems, 2) Investigate whether changes in opioid prescribing patterns reduce suicide attempt and mortality, and 3) Investigate whether changes in opioid prescribing patterns reduce opioid-related suicide attempt and mortality poisonings. Overall, we propose to use an Interrupted Time Series Design, consistent with the parent award, to measure changes in prescribing patterns and suicide outcomes.

Fifty percent of all adolescents will use some form of illicit drugs before the end of high school, 20-25% will meet criterial for depression, and many others will engage in health compromising behaviors like delinquency and violence with consequences for their long-term health. Evidence-based interventions shown to prevent these behavioral health concerns could improve adolescent health trajectories if implemented widely in pediatric primary care. The American Academy of Pediatrics? Bright Futures recommends that pediatricians offer developmentally tailored anticipatory guidance to all parents to support their children’s healthy development, but programs providing guidance are not offered universally. This UG3-UH3 application tests the feasibility and effectiveness of implementing Guiding Good Choices, a universal, evidence-based anticipatory guidance curriculum for parents of early adolescents, in three large, integrated healthcare systems serving socioeconomically diverse families. This intervention reduced adolescent alcohol, tobacco and marijuana use, depression, and general delinquency in two previous rigorous randomized controlled trials. It also strengthened parenting practices and parent-adolescent relationship quality, both broadly protective against behavioral health concerns. Guiding Good Choices has the capacity to achieve population-level impact on adolescent health if made widely available through pediatric primary care. Parents trust pediatricians? advice regarding their children’s well-being, and current research with socioeconomically diverse groups suggests that they are eager to participate in family-focused programs offered in primary care clinics. Building on this body of research, the investigative team, in close cooperation with the NIH Healthcare Systems Research Collaboratory and healthcare systems partners, will conduct a cluster-randomized trial of Guiding Good Choices in 72 pediatric primary care practices. Half will be randomly assigned to offer the program universally to parents of adolescents ages 11 to 12, and half will serve as usual care controls. The study will use a workflow that is easy to adopt, implement, and maintain by primary care clinics to enroll families in the intervention at the adolescent well visit. We anticipate recruiting over 4,500 families into the trial. The team will use the RE-AIM framework to test implementation outcomes and effectiveness, including hypothesized reductions in several behavioral health problems (e.g., substance use initiation, mental health symptoms and diagnoses), and emergency department and inpatient service utilization. We will use data from the EHR and a supplemental behavioral health survey to monitor outcomes up to 3 years post intervention.
We will also assess the feasibility and sustainability of implementing the intervention in each HCS, including health economic evaluation to understand costs in relation to value gained. Throughout the trial the investigative team will engage in ongoing dialog with HCS leaders, pediatricians, and clinic staff to ensure the intervention and implementation process fit the needs of each HCS. We anticipate that evidence of feasibility and effectiveness in three different HCS will foster broad dissemination to achieve public health impact.

Principal Investigator: Jordan Braciszewski, Ph.D.
NIDA CTN-0074: Primary Care Opioid Use Disorders Treatment (PROUD) Trial (3UG1DA040314) Subcontract

Over 20 million US adults and youth suffer from substance use disorders (SUDs) and substance use (SU) related problems. However, most people with SUDs never receive SUD treatment. Historically, research on SUDs has focused on the small minority of patients with SUDs who are seeking, or already engaged in, specialty SUD treatment. The overall goal of the proposed Addictions Research Network (ARN) node of the Clinical Trials Network will be to conduct cutting edge research to improve outcomes in all patients with SU/SUD who are seen in medical settings. The ARN includes 15 large health systems across the US that use the HMO Research Network’s (HMORN’s) Virtual Data Warehouse, providing geographic and racial/ethnic diversity as well as variation in systems of medical and SU/SUD care. The proposed ARN node has 3 broad agendas—1) to evaluate effective practices for identifying, engaging and treating patients with SU or SUDs in medical settings; 2) to develop and test effective, practical ways to implement these practices in a sustained manner as part of routine medical care; and 3) to develop and disseminate innovative research methods on SU and SUDs. Three PIs will lead the ARN node, each with expertise critical to our research agenda. Dr. Weisner, who has more than 25 years of experience leading SUD research in public and private medical settings, will lead the ARN as Senior PI at Kaiser Permanente. She partners at Kaiser Permanente with Dr. Campbell, an expert in research on opioid misuse and patient-centered and comparative effectiveness research, and Dr. Bradley, Senior PI at Group Health, who has 20 years of research experience targeting non-treatment-seeking patients with alcohol misuse and SUDs in medical settings. The ARN will have 3 Cores: 1) an Administrative Core will support all aspects of the ARN node; 2) an Implementation Core will support patient-centered design of practical, sustainable approaches to implementing SUD care in routine medical settings using electronic health records (EHRs), and; 3) an Analytics Core with expertise in programming, biostatistics using EHR data, and economics, will support innovative methods research and study design. ARN work will leverage the HMORN’s 20 years of conducting pragmatic clinical trials and comparative effectiveness research across health systems using EHRs and the nationwide Virtual Data Warehouse. The ARN node will provide a robust foundation for population-based studies—including pragmatic randomized controlled trials, comparative effectiveness studies, and implementation research—that can evaluate long-term health outcomes. Moreover, through our connection to 15 learning healthcare systems, our research will design approaches to improve the quality of care for SU and SUDs in real-world medical settings. In this way—with the other CTN nodes—the ARN node will help build the infrastructure required for the next era of addictions health services research.

Principal Investigator: Amy Loree, Ph.D.
SBI-Tech Michigan: Optimizing SBI Implementation for High Risk Alcohol Use Among Women of Childbearing Age (NU84DD000001) Cooperative Agreement

The purpose of this Notice of Funding Opportunity (NOFO) is to reduce risky alcohol use among women of childbearing age through system-level implementation of alcohol screening and brief intervention (SBI) in health systems providing women’s health services. Risky alcohol use can result in a variety of negative health and social consequences, such as motor vehicle crashes, intimate partner violence, and fetal alcohol spectrum disorders. It is costly, results in over 88,000 deaths annually, and can affect serious medical conditions, such as hypertension, liver disease and certain types of cancer. Health professionals are uniquely positioned to intervene with patients with acute and chronic health conditions caused or exacerbated by risky alcohol use. Alcohol SBI implementation efforts within health systems will focus on development and implementation of: a training and technical assistance plan; alcohol SBI protocols in primary care clinics; system-level approaches that facilitate uptake (e.g., electronic health record integration and performance metrics); an evaluation plan assessing feasibility and impact of system-level implementation; a dissemination plan on promising models and lessons learned; and a sustainability plan. Expected performance outcomes include documenting provider/clinic readiness to conduct alcohol SBI, documenting implementation barriers
and proposed solutions, tracking clinic-level data on alcohol SBI, and assessing the use of system-level strategies.

Center for Individualized and Genomic Medicine Research (CIGMA)

Principal Investigator: Keoki Williams, M.D.

Leveraging Electronic Medical Records to Perform Large-Scale Diabetes Pharmacogenomics among Ancestrally Diverse Patient Populations (R01DK113003)

Diabetes mellitus is a modern-day scourge, affecting an ever-increasing proportion of individuals worldwide, including 26 million Americans currently. Moreover, type-2 diabetes (T2D) disproportionately affects historically disadvantaged U.S. minority groups, as evidenced by the much higher rates of disease and more severe complications among African American individuals. Although there are multiple therapeutic classes of oral medication available for treating T2D, metformin is currently recommended as the first-line therapy. Metformin lowers blood glucose levels by reducing hepatic gluconeogenesis, improving skeletal muscle insulin sensitivity, and limiting intestinal glucose uptake. It has also been shown to be an effective therapy for preventing incident diabetes. Despite being one of the most frequently prescribed drugs worldwide, very little is known about the biologic mechanism(s) through which metformin mediates its effect. This knowledge would be of value therapeutically to better understand and predict treatment response. By extension, even less is known about the activity of metformin among African American individuals, as few studies have included substantial numbers of non-European population groups. This application will help rectify existing knowledge gaps by studying a large and diverse patient population with T2D. Specifically, we will utilize electronic medical record (EMR) data for large-scale diabetes pharmacogenomics. These data have the advantage of being able to account for medication use and drug exposure over time; to provide substantial numbers of individuals for combined and population group specific analyses; and to assess clinical endpoints both retrospectively and prospectively. In this application, we propose the following study aims: 1) To assess whether there are differences in metformin treatment response by self-reported race-ethnicity and genetic ancestry; 2) To use novel, gene-based association approaches to identify both shared and population group specific genetic variants influencing metformin's effect on blood glycemia (i.e., HbA1c levels); and 3) To replicate our findings in a separate group of patients and to include additional exploratory analyses to assess whether the identified genetic variants influence diabetes-related microvascular events, macrovascular events, and adverse drug reactions. The knowledge gained through this study will directly address the goals of Health People 2020 – “achieve health equity, eliminate disparities, and improve the health of all groups.”

Principal Investigator: Keoki Williams, M.D.

Poly-omic Study of Asthma Exacerbations in Diverse Populations (R01HL141845-01A1)

Asthma exacerbations contribute to considerable disease morbidity and account for nearly half of all asthma related costs. Nevertheless, we do not currently have biomarkers that can be used clinically to reliably predict an impending exacerbation. Such measures could transform asthma care if they allowed for the timely recognition, treatment, and prevention of these severe events. It is important to note that certain population groups, such as African Americans and Latinos (particularly Puerto Ricans), suffer disproportionately from these complications with rates of asthma-related emergency department visits, hospitalizations, and deaths nearly 3-5 times higher than those of European Americans. Therefore, it should be surprising that nearly all existing genetic studies of asthma exacerbations have focused on individuals of European descent and have been insufficiently powered to study other groups. Other limitations include analyses which didn’t take into account the timing of events and studies which focused on allergic mechanisms (as opposed to taking an agnostic approach). In this application, we will utilize the enormous amount of whole genome sequence data that will be generated by our Asthma Translational Genomics Collaborative (ATGC) and the NHLBI's Trans-Omic Precision Medicine (TOPMed) program to identify genomic markers of asthma exacerbations. The ATGC comprises 9 cohort studies and 10,840 patients with asthma (7,212 African American individuals and 3,628 Latino individuals). We will use the Study of Asthma Phenotypes and Pharmacogenomic Interactions by Race-Ethnicity (SAPPHIRE) as the discovery set to identify genetic variants associated with time to exacerbation (Aim 1). The SAPPHIRE cohort is ideally suited to serve as the discovery set for this project because of its prospective longitudinal records of events, detailed characterization of participants, and extensive biobank, which includes serum and RNA samples.
Replication of top genetic associations will be performed in the other 7,086 ATGC participants (Aim 2). Transcriptomic data generated from RNA sequencing will be used to identify genes whose expression in whole blood is associated with time to exacerbation (Aim 3a), and variants promoted from Aims 1 & 2 will be assessed as expression quantitative loci for their association with gene expression (Aim 3b). For Aim 4, banked serum will be used to assess the proteome of individuals from phenotype extremes (i.e., serum collected from individuals prior to a severe exacerbation vs. serum from individuals with asthma who don’t experience exacerbations). Using mass spectrometry, we will broadly assess serum for proteins differentially expressed between these groups (i.e., an untargeted proteomic approach), and we will use the information gleaned from the genomic, transcriptomic, and untargeted proteomic analyses to assess specific proteins (i.e., a targeted proteomic approach) for expression differences in additional groups of individuals at phenotype extremes. In short, we are proposing both independent and interdependent “omic” analyses to identify biomarkers of asthma exacerbations in populations at highest risk.

Department of Public Health Sciences

Principal Investigator: Andrea Cassidy-Bushrow, Ph.D.

Delivery Mode, Environment and the Gut Microbiome: Influence on Childhood Body Size (R01HD082147)

Caesarean section (CS) delivery, which accounts for ~32% of all US births, has been associated with offspring obesity. Little is known about the mechanisms linking CS with obesity risk. The gut microbiome, which varies by mode of delivery, is also associated with childhood obesity. In our established racially and socioeconomically diverse birth cohort (WHEALS; Wayne County Health, Environment, Allergy and Asthma Longitudinal Study), the early-life gut microbiome is associated with body mass index (BMI) category at age 2 years; CS is associated with both a distinct early-life gut microbiome and with increased BMI at age 2 years; and the presence of pets in the home, which increases microbial diversity, reduces the association between CS and BMI. Our data provide evidence for a mediating role of the gut microbiome in the CS-obesity relationship. However, to provide stronger evidence requires additional study. This project builds on extant data in WHEALS and on-going data collection in a subset of these children to examine the role of the gut microbiome in the CS-obesity association. Children will be invited for a research clinic visit for comprehensive body size assessment and blood draw at age 10-12 years. Gut microbiome composition and predicted function will be measured in banked early-life (1 and 6 months of infancy) stool samples and in samples from these children at age 10-12 years using the 16S rRNA and ITS2 biomarker genes and the Illumina MiSeq platform. A metabolomics analysis will be conducted in a subset of these stool samples. Adiposity will be measured as BMI at ages 2 and 10-12 years, BMI trajectory from birth to age 10-12 years, and anthropometric, bioimpedance and inflammatory measures at ages 10-12 years. Combined, we anticipate 630 unique children will have 10-year adiposity measures and at least one early-life microbiome measure (~405 with 1 month and ~381 with 6 month stool samples, which includes ~300 children with paired 1 and 6 month samples). Of these children, 400 will also have gut microbiome measured at age 10-12 years. Our specific aims are to: (1) examine if mode of delivery is associated with childhood adiposity; (2) examine if the gut microbiome is associated with childhood adiposity; and (3) examine whether the gut microbiome mediates relationships between mode of delivery and measures of adiposity. Such a complementary “omics” approach has never been applied to the study of childhood obesity and is likely to provide critical insights into disease development in early-life as well as potential targets amenable for intervention.

Principal Investigator: Melissa Davis, Ph.D.
The DARC side of Breast Cancer (R21CA210237)

TNBC is arguably the most deadly BrCa subtype with higher prevalence in pre-menopausal women and in women of African descent. We know that the combined TNBC prevalence and poor treatment options are a likely cause of persistently higher mortality rates in African Americans compared to European Americans in the US. However, we have shown that within African Americans, disparities in BrCa survival are more pronounced within the TNBC category compared to the ER positive groups. These data indicate that unique mechanisms are operating in either tumor biology or host response in women of African descent. The ancient and African- specific Fy- allele alters the regulation of DARC/ACKR1, an atypical chemokine receptor, in a tissue-specific fashion beyond the previously described RBC phenotype. This implicates DARC/ACKR1 in various altered phenotypes in these ancestry groups, specifically as it relates to
chemokine regulation. This project will test the hypothesis that DARC expression in tumor cells alters tissue chemokine levels to modify the host immune response to tumorigenesis, and that absence of DARC expression on blood cells as a result of the African-specific Fy- allele alters circulating chemokine levels, altering the tumor microenvironment and enhancing tumor aggression. Specifically we will; 1- Determine if DARC tumor expression associates with ancestry and altered host immune responses in a pilot BrCa cohort of African Americans and European Americans and 2- Determine if loss of DARC on bone-marrow-derived (bmd) blood cells alters chemokine profiles and tumor immune response, using pre-existing transgenic C3-1Tag BrCa and AckR1-/- mice.

**Principal Investigator:** George Divine, Ph.D.
**Targeted Clinical Trials to Reduce the Risk of Antimicrobial Resistance: Randomized Controlled Trial for Treatment of Extensively Drug-Resistant Gram-Negative Bacilli (Option 5)**

The Gram-negative bacilli organisms Acinetobacter baumannii, Klebsiella spp., Escherichia coli, Enterbactor spp. and Pseudomonas aeruginosa have become a frequent cause of bloodstream infection and pneumonia in the hospital and other healthcare settings. Among these pathogens, antimicrobial resistance has emerged to many classes of antimicrobial agents. Most concerning has been the emergence of resistance to group 2 carbapenems (such as imipenem). In several regions of the world, including Southeastern Michigan, strains of extensively-drug resistant Gram-negative bacilli (XDR-GNB) that exhibit resistance to most, and in some cases all types of available antimicrobial agents, including group 2 carbapenems, have emerged and disseminated. Treatment options for XDR-GNB typically include Colistimethate sodium (referred to as colistin in this study), used alone (monotherapy) or in combination with other agents. Unfortunately, resistance to colistin has begun to emerge in some strains of XDR-GNB, which is a truly concerning development, since colistin is one of the last remaining treatment options for XDR-GNB. No prospective, randomized controlled trials have been conducted to evaluate the clinical efficacy of colistin monotherapy versus colistin-containing combination therapy or the impact of these therapeutic modalities on the emergence of colistin resistance among XDR-GNB. We plan to conduct a double-blind randomized controlled trial including patients with pneumonia and bloodstream infection due to XDR-GNB. After enrollment, subjects will be randomized to receive 14 days of either colistin monotherapy or colistin plus meropenem.

In the Detroit metro area, infections due to XDR-GNB have developed into a regional challenge and common problem. We have assembled a multi-disciplinary team that includes Infectious Diseases researchers, clinicians, infectious diseases pharmacists, microbiologists, epidemiologists and statistical experts to address critically important questions and challenges regarding the management of bloodstream infection and pneumonia due to XDR-GNB. Specifically, we hypothesize that the combination of colistin and imipenem will provide superior efficacy in the treatment of XDR-GNB pneumonia and bloodstream infection and will prevent the emergence of decreased susceptibility to colistin among XDR-GNB strains. We also aim to analyze tools that could be used in "real time" to aid clinicians treating patients with infection due to XDR-GNB. For example, we aim to analyze the association between the presence of in vitro synergy of the colistin and carbapenem (imipenem or meropenem) combination (as determined by E-test) and clinical outcomes; and the association between colistin plasma levels and clinical outcomes and the development of nephrotoxicity.

**Principal Investigators:** Christine Cole-Johnson, Ph.D.
**Human Epidemiology and Response to SARS-COV-2 (HEROS)**
**Core A, Core D, Core E**
**Principal Investigator:** Albert Levin, Ph.D., Core B
**Principal Investigator:** Kimberley Woodcroft, Ph.D., Core C

This request is in response to NOT-AI-20-031 for supplement funding in response to the COVID-19 emergency. COVID-19, the infectious disease caused by SARS-CoV-2, is rapidly affecting humans around the globe. While initial epidemiological data have focused on cases that resulted in severe respiratory disease seen predominantly in adults, little information regarding the infection burden in children is available. This is complicated by the observation that many virologically confirmed cases in children are asymptomatic. Undocumented, and likely infectious, cases could result in exposure to a far greater proportion of the community than would otherwise occur. Indeed, it has been proposed that undocumented
(or silent) infections are the source for almost 80% of documented infections; thus, it is critical to determine the silent and symptomatic infection rate in children. To overcome challenges for clinical study implementation imposed by current healthcare access restrictions, a surveillance study under design will enroll and prospectively observe eligible children, and their family members, that are current participants in our NIH-funded, ongoing, birth cohort studies. These children and their families are known to research staff and as part of their participation in HFHS studies, they have already been exposed to the procedures involved in a surveillance study. We are requesting support for the pediatric studies aligned with our Microbiota and Allergic Asthma Precision Prevention (MAAP2) (PI: Johnson, Owny P01AI089473) to participate in the multi-center survey entitled Human Epidemiology and Response to SARS-CoV-2 (HEROS), Protocol # DAIT-COVID-19-001. Our primary objective is to report the incidence of SARS-CoV-2 infection (detection of virus in nasal secretions) over time in cohort children (index child) and household contacts (caregivers and siblings). A secondary objective is to compare SARS-CoV-2 infection status and antibody development for index children/siblings with atopic conditions (e.g. asthma, eczema) versus children without atopic conditions. As an exploratory aim, we will investigate whether SARS-CoV-2 infection (as determined by virus detected in nasal secretions) is associated with the presence of virus in stool. Our targeted enrollment is 300 families recruited over a 2-week period and followed for a minimum of 6 months. At predetermined intervals, biological samples (nasal swabs, peripheral blood, stool) will be collected by the caregiver at home using materials provided to the family. Symptom and exposure surveys will be completed remotely via a smart phone, on-line, or telephone at the time of biological sample collection. This timely, multi-site study can be rapidly implemented and realistically conducted without necessitating any visits to a clinical research center and will provide invaluable information on the infection burden of SARS-CoV-2 in children.

**Principal Investigators: Christine Cole-Johnson, Ph.D.**
**Microbiota and Allergic Asthma Precision Prevention (MAAP2) Project 1, Project 3, Project 4**
**Principal Investigator: Edward Zoratti, M.D. Project 2**
(P01A1089473)

This application builds on the findings of our initial P01 designed to examine relationships between environmental factors, especially pets, the infant gut microbiota and pediatric allergic asthma. We have shown that: 1) dogs alter the microbial composition of dust in homes, 2) children born into homes with dogs have different developmental patterns of gut microbiota and of IgE, 3) a distinct pattern of gut microbial composition at 1 month of age is related to heightened risk of sensitization to multiple allergens at 2 years and of asthma at 4 years, and this pattern is influenced by numerous maternal characteristics, 4) sensitization to multiple food and inhalant allergens at 2 years is strongly related to asthma at 10 years, 5) the metabolic profiles of stools are related to later allergic sensitization 6) 12,13-DiHOME, a metabolite in stool, promotes development of Th2 lymphocytes and lowers development of Treg lymphocytes in an in vitro assay, and 7) in another study, the meconial microbiota is distinct in neonates born to mothers with asthma. Our complementary mouse studies have shown that: 1) gavaging with dust from homes with dogs reduces lung inflammation from allergen sensitization and from respiratory syncytial virus (RSV) infection, 2) dog dust gavaged mice have increases in Lactobacillus johnsonii in their cecas 3) oral administration of live L. johnsonii confers protection against pulmonary inflammation induced by allergen and RSV, 4) L. johnsonii alters the function of bone marrow-derived dendritic cells, 5) mice orally supplemented with L. johnsonii have altered serum metabolic profiles, and 6) mouse pups born to L. johnsonii-supplemented mothers are protected against allergen challenge and RSV infection. Collectively these findings showing the influence of maternal factors provide the basis for this application's focus on the maternal gut and vaginal microbiotas during pregnancy, and how these relate to infant gut microbial development and risk of allergic asthma. Project 1 focuses on the relationship of maternal environmental and dietary factors, including maternal and infant gut microbiotas, to the child's developing a high-risk for asthma phenotype by age 2 years. Project 2 proposes a detailed examination of relationships between maternal and child microbiota, breast milk composition and IgE development amongst a cohort of pregnancies in which the mother has current allergic asthma. Project 3 synergistically interacts with Projects 1 & 2 and also uses specimens from 10-year-old allergic asthma cases and controls in the initial P01 birth cohort to examine gut microbes producing metabolites associated with a lowered risk of allergic inflammation and how they are transferred from mother and established in offspring. Project 4 will use mouse models to examine the relationships between manipulation of maternal microbiota and immune development in offspring. We anticipate that together these studies will show that interventions directed at the gut microbiota of mothers during pregnancy and of high-risk neonates after birth could reduce the risk of allergic asthma in childhood.
Such findings would provide the foundations of a rational strategy to prevent allergic asthma.

Principal Investigator: Christine Cole-Johnson, Ph.D.
Children’s Respiratory and Environmental Workgroup (CREW) (UG3OD023282) Subcontract, Cooperative Agreement

The grant is part of $157 million in awards announced yesterday by the NIH that launches a seven-year initiative called Environmental Influences on Child Health Outcomes (ECHO). The ECHO program will investigate how exposure to a range of environmental factors in early development – from conception through early childhood – influences the health of children and adolescents.

Individual birth cohort studies have identified risk factors for developing childhood asthma, including environmental exposures in early life such as allergens, pollutants, patterns of infection and colonization with viruses and bacteria, and psychosocial stress. Despite such advances, further progress in understanding the root causes of asthma have been hampered by at least two factors. First, procedures and scientific methods are not standardized across cohorts, making it difficult to compare and validate findings. Second, asthma definitions across cohorts vary considerably. In fact, asthma is a syndrome; there are different subtypes of asthma with distinct clinical features (asthma phenotypes) and likely different etiologies (asthma endotypes). We hypothesize that host factors (genetics, epigenetics) interact with environmental exposures during the prenatal period and early childhood to cause specific endotypes of childhood asthma. We further propose that identification of endotypes and associated molecular biomarkers in early life can provide a new paradigm for asthma prevention. Unfortunately, single cohorts have limited ability to identify asthma endotypes due to small sample size and unique population characteristics. To overcome shortcomings of individual cohorts, investigators leading 12 asthma birth cohorts across the U.S. now propose the establishment of the Children’s Respiratory Research and Environment Workgroup (CREW) consortium. This consortium proposes to identify asthma endotypes and overcome shortcomings of individual cohorts by: 1) providing a large (nearly 9000 births and long-term follow-up of 6000-7000 children and young adults) and diverse national data set, 2) harmonizing data related to asthma clinical indicators and early life environmental exposures, 3) developing standardized measures for prospective data collection across CREW cohorts and other ECHO studies, and 4) conducting targeted enrollment of additional subjects into existing cohorts. This approach will enable collection of samples that are optimized for a systems approach to understanding how environmental and host factors in early life promote the development of specific asthma endotypes. Collectively, the results of this comprehensive research to identify the root causes of asthma vs. resilience and health will go far beyond what can be accomplished by individual cohorts, and thus provide a foundation for future efforts aimed at personalized prevention of chronic childhood asthma.

Principal Investigator: Christine Cole-Johnson, Ph.D., Public Health Sciences and Brian Ahmedani, Ph.D., Center for Health Policy and Health Services
Trans-America Consortium of the Health Care Systems Research Network for the All of Us Research Program (OT2OD026550)

Clinicians throughout history have worked to tailor both prevention and treatment strategies to the individual patient’s needs; it is a fundamental credo to the practice of medicine. However, the vast majority of evidence-based clinical practice is based on research results acquired from measuring the common treatment effect on the “average person” in a restricted patient population with limited data, which we now know does not necessarily apply to numerous patients in the real-world setting. Thus, some patients will benefit from evidence-based treatments and preventative interventions, while others will be harmed by taking medications or undergoing processes and procedures that are at best non-effective and at worst cause serious side effects. However, since the initiation of pharmacogenomics in the mid-90s, the astounding pace of development of the technical and analytic tools to measure individual inherited and acquired biological variations at all physiological levels, as well as efficiently capture a patient’s medical and risk factor history and personal preferences via electronic means, is at a scale never before known.(PMIDs: 26554403, 26804248, 26802434, 26686739, 26769233, 26702339, 26700443, 26764593, 25231862) The current concept of “Personalized Medicine” or “Precision Medicine” in which these tools can be deployed to sharply hone predictions about an individual’s risk for disease or response to treatment, while still in its infancy, has immeasurable potential.(PMIDs:20551152, 26014593, 26810587) Further, the costs for next generation sequencing are expected to continue to decline as technology advances. (PMIDs:
As resources are becoming increasingly constrained, it is important to devote scientific time, energy and dollars to questions that matter to the community and have strong potential for effectively improving medical care, public health and wellness. Hence the need, creation, and continuing development of the All of Us Research Program (AoURP). The promise of Precision Medicine in the U.S. can be most effectively realized on a large scale in the next decades if a research infrastructure is established and accessible to scientists across the nation and includes a large and engaged study population with comprehensive health and lifestyle histories linked to biospecimens. Critically, this population must be diverse, representing minority and other subgroups underrepresented in biomedical research. Further, as our investigators and others have recently published, the need to engage all stakeholders, including patients and providers, into both the research and “integration into practice” aspects of Precision Medicine as it progresses, is widely recognized. Our Consortium objective is to recruit 93,000 participant partners into the AoURP, with a focus on African Americans, Arab Americans, Hispanics, rural residents, persons of low socioeconomic status (SES) and children, with the ability to target other groups of interest as needed. Now that we are rapidly ramping up engagement efforts in preparation for AoURP national launch, we will capitalize on an influx of appropriate resources and our experience in engaging, recruiting and retaining large numbers of participants in epidemiological and clinical cohorts, along with our patient-centered and process improvement approaches, to efficiently maximize recruitment and retention in the AoURP.

Principal Investigator: Lois Lamerato, Ph.D.
US Hospital Vaccine Effectiveness (VE) Network (U01P000974) Subcontract
US Influenza Vaccine Effectiveness Network (U01IP001034) Subcontract

Prevention of hospitalization has long been viewed as a major health benefit of the use of influenza vaccine. This was, in large part, the rationale for the initial vaccination programs targeting the elderly and those with underlying health conditions. However, in the last decade, questions have been raised about the value of such programs. Modern study designs to assess vaccine effectiveness (VE) have required laboratory confirmation of influenza infection, as well as documentation of vaccine receipt and the use of a test-negative design to control for differences in healthcare-seeking behavior between vaccinated and unvaccinated patients. There is a need for current estimates of VE in preventing influenza-associated hospitalization using these methods. We propose estimation of influenza vaccine effectiveness in preventing influenza hospitalization in two health systems in Michigan, where we have been conducting annual assessments of VE in various populations since 2008. We will conduct surveillance at two hospitals and will enroll adult inpatients with acute respiratory infection. Vaccination status will be reported and documented and considered with laboratory-confirmed influenza outcomes to estimate vaccine effectiveness for prevention of hospitalization. Analyses will use a test-negative design; those testing positive for influenza cases those testing negative are controls. Modifiers and confounders of vaccine effectiveness such as age, health status, high-risk health conditions, functional status, frailty, education, time from illness onset to specimen collection, calendar time, and propensity for vaccination will be assessed. In addition to our proposed influenza surveillance and VE assessment, we propose an estimation of the incidence of hospitalization in adults due to respiratory syncytial virus (RSV) and other respiratory viruses. This will allow for the evaluation of bias in influenza VE assessment due to interaction between influenza vaccination, infection, and non-influenza respiratory viruses, and will establish a platform for the future evaluation of RSV vaccines. We will accomplish these additional objectives by expanding our surveillance to months before and after the typical influenza season and evaluating specimens by molecular methods for RSV and other respiratory viruses.

Principal Investigator: Christine Neslund-Dudas, Ph.D., Public Health Sciences, and Michael Simoff, M.D., Pulmonary
Center for Research to Optimize Precision Lung Cancer Screening in Diverse Populations (UM1CA221939) Subcontract

Lung cancer, the most significant cause of cancer deaths in the US, is an urgent public health threat. It disproportionately affects populations that are already plagued by high poverty rates and low education levels. These populations experience both health disparities in the early diagnosis and treatment of cancer and are historically difficult to reach with cancer screening initiatives. While the results from the National Lung Screening Trial (NLST) indicated that low dose CT (LDCT) is an efficacious and cost-effective strategy for lung cancer screening (LCS), many uncertainties exist with respect to how patient, provider, health system,
and societal factors may impact the quality, compliance, effectiveness, and the risk of harms associated with lung cancer screening, within community-based health systems who serve diverse populations. Spanning from Pennsylvania to Hawaii and including five heterogeneous health systems with diverse populations, our proposed PROSPR Research Center, the Center for Research to Optimize Precision Lung Cancer Screening (CPLS), brings together a team of experienced, interdisciplinary researchers and clinicians with long-standing collaborative ties that is well-positioned to pursue research related to the barriers and opportunities associated with the implementation of LCS programs within community settings. The health systems within CPLS include: Henry Ford Health System in Metro Detroit, Kaiser Permanente Colorado, Kaiser Permanente Hawaii, Marshfield Clinic Health System in rural Wisconsin, and University of Pennsylvania Health System. The ultimate goal of CPLS is to identify critical gaps in the LCS process and to design innovative multilevel interventions to reduce lung cancer mortality, particularly among underserved populations. To achieve this goal, CPLS will complete the following specific aims: 1) build a comprehensive data ecosystem by pooling and linking common data elements to capture the entire LCS process and to assess the patient, provider, facility, health system, and societal factors that affect LCS; 2) leverage the CPLS data resource to conduct four high-impact, observational studies of the multilevel factors associated with the LCS process; 3) based on findings from Aims 1 and 2, develop and test interventions to address identifiable gaps in care that may lead to health disparities in LCS, 4) actively participate in Trans-PROSPR research initiatives and collaborate with external investigators via the use of publicly-available CPLS datasets. Our center focuses on the inclusion of diverse, underserved populations that are defined by multiple factors that may adversely impact access to, and utilization of, cancer screening. In response to both the Surgeon General's strong emphasis on the need to reduce lung cancer mortality and the Cancer Moonshot Blue Ribbon Panels focus on reducing the disproportionately high cancer death rates in underserved populations, CPLS will serve as a model for high-impact, translational research to reduce disparities in cancer mortality.

Principal Investigator: Laila Poisson, Ph.D.

Molecular and Clinical Evaluation of the Glioma Patient Experience to Anticipate Modern Outcomes and Guide Patient Care (R01CA222146)

Landmark papers published recently by us, and others, mark the new era of molecular diagnoses and precision therapy for glioma. In the summer of 2016, the World Health Organization (WHO) published updated diagnosis criteria for glioma that include molecular markers, taking a first step toward a molecularly precise diagnosis. It is our long-term goal to capitalize on the longitudinal resources of brain tumor banks to rapidly assess molecular hypotheses for prognosis and treatment of glioma. With the significant contribution of 240 cases from Henry Ford Hospital (HFH), an effort to molecularly and clinically profile glioma was started by The Cancer Genome Atlas (TCGA) project. Capitalizing on our clinically annotated brain tumor bank at HFH, we will focus on therapeutic outcomes, recurrent disease, and extended survival, which were not captured in the TCGA project. For this work, we have constructed an interdisciplinary team of collaborators, with clinical and informatics expertise, to profile an additional 340 glioma cases (WHO grade II-IV). In total, we will assess 700 tumor specimens (FFPE/frozen) from the HFH tumor bank (2001-present), representing both primary and matched progressive disease (Aim 1). Molecular data will be generated by exome sequencing to assess DNA sequence and copy number variants, targeted Sanger sequencing to profile the TERT promoter, and DNA methylation array assays to profile the methylome. Clinical annotation from our tumor bank, including long-term follow-up and therapy regimens, will be added to each of the 550 profiled glioma cases. The resulting comprehensively annotated tumor bank will be an invaluable resource for queries of clinical-molecular associations and the progression of disease, made available to researchers at HFH and beyond. In this proposal we use our database to address two analytical aims: (Aim 2) to carefully design statistical models of prognosis and therapy response among modern diagnosis classes using retrospective records; (Aim 3) to identify genomic differences, per patient, arising over the course of treatment and progression, which we expect will impact therapy decisions and inform standard treatments strategies. As part of the third aim, we will also explore the genomic patterns and clinical response of patients with exceptional survival, which may indicate differential molecular diagnosis or suggest therapeutic avenues for extending survival in others.
Principal Investigator: Benjamin Rybicki, Ph.D.
A New Prospective U.S. Cohort Set Within the Health Care System Institutions to Study Cancer (HHSN2612018000201)

Three mid-western integrated health care systems, HealthPartners (Minneapolis, MN), Henry Ford Health System (Detroit, MI), and Marshfield Clinic Health System (Marshfield, WI), here-forward known as the Great Lakes Consortium for Cohort Studies in Cancer (GLC3), have over a decade of experience working together as part of the NCI funded Cancer Research Network (CRN)(1-21) and its parent consortium the Health Care Systems Research Network (HCSRN)(22-27). These three integrated health care systems (IHCS) have joined together in response to the call by NCI to establish a U.S. cohort of healthy adults. The NCI U.S. Cohort will be recruited, consented on-line and followed for cancer-related outcomes during the ten year period of the contract. Biopspecimens and on-line questionnaires will be captured at baseline and at defined intervals which will be determined in a final protocol designed in collaboration between NCI, Information Technology support contractors, and participating integrated health care systems (IHCS). The overall goal of the NCI U.S. Cohort study is to enroll and follow 150,000 to 200,000 adult members of IHCS without cancer at the time of study enrollment. GLC3 proposes to recruit and enroll 20,000 health plan members (29% African American) across all three sites.

Principal Investigator: Jennifer Straughen, Ph.D.
The Prenatal Origins of Autism Spectrum Disorder (W81XWH191508)
The proposed study will leverage existing data, resources, and biologic samples as well as physician partnerships to conduct a study of 177 ASD cases (62 with archived placental tissue) and 62 frequency matched controls born in Michigan between 2012 and 2017. ASD cases are extensively evaluated by a team of experts at the Henry Ford Health System Center for Autism and Developmental Disabilities (CADD) and have detailed diagnostic information including severity level in their electronic medical record. Neonatal angiogenic profile will be measured in archived dried blood spots from Michigan’s Newborn Screening Program. Placental histopathology and angiogenic markers will be assessed in archived placental tissue from 62 cases and 62 typically developing controls frequency matched on year of birth, preterm status, and sex.

Impact: This proposal could provide evidence for the role of angiogenesis in the etiology of ASD while at the same time opening a door to neonatal risk assessment and earliest possible intervention. If successful, our findings may suggest approaches to minimize specific types of histopathology and/or changes in angiogenic profiles and mitigate their adverse effects in offspring. In addition, our findings may enable us to develop a clinically relevant placental screening tool that can be used for identification of children at highest risk of ASD so that interventions may begin as early as possible. Importantly, our collaboration with physicians at CADD who are currently diagnosing and treating ASD affected children and their families will facilitate the translation of research to practice.

Innovation: The proposed study is innovative not only in its focus on the placenta, but also its focus on processes that have not been traditionally examined in ASD research. The most innovative feature of the proposed study is its emphasis on markers of angiogenesis beginning in the prenatal period and extending into neonatal life. Despite its critical importance for neurodevelopment, ASD research to date has largely overlooked angiogenesis. This proposal is also innovative in that, if successful, it will simultaneously explain the sex specificity of ASD as well as the heterogeneous clinical expression of ASD symptoms. Furthermore, it will open the door to the development of noninvasive neonatal screening and diagnosis.

Principal Investigator: Ganesa Wegienka, Ph.D.
Epidemiology of Allergic Disease Endotypes (R01AI110450)

Pediatric allergy and asthma are a costly public health burden, but so far substantial research efforts have yielded no prevention strategies. A likely reason is that despite longstanding recognition by the medical community that the term ‘asthma’ refers to a collection of diseases, researchers have historically treated the syndrome as a single disease entity. Epidemiologically, the collapse of different phenotypes (observed disease patterns) and endotypes (phenotypes further delineated by pathophysiological processes), into a single category corrupts associations between risk factors and diseases. Thus, progress in allergic disease research has been hampered. Prior attempts have been made to identify such phenotypes and endotypes, but a combination of incomplete data and oversimplified statistical methods have limited progress. We propose to apply sophisticated latent class analyses in a large general risk cohort combined with...
immunological markers to finely discriminate asthma and allergy disease phenotypes and endotypes and then use this information to conduct risk factor analyses. Using this approach in our WHEALS birth cohort, we have already characterized four classes at age 2 years: 1) Low to No Sensitization; 2) Highly Sensitized; 3) Milk and Egg Dominated Sensitization; and 4) Peanut and/or Inhalant allergen – No Milk Sensitization. Total IgE levels varied between the groups, as did the rates of eczema and doctor diagnosis of asthma (at age 4 years). The Highly Sensitized had the greatest rates, the Low to No Sensitization had the lowest rates, and the other two classes had rates intermediate between the Low and High Sensitization groups. These data suggest the use of latent classes, rather than the use of the “traditional” definition of atopy (any allergen-specific IgE (slgE) _0.35 IU/mL), more specifically identifies those on a trajectory for allergic disease, yielding advancement in both allergic disease research and clinical care. Using the predominantly (62%) African American birth cohort WHEALS, we will: Aim 1) Determine which early life allergic disease phenotypes identified at age 2 years are associated with lung function (spirometry and methacholine challenge) at age 10 years; Aim 2) a) Identify the allergic disease endotypes for 10 year old children based on annual report of wheeze; lung function, eNO, obesity, cytokines, and white cell counts and extensive immunophenotyping [assessment of cellular markers to identify and quantify activation of regulatory T cells (Tregs), basophils and dendritic cells (DCs)] at age 10 years; and total IgE and sensitization (slgE and skin prick tests) at ages 2 and 10 years; and, b) Estimate associations between early life risk factors (e.g., delivery type, pet exposure, etc.) and the identified Aim 2a endotypes; and, 3) Compare and contrast the risk factor associations with the endotypes in Aim 2 to the risk factor associations determined using “traditional” definitions of atopy and asthma (doctor diagnosis and medication use and/or symptoms in the last year). Analyses will be performed for all 900 WHEALS cohort children and separately for Black children and White children to assess racial differences.

Principal Investigator: Ganesa Wegienka, Ph.D.
Environmental Risk Factors for Uterine Fibroids: A Prospective Ultrasound Study (R01ES028235)
Study of Environment, Lifestyle, and Fibroids (SELF) (HHSN273201600003I) Subcontract

Uterine leiomyomata (UL), or fibroids, are the most common neoplasms of the uterus and are a major source of gynecologic morbidity. In the United States (U.S.), the lifetime risk of symptomatic UL is approximately 25-30%. UL are the leading indication for hysterectomy, and UL-related costs exceed $34.4 billion annually. Black women are disproportionately affected by UL, with a 3-fold greater risk of diagnosis, earlier age at diagnosis and surgery, and more symptomatic tumors on average than white women. Despite the large public health burden of UL, little is known about its natural history or pathogenesis. Animal data and cross-sectional human studies have provided compelling preliminary evidence of a role for vitamin D in UL development and growth. Exposure to heavy metals such as lead, mercury, and cadmium is widespread, with reproductive-aged women, African Americans, and those of lower socioeconomic status having higher exposure levels than other groups. Funded by the National Institute of Environmental Health Sciences (NIEHS), the Study of Environment, Lifestyle and Fibroids (SELF) is a multi-year prospective cohort study of UL determinants in black women from the Detroit area. In 2011-2012, SELF enrolled 1,696 black women aged 23-34 years who had never been diagnosed with UL. At baseline and every 20 months for a total of 5 years (4 total clinic visits), SELF participants complete interviews, have blood collected for biological measurements, and undergo transvaginal ultrasounds for precise identification and mapping of UL at each visit facilitating accurate determination of UL development and growth (cohort retention >85%). The final planned clinic visits are underway. In this application, we propose to extend follow-up of SELF for an additional five years. One more clinic visit with transvaginal ultrasound, biospecimen collection and detailed exposure assessments via interview will be conducted to achieve the following specific aims: 1) Describe the natural history of UL initiation and growth; calculate age-specific UL incidence; and evaluate changes in tumor characteristics (size, number, and location) over a 10-year period; 2) Assess whether vitamin D status influences UL incidence and growth over a 10-year period; and 3) Evaluate the influence of selected environmental toxicants on UL incidence and growth. Specifically, we will examine the influence of active and passive cigarette smoking on UL incidence and growth; assess exposure to a panel of 13 metals and metalloids (and their mixtures) measured in whole blood and UL incidence and growth over a 10-year period; and determine whether vitamin D status modifies the associations between environmental toxicants and UL incidence. With its prospective design, population of young black women, serial ultrasounds, repeated collection of data on exposures and covariates, and careful analysis of chemical mixtures, SELF is ideal for identifying environmental risk factors for UL. Using methods that overcome the limitations of prior studies, this will be the most definitive study of modifiable environmental risk factors of UL and is likely to have high impact on science, clinical care, and public health policy.
The broad, long-term objective of this project is to enable patients with uterine fibroids (UF) to make informed decisions about management options based on the highest possible quality evidence. To help achieve this objective, we propose a multi-center registry of a geographically, racially, ethnically, and clinically diverse group of women who have received medical or surgical treatment for UF, Comparing Options for Management: Patient-centered Results for Uterine Fibroids (COMPARE-UF), designed to address the following specific aims: AIM 1) Develop the infrastructure necessary to implement large-scale observational comparative effectiveness research (CER) studies of management options for women with UF, including (a) a governance structure, policies, and procedures conducive to collaborative research involving patients, clinicians, methodologists, and other stakeholders, (b) an experienced Research and Data Coordinating Center, and (c) nine geographically diverse Clinical Centers (CCs) representing a broad range of patients and providers. AIM 2) Use this infrastructure to implement 3 projects addressing high-priority evidence gaps related to the effect of different management strategies on patient-centered outcomes. These include PROJECT 1: Comparing management options for symptom relief PROJECT 2: Comparing management options for preserving reproductive function PROJECT 3: Comparing effectiveness in different subpopulations. AIM 3) Evaluate innovative methods for the design, conduct, and analysis of observational comparative effectiveness research in this population. AIM 4) Translate research results into improved patient care, through both traditional peer-reviewed publications and collaborations with stakeholders to integrate the research findings into evidence-based patient decision making tools, clinical practice guidelines, and quality measures.

The average age for a woman to have her first child has been increasing for the last three decades in the United States, making our understanding of ovarian aging and its negative effect on the ovarian reserve, a measure of the capacity of the ovary to produce eggs capable of fertilization. Yet, we know very little about other factors in reproductive-age women that might affect the ovarian reserve, beyond aging itself. This proposal, titled Study of Ovarian Aging and Reserve in Young Women (SOAR), seeks to address the significant gap in our knowledge of factors, particularly modifiable factors, that affect ovarian reserve and might accelerate its decrease in young women. To achieve this goal, we will leverage the ongoing NIEHS Study of Environment, Lifestyle and Fibroids (SELF), which is following a cohort of 1,696 African-American women between the ages of 23-34 years over a five-year period. In this group of young women, we will assess changes in the ovarian reserve by tracking three different measures of the ovarian reserve: anti-Mullerian hormone (AMH), early follicular phase follicle-stimulating hormone (FSH), and antral follicle count (AFC). In addition to collecting survey data, we will also perform oral glucose tolerance testing (OGTT) and anthropometric and bioelectrical impedance analysis (BIA) measurements to more precisely determine the roles of glucose metabolism and obesity on the ovarian reserve. The results of our study will be clinically significant as we currently have limited longitudinal data for counseling women on risk factors for decreased ovarian reserve. Our study design is innovative in that we will use overlapping measures of the ovarian reserve and group-based trajectory modeling to determine correlates associated with decreased ovarian reserve. Specifically, we will determine the demographic, health-behavior, reproductive, and environmental factors associated with decreased AMH (as a measure of the ovarian reserve) over time (Aim 1), determine the association between various measures of obesity and decreased ovarian reserve (Aim 2), and determine the association between glucose dysregulation and decreased ovarian reserve (Aim 3). The proposed prospective longitudinal cohort study will determine the natural history and factors associated with the change in ovarian reserve over time. Further, it will add to the extremely limited data by generating the largest set of longitudinal data on AMH and ovarian reserve in the United States to date, which will benefit all women.
Infertility and spontaneous abortion (SAB) are significant public health problems, affecting up to 25% of reproductive age couples in the United States. Health care costs attributable to infertility and SAB exceed $5 billion per year, and several studies have shown an association between infertility treatments and adverse pregnancy outcomes. Thus, identifying modifiable risk factors for subfertility and SAB is an important public health goal. The potential effects of exposure to endocrine-disrupting chemicals (EDCs) on risk of subfertility and SAB are understudied. The few existing human studies have limitations including small sample size, enrollment after conception, retrospective study design, suboptimal assessment of exposure and outcome, inadequate control for potential confounding variables, and limited generalizability. The proposed study will prospectively assess the relation of exposure to selected EDCs, including phenols, phthalates, and per- and poly-fluoroalkyl substances (PFAS), to risk of subfertility and SAB in a preconception subcohort of 950 pregnancy planners. We will use data from two NICHD-supported prospective cohorts of pregnancy planners in North America and Denmark. With web-based recruitment and data collection, we have enrolled over 17,000 women attempting pregnancy into these cohorts. In a subset of 200 participants, we have successfully pilot tested in-person collection of urine and blood specimens during the preconception and early pregnancy periods. In this application, we propose to expand in-person biospecimen collection, increasing the number of women with preconception and early pregnancy urine and blood samples from 200 to 950. At each of our three biospecimen collection sites (Boston, Detroit, and Aarhus), we will enroll 250 women and collect three urine samples and one blood sample (in preconception) and three urine samples and one blood sample (in early pregnancy). We will ship the samples to the CDC for the analysis of urinary phthalates, urinary phenols, and serum PFAS. To increase cost efficiency, we will pool three urine samples in each exposure window before assaying for phthalates and phenols; and we will assay one preconception blood specimen for PFAS, a persistent chemical, in analyses of subfertility and SAB. Finally, we will conduct a pilot study among 100 U.S. participants to assess the feasibility of collecting urine by mail, which would allow us to take advantage of our full geographically diverse cohort in the future. Strengths of this application include the prospective design, preconception enrollment of pregnancy planners, repeated measurement of exposure during preconception and early pregnancy, excellent control for confounding via bimonthly prospective data collection on a wide range of covariates and use of the latest analytic methods for mixtures modeling. We have generated compelling preliminary data to support our aims. The present grant is cost-effective in leveraging already-established cohort studies with data collection and follow-up supported by other grants. The results generated will be translatable by directly informing future regulatory decisions about EDC standards in a manner that could reduce rates of infertility and SAB.